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Biochemical and Structural Studies of Yeast Vps4 Oligomerization

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Abstract

The ESCRT pathway functions in vesicle formation at the multivesicular body, the budding of enveloped RNA viruses such as HIV-1, and the final abscission stage of cytokinesis. As the only known enzyme in the ESCRT pathway, the AAA ATPase Vps4 provides the energy required for multiple rounds of vesicle formation. Like other Vps4 proteins, yeast Vps4 cycles through two states: a catalytically inactive disassembled state that we show here is a dimer, and a catalytically active higher order assembly that we have modeled as a dodecamer composed of two stacked hexameric rings. We also report crystal structures of yeast Vps4 proteins in the apo- and ATPyS-bound states. In both cases, Vps4 subunits assembled into continuous helices with six-fold screw axes that are analogous to helices seen previously in other Vps4 crystal forms. The helices are stabilized by extensive interactions between the large and small AAA ATPase domains of adjacent Vps4 subunits, suggesting that these contact surfaces may be used to build both the catalytically active dodecamer and catalytically inactive dimer. Consistent with this model, we have identified interface mutants that specifically inhibit Vps4 dimerization, dodecamerization, or both. Thus, the Vps4 dimer and dodecamer likely form distinct but overlapping interfaces. Finally, our structural studies have allowed us to model the conformation of a conserved loop (Pore Loop 2) that is predicted to form an argininerich pore at the center of one of the Vps4 hexameric rings. Our mutational analyses demonstrate that Pore Loop 2 residues Arg241 and Arg251 are required for efficient HIV-1 budding, thereby supporting a role for this "arginine collar" in Vps4 function.

Keywords

Vps4; AAA ATPase; Oligomerization; MVB pathway; X-ray Crystallography

Introduction

The processes of vesicle budding into multivesicular bodies (MVBs), retrovirus release, and cytokinesis share a common requirement for an overlapping subset of cellular machinery, termed the Endosomal Sorting Complexes Required for Transport (ESCRT)^{1; 2; 3}. The ESCRT machinery may perform analogous functions at the final stages of all three processes, where a membrane fission event is required to resolve a thin, cytosol-containing membrane

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tubule. At steady state, ESCRT factors bind weakly to membranes throughout the cell⁴, but they can be recruited to function at specific sites of membrane remodeling including the late endosome, the plasma membrane, and the midbody. Most ESCRT factors function as subunits of one of four different complexes, termed ESCRT-0, -I, -II, and -III. Although their functions are not yet fully defined, ESCRT-0, -I, and -II appear to act as adaptors, binding directly to both membranes and protein cargoes as they are sorted into vesicles or virions. ESCRT-III subunits, in contrast, appear to co-assemble into membrane-associated filaments that play a more direct role in membrane remodeling, possibly mediating membrane extrusion and/or fission⁵.

Once assembled on membranes, the ESCRT machinery is released by the action of the Vps4 ATPases, the only known enzymes in the ESCRT pathway. A direct role for Vps4 function in the ESCRT pathway is indicated by a number of observations, including: 1) Vps4 localizes to endosomal membranes⁶ and to the midbody during cytokinesis^{7; 8}, 2) MVB protein sorting and intralumenal vesicle formation are inhibited in the absence of Vps4 or upon expression of dominant negative Vps4 that cannot bind or hydrolyze ATP^{6; 9; 10; 11; 12}, and ³) Depletion or dominant inhibition of Vps4 blocks late stages of both cytokinesis^{7; 8} and enveloped virus budding (reviewed in ^{13; 14; 15}). At a minimum, Vps4 ATPase activity appears to be required to release and recycle the assembled ESCRT machinery, because the other ESCRT factors assemble but are not released in the absence of Vps4 activity¹⁶. Vps4 activity may also be mechanistically coupled to membrane deformation and/or fission, though such coupling has not been demonstrated experimentally.

Vps4 proteins belong to the large and diverse family of AAA ATPases (<u>ATPases Associated</u> with diverse cellular <u>Activities</u>) (reviewed in ¹⁷; 18; 19; 20; 21; 22; 23). AAA ATPases can contain either one (class I) or two (class II) ATPase cassettes, and Vps4 is a class I ATPase. Yeasts express a single Vps4 protein, whereas humans and other mammals express two closely related Vps4 proteins, termed VPS4A and VPS4B/SKD1²⁴. As summarized in Fig. 1a, Vps4 proteins contain an N-terminal MIT domain that binds ESCRT-III protein substrates, a central ATPase cassette composed of large and small domains, and a three-stranded antiparallel sheet (the β domain) inserted within the small ATPase domain ²⁵; 26; 27; 28. On its own, the linker between the MIT and ATPase regions lacks a persistent structure²⁶, and can be hyper-susceptible to proteolysis in the context of the full length protein. This linker undergoes nucleotide-dependent changes in proteolytic susceptibility²⁸, however, suggesting that it serves as a semi-flexible tether that can adopt different conformations.

Most AAA ATPases function as closed hexameric rings, although other arrangements are known²⁹. Ring formation is mediated by the AAA ATPase cassette, and ATP binding typically promotes ring assembly because nucleotides bind between and contact adjacent subunits¹⁸. Vps4 is typical in this regard because ATP binding promotes enzyme assembly^{16; 25}, and Vps4 appears to have a canonical ATP binding site (although high resolution structures of ATP-bound Vps4 proteins have not previously been reported). Upon ATP binding, Type I ATPases generally assemble into single ring structures, whereas Type II ATPases generally form double ring structures in which the two ATPase cassettes form separate, stacked rings (e.g., 21; ³⁰). The Type I Vps4 enzymes are atypical in this regard in that they appear to form double stacked rings^{16; 25; 27; 31}. Crystal structures of double ring Vps4 complexes are not yet available, and current models for the stoichiometry and structures of this assembly have therefore been deduced from gel filtration ^{16; 25}, protein crosslinking^{16; 31}, mutagenesis²⁵, single particle cryo-EM reconstructions^{27; 31}, and modeling studies^{25; 27}. Vps4 proteins also bind an activator, Vta1/SBP1/LIP5, which contacts the β domain and promotes assembly and ATPase activity^{25; 31; 32; 33; 34; 35; 36; 37; 38}. Recent biochemical and EM studies of the yeast Vps4-Vta1 supercomplex suggest that Vta1 may bind just one of the two rings in the double-ring assembly³¹.

Although there is general agreement that Vps4 enzymes drive the ESCRT pathway by cycling between an inactive, low molecular weight state and an active, high molecular weight state, uncertainties surround the structures and stoichiometries of both the active and inactive conformations. For example, the inactive state of Vps4 has been described as both a monomer²⁸ and a dimer^{16; 25; 27}, and the nature of the putative dimer is not clear. Similarly, two different single particle reconstructions of the fully assembled core enzyme at moderate resolution have been interpreted as showing a dodecamer³¹ or a tetradecamer²⁷, and these studies also differ on other aspects of the global architecture. In an effort to characterize the architectures and oligomeric states of Vps4 further, we have determined two new crystal structures of the *S. cerevisiae* Vps4 core enzyme, performed analytical ultracentrifugation analyses that define the oligomeric state of the inactive Vps4 complex, and carried out extensive mutagenesis aimed at discriminating between different Vps4 assembly models and identifying residues that perform key functions in the active and inactive states.

Results

Structure Determinations

Two different Saccharomyces cerevisiae Vps4 constructs that spanned residues 104-437 and 122-437 were expressed in E. coli, crystallized, and structurally characterized by X-ray crystallography using the human VPS4B structure as an initial search model²⁵. Both constructs lacked N-terminal MIT domains and segments of the ensuing linker (see Fig. 1), and both also contained the E233Q mutation, which allows ATP binding but blocks ATP hydrolysis¹⁶. Vps4₁₂₂₋₄₃₇ crystallized in space group P6₅22, with one protein molecule per asymmetric unit and a sulfate ion bound at the active site. This structure was refined at a resolution of 2.7 Å and an R_{free} value of 28.7% (Table 1). Vps4104-437 crystallized in space group P212121 with three molecules per asymmetric unit and was refined at a resolution of 3.25 Å to an Rfree value of 28.9%. Molecules A and B showed good density for ATP_YS at the active site (Fig. 2), whereas molecule C also had active site electron density but clearly lacked density for the adenine nucleoside. The density could be adequately modeled as an ethylene glycol molecule in complex with a magnesium ion, and this is therefore our tentative assignment for the bound ligand in this case. Crystallographic statistics are summarized in Table 1, and both structures are generally well ordered, albeit at modest resolution. As detailed in Table 2, the ordered regions of all constructs begin at approximately residue 122, and we therefore refer to the proteins interchangeably as $Vps4_{\Delta MIT}$, with their bound ligands designated as "SO₄" (crystal form 1), "ATPyS" (molecules A and B in crystal form 2), and "ethylene glycol-Mg²⁺" (molecule C in crystal form 2).

