1	Structural basis of CHMP2A-CHMP3 ESCRT-III polymer assembly and membrane
2	cleavage
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28 Abstract

The endosomal sorting complex required for transport (ESCRT) is a highly conserved protein 29 machinery that drives a divers set of physiological and pathological membrane remodeling 30 31 processes. However, the structural basis of ESCRT-III polymers stabilizing, constricting and 32 cleaving negatively curved membranes is yet unknown. Here we present cryo electron microscopy 33 structures of membrane-coated CHMP2A-CHMP3 filaments of two different diameters at 3.3 and 34 3.6 Å resolution. The structures reveal helical filaments assembled by CHMP2A-CHMP3 35 heterodimers in the open ESCRT-III conformation, which generates a partially positive charged membrane interaction surface, positions short N-terminal motifs for membrane interaction and the 36 C-terminal VPS4 target sequence towards the tube interior. Inter-filament interactions are 37 electrostatic, which facilitate filament sliding upon VPS4-mediated polymer remodeling. 38 Fluorescence microscopy as well as high speed atomic force microscopy imaging corroborate that 39 CHMP2A-CHMP3 polymers and VPS4 can constrict and cleave narrow membrane tubes, thus 40 41 acting as a minimal membrane fission machinery.

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44 Introduction

The endosomal sorting complex required for transport (ESCRT) machinery catalyzes many divergent membrane remodeling processes including the formation of multivesicular endosomes, cytokinesis, nuclear envelope reformation, membrane repair, autophagy, exosome biogenesis, neuronal pruning, dendritic spine maintenance, enveloped virus budding, release of peroxisomes and of recycling endosomes (Allison et al., 2013; Henne et al., 2013; Loncle et al., 2015; Mast et al., 2018; Olmos and Carlton, 2016; Sadoul et al., 2018; Scourfield and Martin-Serrano, 2017; Votteler and Sundquist, 2013; Zhen et al., 2021).

52 Common to all ESCRT-catalyzed processes in eukaryotes, archaea and bacteria is the recruitment of ESCRT-III proteins that polymerize to generate and/or to stabilize membranes with 53 54 either flat, negatively or positively curved geometries (Bertin et al., 2020; Caillat et al., 2019; Gupta et al., 2021; Junglas et al., 2021; Liu et al., 2021; McCullough et al., 2018; Moser von Filseck et 55 al., 2020; Pfitzner et al., 2021). The principal function of the polymers is to induce membrane 56 constriction via outside-in fission of tubular structures with ESCRT-III protein coats on the outside 57 of a membrane tube or inside-out fission with ESCRT-III polymers assembled within membrane 58 59 neck/tube structures formed during vesicle and virus budding or at the cytokinetic midbody (Caillat et al., 2019; Harker-Kirschneck et al., 2022; McCullough et al., 2018; Nguyen et al., 2020; Pfitzner 60 et al., 2021; Remec Pavlin and Hurley, 2020). 61

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Humans express eight ESCRT-III proteins that can comprise several isoforms per 62 member, corresponding to seven homologues in S. cerevisiae (in parentheses) named 63 CHMP1A/B (Did2), CHMP2A/B (Vps2), CHMP3 (Vps24), CHMP4A/B/C (Snf7), CHMP5 (Vps60), 64 65 CHMP6 (Vps20), CHMP7 and CHMP8/IST1 (Ist1) (McCullough et al., 2018). S. cerevisiae 66 assembles two ESCRT-III subcomplexes, Vps20/Snf7 that in turn recruits Vps24/Vps2 (Babst et 67 al., 2002) consistent with CHMP4 recruiting CHMP3 and CHMP2A (Morita et al., 2011). Notably, 68 Vps24 (CHMP3) and Vps2 (CHMP2) have been suggested to block Snf7 (CHMP4) polymerization 69 and cap ESCRT-III assembly prior to recycling (Saksena et al., 2009; Teis et al., 2008). Consistent with the idea of a core ESCRT-III, HIV-1 budding can be catalyzed with a minimal set of one 70 CHMP4 and one CHMP2 isoform (Morita et al., 2011). Although CHMP3 is not strictly required, 71 72 truncated versions thereof exert a potent dominant negative effect on HIV-1 budding (Zamborlini et al., 2006) and CHMP3 synergizes HIV-1 budding efficiency with CHMP2A but not with CHMP2B 73 (Effantin et al., 2013). Thus ESCRT-III CHMP4, CHMP2 and CHMP3 constitute a minimal 74 machinery that together with VPS4 catalyzes membrane fission from within membrane necks as 75 suggested by in vitro reconstitution (Schoneberg et al., 2018). 76