Vps4 Structures

As expected, the Vps4 structures reported here resemble published structures of human VPS4B $_{\Delta MIT}$ -SO $_4^{25}$, yeast (His) $_6$ Vps4 $_{\Delta MIT}^{27}$, and yeast Vps4 $_{\Delta MIT}$ -ADP²⁸. For example, the ATPase cassettes of our yeast Vps4 $_{\Delta MIT}$ -SO $_4$ and human VPS4B $_{\Delta MIT}$ -SO $_4$ structures overlap with a C α root mean square deviation (RMSD) between ordered residues of just 1.2 Å (Fig. 1c). Nevertheless, our two new structures have several ordered loops that have not been seen in previous structures, and the new structures extend the existing ranges of bound ligands, interdomain angles, and intermolecular packing interactions.

As in other AAA ATPases, the ATPase cassette of Vps4 is composed of two domains. The larger amino-terminal domain has six α -helices (helices 1–5 and 10) that pack on either side of a six-stranded β sheet. The first strand of the sheet (β'), which is found in only a subset of AAA ATPases, runs antiparallel to the remaining five strands (strands 1–5). The smaller carboxyl terminal domain is an antiparallel four-helix bundle (helices 6–9) (Fig. 1b). The two domains are structurally distinct, although the final helix extends away from the smaller domain

to pack along one edge of the large domain sheet. Vps4 is unique amongst known AAA ATPases in that a three-stranded β -sheet, called the β domain, is inserted between helices 8 and 9 of the four-helix bundle. The β domain contributes to Vta1 binding²⁵ and enzyme assembly³⁹.

Both of our Vps4 crystal forms were grown in the presence of $ATP\gamma S$, and two of the three molecules in crystal form 2 (A and B) show clear electron density for bound ATPyS nucleotides (Figures 2a, b). In these two molecules, the nucleotide binds in a site between the large and small domains, and makes all of the expected contacts with the Walker A/P Loop (¹⁷³GPPG<u>TGKSY</u>¹⁸¹), Walker B (²³⁰FIDEVD²³⁵), and sensor 1 (²⁷⁵ATN²⁷⁷) elements¹⁸ (Figure 2a). In particular, the adenine ring adopts the anti conformation, and stacks between Tyr181 (Walker A) and Met307 (helix 6) 40; 41; 42; 43; 44. The ATP γ S triphosphate hydrogen bonds with backbone amide NH groups in the P Loop (¹⁷³GPPGTGKSY¹⁸¹, underlined residues make H bonds), and the phosphates also contact the conserved Lys179 (Walker A/P Loop, β -phosphate salt bridge), Asn277 (sensor 1, γ -sulfate) and Pro175 (P Loop, γ -sulfate). A bound Mg²⁺ ion is coordinated by oxygen atoms from the β and γ phosphates, and by the conserved side chains of Ser180 (P Loop) and Asp232 (Walker B). The structure is consistent with the observation that the K179A mutation inhibits ATP binding (and higher order assembly)¹⁶. The Walker B element also contains another highly conserved acidic residue (Glu233), which is thought to function as the catalytic base that activates a nucleophilic water^{16; 25}. This residue was mutated to Gln in the Vps4_{Δ MIT} constructs to minimize ATP hydrolysis, and the Gln233 side chain projects out of the active site where it contacts Ser198 and Ser200. We have not attempted to compare the positions of the active site nucleotides and side chains with those of the ADP-bound Vps4 structure because both complexes were solved at modest resolution and because the bound ADP ligand has a very high B factor $(179 \text{ Å}^2)^{28}$.

Our remaining two Vps4 structures do not exhibit clear density for bound ATP γ S nucleotides. Specifically, Molecule C in crystal form 2 has density in the phosphate binding sites that likely corresponds to an ethylene glycol molecule (which presumably came from the 48 % ethylene glycol crystallization solution) in complex with a magnesium ion, which is coordinated by the conserved Asp232 and Ser180 side chains and a hydroxyl group of the ethylene glycol. The single unique molecule in the VPS4_{ΔMIT}-SO₄ crystal form also lacks nucleotide density, but has density within the β and γ phosphate binding sites that likely corresponds to a sulfate ion from the 1.6 M ammonium sulfate crystallization solution. A sulfate ion is also seen in the same position in the structure of human VPS4B_{ΔMIT}-SO₄²⁵.

The two different crystal forms reported here extend the range of interdomain angles seen for Vps4, and further show how the different domains can move relative to one another (Fig. 3 and Table 2). As has been noted previously²⁸, relative domain motions are centered about two different hinges. The first hinge is Pro297, which is located within the linker between the two domains of the ATPase cassette. Molecule A in crystal form 2 has the most closed hinge I angle seen to date (115°), whereas molecule C has the most open angle (134°). The second hinge is evident in overlays of the core of the small domain (Fig. 3 and data not shown). This hinge angle can vary by up to 16°, and this variation allows the β domains to project in different directions from the small ATPase domain, which may be required to accommodate different β domain packing arrangements in the two rings of the dodecamer³¹.

As in all previously reported Vps4 crystal structures, the subunits in our two crystal forms pack as continuous helices that repeat every six subunits. In crystal form 1, the helix follows the 6_5 crystallographic screw axis, and every subunit is therefore in an equivalent position. In crystal form 2, the two ATP γ S-bound Vps4 molecules are nearly equivalent, whereas the more open third molecule sits in a slightly different position within the spiral. Analogous 6-fold (or

pseudo 6-fold) screw arrangements have been seen in the crystal structures of other AAA ATPases, which presumably reflects their propensity to form six membered rings⁴⁵; 46. As discussed below, adjacent helices are connected by different types of two-fold symmetric packing interactions in the different Vps4 crystal forms.

Nucleotide-free Vps4 Dimerizes in Solution

The oligomeric state of the inactive, low molecular weight Vps4 complex has been controversial, and we therefore performed gel filtration and analytical ultracentrifugation analyses to examine the oligomerization of nucleotide-free Vps4 proteins. Pure recombinant Vps4 (MW=48,803 g/mol) and Vps4_{Δ MIT} (MW=37,700 g/mol) proteins eluted from a Superdex S200 gel filtration column as single species with apparent molecular weights of 127 kDa and 88 kDa, respectively. Thus, the Stokes radii of both constructs were most consistent with dimers, but could possibly be explained by extended monomers. Equilibrium sedimentation analyses were used to distinguish between these two possibilities because this approach provides shape-independent measures of protein solution mass.

As shown in Fig. 4 and Supplemental Fig. 2, the radial distributions of full length Vps4 (panel 1) and Vps4_{AMIT} (panel 2) fit well to single-species dimer models, but not to monomer or trimer models. In each case, sedimentation data were collected at three different protein concentrations and two rotor speeds, yielding a total of six data sets for each protein. The six data sets were globally fit to single species models in which the molecular weight was allowed to float, and also to monomer, dimer, or trimer models with fixed molecular weights. The distribution of Vps4 fit a protein with a predicted mass of 96,210 g/mol, which matched the mass expected for a Vps4 dimer ($M_{obs}/M_{monomer} = 1.97$). Data for three initial concentrations of Vps4 centrifuged at 12,000 RPM are shown together with the global fits (Fig. 4, panel 1), and the small, random residuals indicate that the data were satisfactorily fit by a simple single species model (lower panels). Global fits to the three data sets collected at 16,000 RPM were also satisfactory (data not shown). Similarly, when fixed monomer, dimer, or trimer molecular weights were used during the global fitting procedure, the dimer model was clearly the best fit to the data (Supplemental Fig. 2). Likewise, sedimentation data for the $V_{ps4_{AMIT}}$ protein estimated a solution mass of 82,435 g/mol ($M_{obs}/M_{monomer} = 2.19$, Fig. 4, panel 2), and the dimer model was also the best fit of the different possible fixed molecular weight models (Supplemental Fig. 2).

As in most of our other experiments, these analyses utilized Vps4 proteins with E233Q mutations because the constructs were also used to examine the effects of ATP binding in the absence of hydrolysis. To ensure that the E233Q mutation did not alter the oligomerization state, gel filtration analyses were also performed on wild type Vps4_{Δ MIT} proteins (both Vps4₁₀₁₋₄₃₇ and Vps4₁₂₄₋₄₃₇). In both cases, these proteins eluted with the same retention times as their E233Q analogues, indicating that the E233Q mutation did not affect the oligomerization state (not shown). We therefore conclude that in the absence of nucleotide, Vps4 and Vps4_{Δ MIT} proteins form stable dimers in solution.

The Vps4 Dimer Interface

We reasoned that recurring crystal packing interactions might recapitulate aspects of Vps4 dimerization in solution, and therefore examined all of the intermolecular lattice contacts in the different known Vps4_{Δ MIT} crystal forms. The different interfaces are summarized in Table 2 and Supplemental Figure 1, and subsets of the interfaces are shown in Fig. 5a. The various yeast Vps4 crystal lattices show six different classes of dimer interfaces, five of which exhibit two-fold rotational symmetry (Interfaces 1–5) and one of which lacks symmetry (Interface 6). Interface 1, the largest of the two-fold symmetric interfaces, involves contacts between residues in helix 3 and strand 2 in adjacent large domains. This interface is seen in three different crystal

forms, and buries an average of $620\pm130\text{\AA}^2$ /subunit. Variants of Interface 1 were seen in our Vps4 $_{\Delta MIT}$ -SO₄ crystals (480Å²) and also in two other Vps4 $_{\Delta MIT}$ crystal forms, one with a bound sulfate (2QP9) and one without a bound ligand (2RKO) (740Å² and 640Å², respectively). We note that although similar contact surfaces are used to create Interface 1 in these three crystal forms, the interactions differ significantly in detail (see Supplemental Figure 1).

Hartmann et al. have argued that the crystallographic Vps4 Interface 1 also forms in solution based upon their report that the Q216A interface mutation blocks Vps4_{Δ MIT} dimerization in solution²⁷. In our hands, however, the Q216A mutation did not alter the gel filtration mobility of Vps4_{Δ MIT}, indicating that this mutation did not inhibit dimer formation in solution (Fig. 5b, panel 1, compare red and black chromatograms). This conclusion was confirmed in analytical ultracentrifugation experiments, which showed that the Vps4_{Δ MIT}, Q216A protein remained dimeric over the entire concentration range tested (Fig. 4, panel 3, MW_{obs} = 71,668 g/mol, MW_{obs}/MW_{monomer} = 1.90). The Vps4_{Δ MIT} mobility was similarly unaffected by two other Vps4 Interface 1 mutations (N224A and M220D, data not shown), and we therefore conclude that Interface 1 does not mediate Vps4 dimerization in solution.

Four other classes of two-fold symmetric subunit contacts were seen in at least two different Vps4_{Δ MIT} crystal forms (Interfaces 2–5). However, these interfaces were approximately half the size of Interface 1, with average buried surface areas of: 350±70, 280±130 Å², 320±150 Å², and 360±120 Å², respectively. None of these interfaces occurred in all of the reported crystal forms, and the residues within Interfaces 2, 4, and 5 are not well conserved across species. Interface 3 residues are well conserved, but this is likely explained by the fact that they form the "Rossmann fold" element of the nucleotide binding site. These observations imply that Interfaces 2–5 are also unlikely to mediate Vps4 dimerization in solution.

Interface 2 connects the two small domains of Vps4 and primarily involves contacts between residues in helices 7 and 9 in adjacent small domains. This interface is present in our Vps4_{Δ MIT}-SO₄ crystal structure (buried surface area, 330Å²), and also in Vps4_{Δ MIT} crystals with a bound sulfate (2QP9) and without a bound ligand (2RKO) (430Å², and 290Å², respectively). Xiao et al. reported that two mutations within this interface (M330D and L407D) did not alter the gel filtration mobility of Vps4_{Δ MIT}²⁸. In good agreement with this report, we also found that the L407D mutation had no effect on the gel filtration mobility of Vps4_{Δ MIT} (Fig. 4b, panel 2). These data confirm that Vps4 Interface 2, like Interface 1, does not mediate Vps4 dimerization in solution.

These observations indicate that none of the symmetric dimer interfaces seen in more than one Vps4 crystal form are likely to mediate Vps4 dimerization in solution. We therefore considered the possibility that Vps4 might form an asymmetric dimer in solution. In particular, all known Vps4 crystal forms contain an asymmetric sixth interface that is larger than any of the two-fold symmetric interfaces (760±80 Å²). Interface 6 connects the small domain of one subunit with the large domain of an adjacent subunit, and is used to create the continuous helix of Vps4 molecules. The interface is complex, and involves contacts between helix 1 and the helix 1/ strand 1 loop on one side of the interface, and helices 7–9 and the C-terminal edge of the β domain on the other. Additional contacts between adjacent large domains are also made between the two molecules that bind ATP- γ S in our crystal form 2 (between helix 5 on one side of the interface and the strand 2-helix 3 loop on the other). These additional contacts result from closure of the interdomain angle upon ATP γ S binding (see Fig. 3).

The requirement for Interface 6 in Vps4 dimerization was tested using four different point mutations (L151D, I351A, I354D, and W388A). We initially changed the highly buried L151, I351 and I354 residues to aspartate in order to robustly disrupt the interface. However, the

I351D protein was only partially soluble, and we therefore replaced this residue with alanine. W388 was changed to alanine rather than aspartate because this dramatic reduction in side chain volume was expected to yield a sufficient change in interface stability. Strikingly, every one of these mutations reduced the gel filtration mobility of Vps4_{ΔMIT} to that expected for a monomer (compare red and black curves in Fig. 5, panels 3 (L151D) and 4 (W388A), and similar data not shown for the I351A, I354D mutants). Analytical ultracentrifugation experiments confirmed that Vps4_{ΔMIT}, L151D was indeed monomeric in solution (Fig. 4, panel 4, $MW_{obs} = 38,815$ g/mol, $MW_{obs}/MW_{monomer} = 1.03$)). Thus, residues used to create the asymmetric Interface 6 seen in all known Vps4_{ΔMIT} crystal forms are also required for solution dimerization. As discussed below, we propose that the Vps4 solution dimer interface is likely to be a variant of Interface 6 in which the subunit conformations are altered so that the assembly does not propagate into a polymer.

Requirements for Vps4 Dodecamerization

Mutant Vps4_{Δ MIT} constructs that can bind but not hydrolyze ATP (e.g., VPS4_{Δ MIT}, E233Q) form stable higher order assemblies that have been modeled as double rings, with six³¹ or seven²⁷ subunits per ring. Conversion of Vps4_{Δ MIT} dimers into higher order structures (here termed dodecamers) can also be analyzed by gel filtration chromatography (Fig. 6, panel 1), which provides a convenient assay for mutations that inhibit Vps4 ring formation.

We previously proposed a homology model for one of the two rings of the Vps4_{ΔMIT} dodecamer. This model was created by substituting Vps4 subunits in place of the crystallographically characterized p97 D1 subunits in their hexameric ring conformation²⁵; ⁴⁷. This model for the Vps4 ring generates subunit interfaces that are similar, but not identical to the crystallographically defined Vps4 Interface 6. Differences between the crystallographic interface and the homology model are illustrated in Fig. 6a, where the upper model shows the p97-based homology model and the lower model shows the spiraling "hexamer" of crystal form 1 (crystal form 2 is very similar). The p97 D1 homology model predicts that ring formation will be dependent upon residues located at the center of crystallographic Interface 6 (e.g., Leu151), but not upon residues located at the exterior of Interface 6 that are separated when the subunit orientation is adjusted to match the p97 D1 ring (e.g., Trp388, see insets). We therefore probed the validity of the p97 D1 homology model by testing the assembly properties of VPS4_{ΔMIT} proteins with L151D and W388A point mutations.

As expected, the Vps4 $_{\Delta MIT, L151D}$ mutant remained monomeric in both the presence and absence of ATP, indicating that this mutation blocks both dimerization and dodecamerization (compare panels 1 and 2 in Fig. 6b). We note that the inability of this mutant to assemble into any higher order structure implies either that formation of the second ring in the Vps4 dodecamer depends upon forming the first ring (our preferred explanation) or, alternatively, that Leu151 makes important contacts in both rings of the dodecamer. In contrast, the Vps4_{ΔMIT, W388A} mutant was monomeric in the absence of ATP, but assembled into stable dodecamers upon addition of ATP (panel 3). This result supports the idea that one of the two rings in the Vps4_{AMIT} dodecamer resembles the hexameric ring formed by the homologous p97 D1 domain. This experiment also indicates that the Vps4_{AMIT} dimer is not an obligate onpathway intermediate for forming the Vps4 dodecamer, because the W388A mutation inhibits dimerization but not dodecamerization. Conversely, we previously showed that the Vps4_{R352A} mutant dimerized normally, but failed to form higher order assemblies in the presence of ATP²⁵. Arg352 sits within the subunit interface of our p97 D1 homology model, but is exposed at the edge of Interface 6 in the different Vps4 crystal structures. Thus, W388A and R352A are examples of mutations that allow formation of one Vps4 state but inhibit formation of the other, and reflect the use of overlapping but distinct interfaces for dimer and dodecamer formation.

A Functional Role for Pore Loop 2

Our model for the p97 D1 homology model for the hexameric ring of Vps4 places two different loops near the center of the narrow pore (Fig. 7). Analogous loops are also present in other AAA ATPases, and they have been termed Pore Loop 1 (²⁰⁶WMG²⁰⁸, green in Figures 7a, b) and Pore Loop 2 (²⁴¹RGEGESEASRR²⁵¹, blue) ⁴⁸; ⁴⁹; ⁵⁰. A section of Pore Loop 2 is also known as the "Arginine Collar", owing to the presence of three conserved arginine residues (Arg241, Arg250, and Arg251 in Vps4)⁵¹; ⁵²; ⁵³. Unlike previous Vps4 structures, Pore Loops 1 and 2 are both well ordered in the three crystallographically independent molecules of Vps4 crystal form 2. Pore Loop 1 forms a short two turn helix between strand 2 and helix 3, and the conserved Trp206/Met207 dipeptide forms a hydrophobic patch that sits above the pore in the hexamer model, with Met207 projecting into the central cavity (green in Figs. 7a, b). Pore Loop 2 forms an extended loop between strand 3 and helix 4, and the conserved Pore Loop 2 Arg residues at positions 250 and 251 form a positively charged "collar" that sits immediately beneath the hydrophobic Pore 1 residues (blue in Fig. 7a, upper panel). Charged Pore 2 residues also serve to restrict the diameter of the pore itself to ~10 Å, with Glu243 and 245 (red), forming a negatively charged ring on the Pore Loop 1 side, and Arg241 (blue), forming a positively charged ring on the opposite side (Fig. 7a, lower panel).

We have shown previously that point mutations in Pore Loop 1 residues of both human VPS4A and VPS4B inhibit HIV-1 release and infectivity, implying that this element is important for Vps4 protein function²⁵. To examine the functional importance of Pore Loop 2 residues, we assayed the efficiency of HIV-1 release and infectivity from cells expressing exogenous wild type VPS4A or VPS4A proteins with mutations in the two most highly conserved Pore Loop 2 arginines (R236A or R246A in VPS4A), the central Pore Loop 2 glutamate (E240A), or the ATP binding site (K173Q, positive control)¹⁶. As shown in Figure 7c, overexpression of VPS4A_{R236A} and VPS4A_{R246A} inhibited the release of an HIV-1 vector, as indicated by reductions in the levels of the virion-associated MA and CA proteins released into the supernatant (upper panel, compare lanes 4 and 6 to lane 3) and in viral titers (lower panel). In both cases, titer reductions were substantial (48- and 73-fold, respectively, note the log scale in Fig. 7c). Virion release and titers were also impaired by overexpression of the VPS4A_{E240A} mutant, but in this case the reduction was modest (4-fold). Control experiments behaved as expected in that overexpression of the VPS4AK1730 ATP binding mutant strongly inhibited virion release and titer (lanes 7, 470-fold titer reduction), overexpression of the wild type VPS4A protein did not impair virion release (compare lanes 2 and 3), and cellular expression levels of the HIV-1 CA and GFP-VPS4A proteins were equivalent in all cases (panels 2 and 3, respectively). We also demonstrated that the analogous Pore Loop 2 point mutations did not affect yeast Vps4E2330 protein dodecamerization in vitro, implying that these residues are not critical for Vps4 assembly (data not shown). We therefore conclude that the conserved arginine residues of Pore Loop 2 play important functional roles in Vps4 enzymes, and that the central Glu residue also contributes, albeit to a lesser extent.

DISCUSSION

Vps4 Oligomerization

At steady state, Vps4 is distributed throughout the cytoplasm in an inactive state, but can be recruited to membrane sites of action where it binds ATP and associates into enzymatically active higher order complexes¹⁶. Our studies imply that Vps4 is likely a dimer in its inactive state. Specifically, we found that Vps4 forms stable dimers in solution, as evidenced by the unambiguous fit of equilibrium sedimentation data to a single species dimer model and by the identification of a series of different point mutations that block Vps4_{ΔMIT} dimerization. As is true for any dimeric protein, however, Vps4 will dissociate into monomeric species at sufficiently low protein concentrations. Indeed, we observed both monomeric and dimeric

species for the Human VPS4B protein²⁵, presumably because the human VPS4B homodimer is less stable than the yeast Vps4 homodimer. Moreover, we observed that even the yeast Vps4 protein exhibited concentration-dependent mobility in gel filtration experiments performed at low micromolar concentrations, indicating that monomeric yeast Vps4 species can also be detected at low protein concentrations under non-equilibrium conditions (not shown). These observations raise the possibility that monomeric Vps4 species may accumulate in vivo if the protein concentration is sufficiently low. Nevertheless, the dissociation constant for the yeast Vps4 dimer must be less than ~5 μ M in order to allow an adequate fit by a single species dimer model over the concentration ranges tested (Fig. 4 and data not shown).

ATP binding promotes formation of higher order Vps4 assemblies, provided that nonhydrolyzable ATP analogs such as ATP γ S are used, or that the Vps4 protein is mutated to prevent nucleotide hydrolysis (or both). These higher order assemblies appear to be dodecamers composed of two non-equivalent hexameric rings. The conclusion that VPS4 likely forms hexameric rings is based upon: 1) six-fold symmetry observed in cryo-EM reconstructions of assembled Vps4_{ΔMIT} and Vps4 complexes³¹, 2) the finding that equilibrium sedimentation data for the Vps4 assembly is best fit by a dodecamer model²⁵, and 3) strong homologies between Vps4 and the related p97 D1 and spastin proteins, both of which form hexameric rings⁵²; 54; 55. The Vps4 dimer/oligomer assembly cycle is therefore reminiscent of NtrC, another AAA ATPase that also forms catalytically inactive dimers, and then assembles into active hexameric rings upon ATP binding⁵⁶; 57. In both cases, the dimers are "off pathway" with respect to hexameric ring assembly, and therefore appear to represent autoinhibited states that negatively regulate enzymatic activity. There is no indication, that the inhibitory dimers of NtrC and Vps4 proteins are structurally related, however, because the NtrC dimer subunits associate in a two-fold symmetric, head to tail orientation that is completely distinct from the orientations of subunits in the hexamer.

Models for Vps4 Assemblies

All Vps4_{AMIT} crystal forms reported to date show a similar packing arrangement in which the subunits form a continuous helix with a six-fold screw axis. These helices, in turn exhibit different types of two-fold symmetric packing interactions in different crystal forms, but our experiments indicate that none of these symmetric crystallographic interfaces is likely to mediate Vps4 dimerization in solution. In contrast, Vps4 dimerization is inhibited by a series of different mutations within the interface used to form the continuous helix (e.g., I351A, I354D, L151D and W388A). This leads to the surprising conclusion that the Vps4 dimer is likely formed by an asymmetric interface. We suggest that this interface likely resembles, but is not identical to, the asymmetric dimer interface that mediates Vps4 helix formation in all crystal forms reported to date. If correct, this model begs the question why the interaction does not repeat infinitely to form a helical (or linear) polymer in solution. Our preferred explanation is that the two subunits of the asymmetric dimer have different interdomain angles, making the dimer incompatible with repeating interactions. This type of asymmetric interaction is hinted at in the crystal structure of the Vps4-ATPyS complex, where differing interdomain angles in the three different subunits produce distinct dimeric interactions between the A:B, B:C, and C:A molecule pairs. However, a more precise understanding of this structure will clearly require high resolution crystals of a discrete Vps4 dimer.

High resolution structural studies will also be required to reveal the precise conformation of the Vps4 dodecamer. In this case, however, moderate resolution cryo-EM reconstruction of dodecameric Vps4 assemblies are available, and these structures reveal that the two hexameric rings adopt very different conformations³¹. The structure of the more constricted ring appears to be consistent with a homology model based upon the hexameric ring formed by the D1 AAA ATPase cassette of the related p97 ATPase⁴⁷. Although the resolution of the Vps4_{ΔMIT}

reconstruction is modest (~25Å), the homology model matches the reconstructed ring in overall shape and dimensions, and the six Vps4 β -domains appear to protrude from the exterior of reconstructed ring density as predicted by the homology model. Moreover, several mutations in the predicted hexamer interface block higher order Vps4 assembly, providing biochemical support for the homology model. It is noteworthy that some of these mutations also inhibit Vps4 dimerization (L151D), whereas others do not (R352A)²⁵, conversely some mutations block dimerization but not assembly (W388A). Thus, our data indicate that the dimer and hexameric ring interfaces utilize overlapping, but non-identical, surfaces.

Comparisons to Previous Vps4 Models

Our conclusions differ in important ways from two previous analyses of the unassembled Vps4 protein. Specifically, Xiao et al. argued that the low molecular weight Vps4 species formed in the absence of nucleotide was a monomer rather than a dimer²⁸. This conclusion was based in part on their observation that mutations in the symmetric crystallographic Vps4 dimer Interface 2 did not alter the Vps4 gel filtration mobility. Similarly, we also found that the interface mutation L407D did not alter the gel filtration mobility of Vps4_{ΔMIT} (Fig. 5, panel 2) However, Xiao et al. did not consider the alternative possibility that Vps4 might dimerize through a variant of the asymmetric Interface 6 (Figs. 4 and 5), whereas we found that mutations within this interface blocked Vps4 dimerization as analyzed both by gel filtration chromatography and analytical ultracentrifugation. Hence, our data demonstrate that Vps4 is primarily dimeric in its low molecular weight form.

Our analysis also differs from that of Hartmann et al., who argued that the Vps4 dimer consists of two molecules held together by the symmetrical crystallographic Interface 1²⁷. This conclusion was based upon their report that the Q216A mutation appeared to inhibit Vps4_{Δ MIT} dimerization, as analyzed by gel filtration chromatography. In our hands, however, Vps4_{Δ MIT} dimerization was not inhibited by the Q216A mutation or by other mutations within Interface 1. Furthermore, we do not believe that the data presented in Fig. 6 of Hartmann et al. constitute compelling evidence that the Q216A mutation inhibits Vps4_{Δ MIT} dimerization in solution.

These differences have important implications for models of higher order Vps4 assembly because Hartmann et al. used the Interface 1 dimer as a constraint in creating their model for the fully assembled Vps4 double ring²⁷. This constraint compelled a model in which the two Vps4 rings pack in a head-to-head orientation. However, our data indicate that the crystallographic dimer Interface 1 does not actually form in solution, as we find that Interface 1 mutants such as Q216A dimerize in solution (Fig. 5b). Furthermore, we find that the Q216A mutant also dodecamerizes normally, implying that that Gln216 does not make an essential contact in the fully assembled enzyme, as would be required by the head-to-head model (see Supplemental Fig. 3). We also find the head-to-head model unattractive because: 1) it differs from the ring orientations seen in all other well characterized Type II AAA ATPases, 2) it would require that the substrate-binding MIT domains emanate from the center of the double ringed structure (rather than from the end(s), as is seen for substrate binding domains of other AAA ATPases), and 3) it requires the ATPase activities of the two rings to work in opposition, which is difficult to envision if substrates are pumped up and through the rings.

Our cryo-EM model for the Vps4 dodecamer indicates instead that the two hexameric rings adopt very different conformations, the more constricted of which resembles the p97 D1 homology model. In this model, the Vps4 subunits interact through a surface that is similar, but not identical to the crystallographically defined Interface 6, which can explain why only a subset of mutations within Interface 6 inhibit Vps4 assembly. Vps4 residues that make up the intersubunit interaction surface in the p97 homology model are shown in detail in Supplemental Fig. 4. The conservation of many of these interface residues between the Vps4, p97 D1, and

spastin proteins, further supports the homology model. This model can also explain why Vps4 becomes an active ATPase upon assembly because hexamerization would position each ²⁸⁸RR²⁸⁹ dipeptide into the active site of an adjacent subunit, in close proximity with the γ -phosphate of the ATP. The use of such "arginine fingers" (sometimes also termed the second region of homology (SRH)) is a recurring feature in AAA ATPase active sites¹⁸, and serves an important catalytic role because the arginine(s) interacts with the γ -phosphate, making it a better leaving group. However, in the spiraling hexamers seen in crystal form 2 and related Vps4 crystal forms, the SRH residues of the adjacent subunit are located at least 12Å away from the γ -sulfate and therefore do not make interactions that would promote ATP hydrolysis. In our p97-based Vps4 hexamer homology model, these residues move much closer to the γ -sulfate, with the Arg288 residue brought to within 3.5Å of a γ -sulfate oxygen, where it could contact the terminal phosphate/sulfate and perform its expected role in catalysis.

A Functional Role for Vps4 Pore Loop 2

The central pores of AAA ATPases are typically defined by two adjacent loops that project into the central cavity. We have previously shown that hydrophobic Pore Loop 1 residues are required for VPS4A function, as judged by the ability of Pore Loop 1 mutants to dominantly inhibit HIV-1 release and infectivity 25 . In the current study, we used the same approach to show that Pore Loop 2 residues that help create the arginine collar are also required for human VPS4A function. Specifically, we found that overexpression of either VPS4A_{R236A} or VPS4A_{R246A} inhibited HIV-1 vector release and reduced viral titers more than 40-fold, showing that these mutant proteins were potent dominant negative inhibitors of HIV budding. As illustrated in Fig. 7, the two arginine residues appear to perform different functions, with VPS4A Arg236 (equivalent to yeast Vps4 Arg241) forming part of the positively charged collar that sits beneath the Pore 1 loop, and VPS4A Arg246 (equivalent to yeast Vps4 Arg251) lining the opposite side of the pore itself. Equivalent arginine residues are also important for the activity of other AAA ATPases, including the two closest relatives of VPS4; p97⁵¹, and spastin^{52; 53}. The functional importance of these conserved residues highlights the similarities between Vps4 and the other two related AAA ATPases, and suggests that conserved residues within the arginine collar may perform analogous mechanistic functions, possibly by binding the substrate and/or by assisting in protein denaturation 51 as polypeptide substrates are transferred through the central pore.

MATERIALS AND METHODS

Cloning

DNA encoding Vps4 from *S. cerevisiae* was amplified by PCR from yeast genomic DNA, and directionally cloned into the pET151 *E. coli* expression vector (Invitrogen). Constructs expressing mutant proteins were created by Quick-change mutagenesis (Stratagene) and verified by DNA sequencing.

Protein Expression and Purification

Vps4_{122–437, E233Q} and Vps4_{104–437, E233Q} were expressed in BL21 Codon+ *E. coli* cells (Stratagene) in ZY autoinduction media for 6 h at 37°C and then overnight at 21°C. Cells were harvested by centrifugation and lysed by resuspension in 10 mg/ml lysozyme in lysis buffer (10 mM imidazole, 300 mM NaCl, 50 mM Tris pH 7.4, 5% glycerol and protease inhibitors (PMSF, aprotinin, leupeptin, pepstatin), 45 min, 4°C), followed by sonication. The lysate was clarified by centrifugation (45 min, 35 000 × *g*) and the soluble Vps4 proteins were bound to a Ni²⁺ sepharose column (Amersham), washed with 10 column volumes of lysis buffer, and eluted in 75 mM imidazole (in lysis buffer). Fractions were assayed for Vps4 by SDS-PAGE, pooled, dialyzed into 100 mM NaCl, 25 mM Tris pH 7.4, and 1 mM DTT, and purified by anion-exchange chromatography (Q-sepharose, Invitrogen, 100–500 mM NaCl gradient in 25

mM Tris pH 7.4, and 1 mM DTT). Vps4 fractions were pooled, and the $(His)_6$ affinity tag was removed by incubation with TEV protease (~1 mg/100mg protein, 20 h at 4°C), followed by dialysis into 20 mM Tris pH 7.4, 100 mM NaCl, and removal of unprocessed protein by Ni²⁺ sepharose chromatography. The cleaved protein was collected in the flow through, concentrated, and purified to homogeneity by gel filtration chromatography (SD200, Amersham, in 50 mM NaCl, 25 mM Tris pH 7.4, 1mM DTT). Yields were typically 25 mg/l culture.

Gel Filtration Chromatography

The gel filtration chromatography experiments shown in Fig. 5, Fig. 6, and Supplemental Fig. 3 were performed on 150 μ M Vps4 proteins using a Superdex 200 column in 100 mM NaCl, 25 mM Tris–HCl (pH 7.5), and 1 mM DTT in the presence or absence of ATP. Both columns were calibrated using molecular weight standards (Biorad).

Analytical Ultracentrifugation

Equilibrium sedimentation experiments on full length Vps4_{E233Q}, Vps4_{Δ MIT}, E233Q and Vps4_{Δ MIT}, E233Q with L151D and Q216A interface mutations, were performed on an XL-A analytical ultracentrifuge (Beckman Coulter). Protein concentrations ranged from 5–30 µm (110-µl) and the corresponding buffer blanks (120 µl; 50 mM NaCl, 25 mM Tris pH 7.5) were run in parallel in sample cells fitted with six-channel equilibrium centerpieces. All data were collected at 4°C and at two rotor speeds for each protein: Vps4 at 12,000 and 16,000 rpm, for Vps4_{Δ MIT} at 14,000 and 18,000 rpm, and for Vps4_{Δ MIT} interface mutants L151D and Q216A at 16,000 and 20,000 rpm. The attainment of equilibrium after 24 h was confirmed by comparing repetitive absorption scans. Theoretical molecular weights, partial specific volumes (v_{bar}), and molar extinction coefficients at 280 nm of the Vps4 variants were calculated using the program SEDNTERP (Version 1.09, available on the World Wide Web)⁵⁸. All data were analyzed using the program Heteroanalysis⁵⁹.

Crystallization

Crystals were grown by sitting drop vapor diffusion at 21°C. Vps4 $_{104-437, E233Q}$ crystallized in drops comprising 2 µl of 13 mg/ml protein, 50 mM NaCl, 25 mM Tris pH 7.4, 1 mM DTT, 4 mM MgCl₂, and 2 mM ATPγS, mixed with 1 µl of reservoir solution (48% ethylene glycol (vol/vol), Bis-Tris pH 5.0). Crystals were cryo-cooled by plunging into liquid nitrogen directly from the crystallization solution. Vps4_{122-437, E233Q} was crystallized in drops comprising 2 µl of 12 mg/ml protein, 100 mM NaCl, 25 mM Tris pH 7.4, 1 mM DTT, 4 mM MgCl₂, and 1 mM ATPγS, mixed with 2 µl reservoir solution (1.6 M ammonium sulfate, 0.1 M Bis-Tris pH 5.5). Crystals were briefly transferred to cryoprotectant (mother liquor made up with 20% glycerol), suspended in a nylon loop, and cryo-cooled by plunging into liquid nitrogen.

Structure Determination and Refinement

Diffraction data from Vps4_{104–437} and Vps4_{122–437} crystals were collected at Beamline X25 of the National Synchrotron Light Source, Brookhaven National Laboratory, and scaled and integrated by using the HKL2000 suite⁶⁰. Crystal structures were determined by molecular replacement using PHASER⁶¹. A polyalanine/homology search based on the human VPS4B structure (pdb code 1XWI)²⁵ was used to determine Vps4_{122–437}, E233Q, and this refined structure was used to determine the structure of Vps4_{104–437}, E233Q. Model building was performed with COOT⁶². Structures were refined using REFMAC5⁶³ with TLS refinement using TLSMD⁶⁴ and TLSANL⁶⁵ in the CCP4 suite⁶⁶. Structures were analyzed with PROCHECK⁶⁷ and figures were generated using PYMOL⁶⁸.

Infectivity Assays

293T cells were seeded in 6-well plates at 8×10^5 cells per well and co-transfected with a pEGFP-VPS4A expression vector and an HIV-1 vector system (packaging and transfer plasmids kindly provided by D. Trono, and envelope plasmid kindly provided by J. Burns). Briefly, 12 µl Fugene 6 Transfection Reagent (Roche Diagnostics) was combined with 85.14 µl Optimem (no additives) and incubated at room temperature for 5 minutes. 1 µg of pCMVDR8.2, 1 µg of pWPTS-nlsLacZ, 0.36 µg phCMV-VSVG, and 0.5 µg of wild type or mutant pEGFP-VPS4A was combined with the transfection mixture (final volume of 100 µl) and incubated at room temperature for 20 minutes. Virions were harvested ~48 h post-transfection and used to transduce the reporter HeLa-M cell line as described previously³⁸ or used in Western blotting experiments to analyze virion release.

Virion Release Assays

Virion-containing supernatants and proteins from transfected cells were prepared for Western blotting as described⁶⁹; ⁷⁰. Primary antibodies used were: rabbit anti-HIV-1 MA (1:1000, D. Trono), rabbit anti-HIV-1 CA UT415 (1:1000, Covance), rabbit anti-VPS4A UT289 (1:000, Covance), and mouse anti- γ -Tubulin GTU-88 (1:5000, Abcam). Secondary antibodies used were: goat anti-rabbit ALEXA680nm (1:20,000, Molecular Probes) and donkey anti-mouse IR800nm (1:10,000, Rockland). Proteins were detected by imaging fluorescent secondary antibodies on an Odyssey scanner (LiCor Inc.).

Protein Data Bank Accession Numbers

Atomic coordinates and structure factors for yeast Vps4 $_{\Delta MIT}$ have been deposited in the RCSB Protein Data Bank with accession number 3EIE for the SO₄-bound form and 3EIH for the ATP γ S-bound form.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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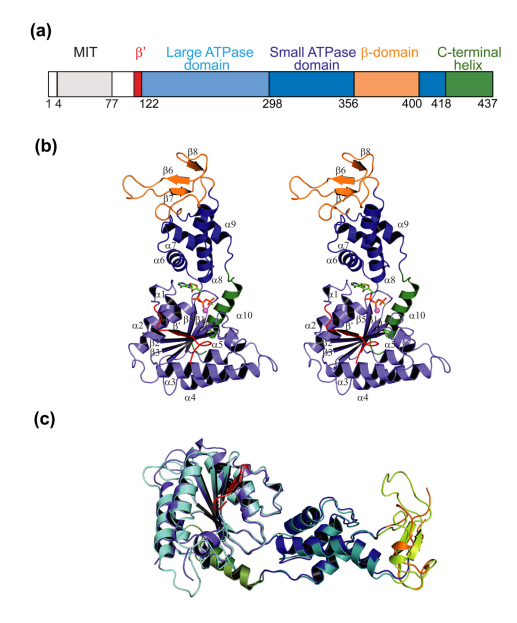


Figure 1.

Structure of yeast Vps4.

(a) Domain organization of the human and yeast Vps4 proteins. Numbering scheme corresponds to the yeast Vps4 protein.

(b) Stereoview of the Vps4_{Δ MIT} structure. This structure corresponds to molecule A in the ATP γ S bound state in crystal form 2. Domains are color coded as in part (a). Note that the small helix spanning residues 199–206 is not numbered in order to maintain consistency with previous conventions.

(c) Superposition of yeast Vps4_{Δ MIT} (in crystal form 1) and human VPS4B_{Δ MIT} (lighter shades) in their sulfate-bound states. The two proteins overlay with an RMSD of 1.2 Å over 225 Ca positions within the AAA ATPase cassette.

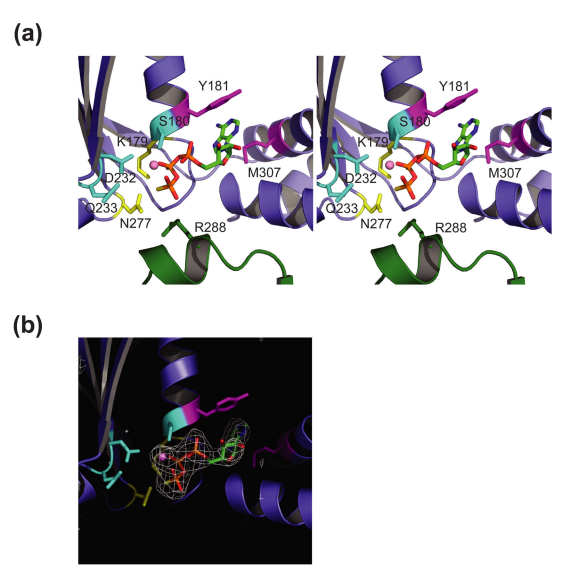


Figure 2.

Vps4 nucleotide binding pockets.

(a) Stereoview of the Vps4-ATP γ S nucleotide binding site of molecule A in crystal form 2. Active site Vps4 residues are color coded according to their functional roles in Mg²⁺ coordination/ATP hydrolysis (cyan, S180, D232 and E/Q233), adenine ring stacking (magenta, Y181 and M307) and phosphate sensing (yellow, K179 and N277) with the "arginine finger" residue R288 from an adjacent molecule in the modeled hexamer shown in green. Note that the E233Q mutant was used here and throughout to allow ATP/ATP γ S binding while inhibiting hydrolysis.

(b) Electron density for the ATP γ S nucleotides in molecule A of Vps4_{Δ MIT} crystal form 2. The densities show (F₀ - F_C) omit maps contoured at 2.5 σ .

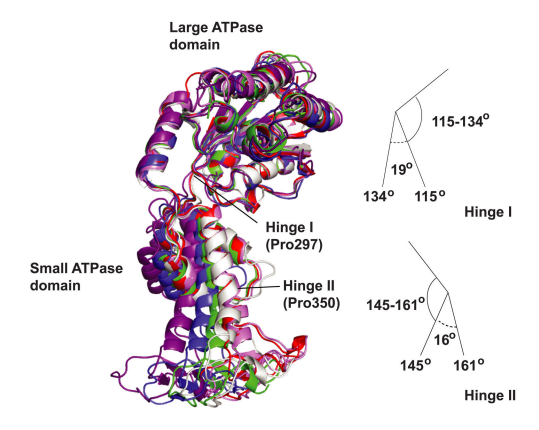


Figure 3.

Interdomain flexibility in different Vps4 $_{\Delta MIT}$ crystal structures.

Superposition of six independent structures of Vps4_{ΔMIT} in complex with: no ligand (2RKO, white), sulfate (crystal form 1, green), phosphate (2QPA, molecule B, blue), ADP (2QPA, molecule A, red), ATPγS (crystal form 2, molecule A, pink), and ATPγS (crystal form 2, molecule C, purple). Superposition on the large ATPase domains reveals that the small ATPase domain can rotate by up to 19° about a hinge angle located in the linker between the two domains and centered at residue Pro297. A second hinge within the small ATPase domain is also evident in the middle of the extended helix α 8, centered about residue Pro350. These two hinge angles were defined by first finding the centers of masses of the large ATPase domain (residues 122–298 and 418–433), the core of the small domain (300–350; 402–414), and the β domain (358–399) using the program 6d_moleman2. Hinge angle I was then calculated as the angle between the large and small domains, with Pro297 C α at the apex, and hinge angle II was calculated as the angle between the small domain and the β domain, with Pro350 C α at the apex.

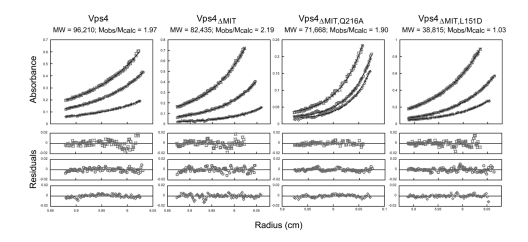


Figure 4.

Vps4 proteins dimerize in solution

Representative equilibrium sedimentation profiles for full length Vps4 (panel 1), Vps4_{Δ MIT} (panel 2), Vps4_{Δ MIT}, Q216A (panel 3, Interface 1 mutant), and Vps4_{Δ MIT}, L151D (panel 4, Interface 2 mutant). Sedimentation data are plotted as absorbance versus the distance from the center of the axis of rotation (radius). To simplify the plot, the radius was normalized so that the data from all three sectors (i.e. three different concentrations) overlap. In each case, data from three protein concentrations and one speed are displayed (open symbols) along with the best single ideal species fit (solid lines). The best fits for each protein were derived from global fits to data collected at three concentrations and two speeds. The residuals (differences between the raw absorbance data and the fit are shown below each panel.



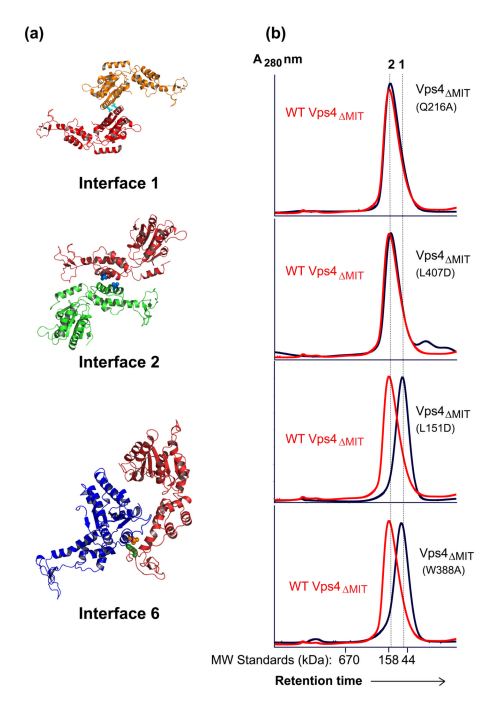


Figure 5.

Mutational analyses of crystallographic Vps4 dimer interfaces.

(a) Crystallographic Vps4 dimer interfaces. Interface 1 is a symmetric interface between two large ATPase domains (residue Q216 is shown in cyan), Interface 2 is a symmetric interface between two small ATPase domains (residue L407 is shown in blue), and Interface 6 is an asymmetric interface between the large and small ATPase domains (residues L151 and W388 are shown in orange and green, respectively).

(b) Gel filtration chromatograms of Vps4_{Δ MIT} proteins with the following mutations: Q216A (Interface 1), L407D (Interface 2), L151D (Interface 6), and W388A (Interface 6). Vps4_{Δ MIT} proteins used here and elsewhere contained the E233Q mutation, which allowed

ATP binding but inhibited hydrolysis. For reference, the elution profile of the "wild type" Vps4_{Δ MIT, E233Q} protein is shown in red in each panel, elution positions for monomeric (1) and dimeric (2) proteins are shown as dotted vertical lines, and the elution positions of molecular weight standards are shown below the chromatograms. Vps4 protein concentrations were 150 μ M in all cases. Note that at low micromolar concentrations, the dimeric proteins exhibited concentration-dependent mobilities (not shown), indicating that appreciable concentrations of monomers could accumulate under these low protein and non-equilibrium conditions.

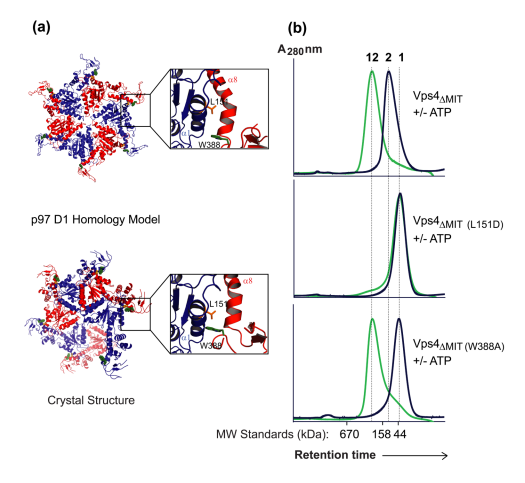


Figure 6.

Mutagenesis test of the Vps4 $_{\Delta MIT}$ hexamer homology model.

(a) The "p97 D1 Homology" model for a Vps4_{Δ MIT} hexameric ring within the dodecameric enzyme (upper) is shown together with the arrangement of Vps4 along the crystallographic six-fold screw axis (lower). The p97 D1 Homology Model was created by superimposing Vps4_{Δ MIT} subunits onto the crystal structure of the hexameric ring of the homologous p97 D1 protein⁴⁴. The Trp388 (green) and Leu151 (orange) residues are shown explicitly in homology model and crystal structure to illustrate how the Leu151 residue contributes to the interface in both models whereas the Trp388 residue contributes to the interface in the crystal structure but not in the p97 D1 Homology Model (see insets).

(b) Gel filtration chromatograms of wild type and mutant Vps4_{Δ MIT} proteins in the absence (black) or presence (green) of ATP. For reference, elution positions for monomeric (1), dimeric (2), and dodecameric (12) proteins are shown as dotted vertical lines, and the elution positions of molecular weight standards are shown below the chromatograms. Note that upon addition of ATP, the wild type Vps4_{Δ MIT} protein converts from a dimer to a dodecamer,

Vps4 $_{\Delta MIT, L151}$ remains a monomer, and Vps4 $_{\Delta MIT, W388A}$ converts from a monomer to a dodecamer.

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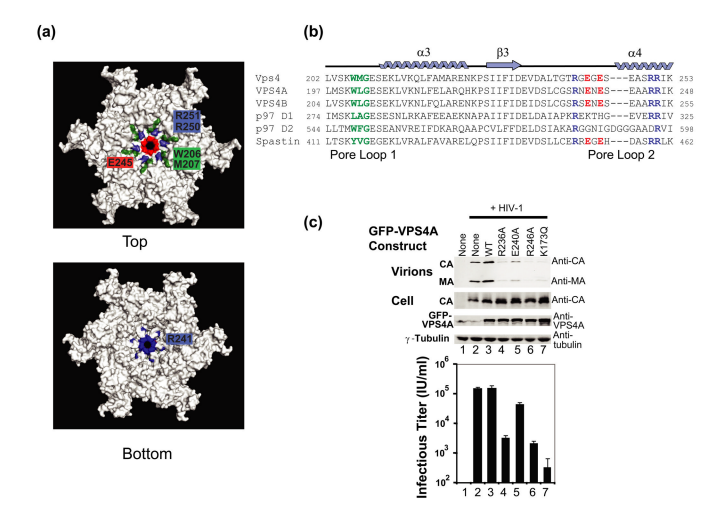


Figure 7.

VPS4A Pore Loop 2 residues are required for HIV budding.

(a) Two views of the p97 D1 Homology model for a hexameric ring of yeast Vps4_{Δ MIT} in the ATP γ S-bound state. Pore loop 1 residues ($_{206}$ WMG $_{208}$) are shown in green, and a subset of the charged Pore Loop 2 residues are shown in blue (basic residues) or red (acidic residues). (b) Sequence alignments highlighting Pore Loop 1 (green) and Pore Loop 2 (blue, basic, and red, acidic) residues from human and yeast VPS4 proteins. Equivalent residues from the related p97 and Spastin AAA ATPase cassettes are highlighted following the same color scheme (except for non-conservative changes).

(c) Vps4 Pore Loop 2 mutations dominantly inhibit HIV-1 release and infectivity. 293T cells were co-transfected with an empty vector control (lane 1) or with a proviral HIV-1 expression vector (lanes 2–7) together with the designated GFP-VPS4A expression constructs. The top two western blots show the levels of virion-associated CA and MA proteins released into the supernatant (Panel 1, Virion) or expressed in the cells (Panel 2, Cell). The bottom two western blots show cellular levels of GFP-VPS4A and of an γ -Tubulin control. The bottom panel shows vector titers in the supernatants under the different conditions (infectious units/ml). Note that expression of the different Pore Loop 2 mutant GFP-VPS4A constructs (lanes 3–6) or of the VPS4A ATPase mutant (lane 7, positive control) dominantly inhibited virion release and titers. Infectivity values are the mean of measurements from quadruplicate repeats and error bars indicate standard deviations.

Data collection and refinement statistics

	Vps4 _{122-437, E233Q} Crystal form 1	Vps4 _{104-437, E233Q} Crystal form 2
Data collection		
Space group	P6 ₅ 22	P2 ₁ 2 ₁ 2 ₁
Cell parameters (Å)	a = 110.8	a = 77.0
	b = 110.8	b = 119.8
	c = 169.1	c = 156.8
Wavelength (Å)	1.00	1.00
Resolution (Å)	30-2.7	30-3.25
Outer Shell (Å)	2.77-2.7	3.37-3.25
Number of Reflections		
Total observed	117,584	121,837
Unique	17,628	23,550
Completeness (%)	99.8 (97.5)	99.8 (100)
$\mathbf{R}_{\mathrm{sym}}\left(\% ight)^{a}$	11.1 (52.6)	12.0 (62.9)
Mean I/o (I)	9.8 (2.3)	8.0 (1.9)
Refinement statistics		
R_{cryst}/R_{free} (%) ^{b,c}	24.1/28.7	21.6/28.9
RMSD from ideal geometry d		
Bond lengths (Å)	0.012	0.017
Bond angles (°)	1.489	1.970
Average B-factor (Å ²)	51.5	86.6
Phi/psi angles (non Gly/Pro)		
Most favorable region (%)	86.7	77.4
Additional allowed region (%)	10.7	20.5
Generous allowed region (%)	2.7	1.6
Disallowed region (%)	0.4	0.7

Values in parenthesis are for the highest resolution shell.

 $^{a}\mathrm{R}_{sym}$ = (|(ΣI – <I>)|)/(ΣI), where <I> is the average intensity of multiple measurements.

 ${}^{b}\mathbf{R}_{cryst} = \boldsymbol{\Sigma}_{hkl} ||\mathbf{F}_{obs}(\mathbf{hkl})|| - \mathbf{F}_{calc}(\mathbf{hkl})||/\boldsymbol{\Sigma}_{hkl}|\mathbf{F}_{obs}(\mathbf{hkl})|$

 C R_{free} = the crossvalidation R factor for 5% of reflections against which the model was not refined

 d Geometry was analyzed in PROCHECK⁶⁷.

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NIH-PA Author Manuscript	Table 2	The intermolecular lattice contacts in all known Vps4 _{ΔMIT} structures
NIH-PA Autho		The intermolecular

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Structure	PDB	Mol/AU	Ligand bound	Hinge angle I	Hinge angle II	Total buried area per monomer [Å 2]
Vps4 (126–433)	2RKO	-	apo	116	157	Interface 1 – 637 Interface 2 – 290 Interface 3 – 156 Interface 6 – 722
Vps4 (129–433)	2QP9	1	SO_4	116	161	Interface 1 – 740 Interface 2 – 431 Interface 3 – 284 Interface 6 - 740
Vps4 (122-433)	This work	-	SO_4	118	156	Interface 1 – 477 Interface 2 – 333 Interface 3 – 409 Interface 6 – 767
Vps4 Molecule A (119–435) Molecule B (119–435) Molecule C (123–432)	2QPA	Ś	ADP PO4 PO4	116 119 124	147 160 153	Interface 2(AB) - 457 Interface 4(BC) - 216 Interface 5(AA) - 280 Interface 5(AB) - 240 Interface 6(AB) - 701 Interface 6(AC) - 673 Interface 6(BC) - 721
Vps4 Molecule A (119–437) Molecule B (120–435) Molecule C (123–437)	This work	κ	ATP/S ATP/S Ethylene glycol- Mg ²⁴	115 118 134	145 155 150	Interface 2(AB) - 676 Interface 4(BC) - 426 Interface 5(AA) - 480 Interface 5(BC) - 451 Interface 6(AB) - 895 Interface 6(AC) - 772 Interface 6(BC) - 720