77 ESCRT-III proteins adopt a closed conformation in the cytosol (Bajorek et al., 2009; Muziol et al., 2006; Xiao et al., 2009). Membrane recruitment via ESCRT-I, ESCRT-II or Alix/Bro1 (Im et 78 79 al., 2009; McCullough et al., 2008; Pineda-Molina et al., 2006; Tang et al., 2016) is thought to 80 induce ESCRT-III activation, which entails opening of the closed conformation (Lata et al., 2008a; Shim et al., 2007; Zamborlini et al., 2006) to an extended open polymerization-competent 81 conformation as first shown for CHMP1B (McCullough et al., 2015). The CHMP1B polymer 82 83 stabilizes positively curved membranes and can co-polymerize with IST1 in the closed conformation thereby forming an outer layer on top of the inner open conformation CHMP1B layer 84 (McCullough et al., 2015), whose interplay leads to membrane tube thinning and cleavage (Cada 85 et al., 2022; Nguyen et al., 2020). 86

CHMP4 homologues, Snf7 and shrub, adopt similar open conformations within crystalline 87 88 polymers (McMillan et al., 2016; Tang et al., 2015). Notably in vitro CHMP4 polymers interact with flat membranes, (Chiaruttini et al., 2015; Mierzwa et al., 2017; Pires et al., 2009) and stabilize 89 positively curved membranes in the presence of CHMP2A (Vps2) and CHMP3 (Vps24) (Bertin et 90 al., 2020; Moser von Filseck et al., 2020). Furthermore, CHMP4 was proposed to interact with 91 negatively curved membranes (Lee et al., 2015) and CHMP4 spirals have been imaged within 92 membrane tubes in vivo (Cashikar et al., 2014; Hanson et al., 2008) leading to the model of 93 94 ESCRT-III spiral springs assembling on flat membranes driving membrane deformation (Chiaruttini et al., 2015). 95

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The conservation of the structural principle of the open ESCRT-III conformation is further underlined by the structures of plastid and bacterial membrane repair proteins Vipp1 and PspA, which both stabilize positively curved membranes and corroborate the conservation of the ESCRT-III machinery throughout all kingdoms of life (Gupta et al., 2021; Junglas et al., 2021; Liu et al., 2021).

101 While these structures demonstrate filament formation to stabilize positively curved 102 membranes only low resolution models of filaments stabilizing negatively curved membranes have 103 yet been imaged revealing single and multi-stranded polymers in vitro (Bajorek et al., 2009; Bertin et al., 2020; Chiaruttini et al., 2015; Dobro et al., 2013; Henne et al., 2012; Lata et al., 2008b; 104 Mierzwa et al., 2017; Moriscot et al., 2011; Moser von Filseck et al., 2020; Pires et al., 2009) and 105 106 in vivo (Bodon et al., 2011; Cashikar et al., 2014; Goliand et al., 2018; Guizetti et al., 2011; Hanson et al., 2008; Mierzwa et al., 2017; Sherman et al., 2016). Consistent with their central role in 107 membrane fission catalyzed from within membrane necks, ESCRT-III CHMP2A and CHMP3 form 108 helical tubular structures with defined diameters in vitro, which have been suggested to stabilize 109 negative membrane curvature (Effantin et al., 2013; Lata et al., 2008b). VPS4 constricts these 110 111 filaments producing dome-like end caps prior to complete polymer disassembly in vitro (Maity et al., 2019) in agreement with permanent ESCRT-III turn-over in vivo (Adell et al., 2014; Adell et al., 112 2017; Mierzwa et al., 2017). Notably, S. cerevisiae Vps2 and Vps24 form similar helical tubes that, 113 114 however, seem to require Snf7 for polymerization (Henne et al., 2012).

Because most ESCRT-catalyzed processes act on negatively curved membranes to catalyze inside-out membrane fission, we set out to determine the structural basis of ESCRT-III stabilizing negatively curved membranes. We reconstituted CHMP2A-CHMP3 polymers within membrane tubes, solved their structure by cryo electron microscopy and show that VPS4 can indeed constrict CHMP2A-CHMP3 membrane tubes to the point of fission, corroborating that CHMP2A and CHMP3 form a minimal membrane fission complex powered by VPS4 and ATP.

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122 **Results**

123 Structure of the CHMP2A-CHMP3 polymer assembled within membrane tubes

124 CHMP3 full length (residues 1-222) and C-terminally truncated CHMP2A (residues 1-161) were 125 assembled into helical tubular structures as described (Lata et al., 2008b). After removal of the N-126 terminal tags, the tubular structures were coated with lipid bilayer, which tightly associated with 127 the protein layer as shown by cryo electron microscopy (cryo-EM) (**Figure S1A**). 2D classification 128 of the manually picked tube-like structures generated a dataset composed of 5 different diameters 129 ranging from 380 Å to 490 Å, with the 410 Å (51.4%) and 430 Å (31.2%) diameters representing

the most populated classes (Figure S1B). The power spectra of the segments of the class 130 131 averages of the two diameters (Figures S1C and D) were then employed to explore possible helical symmetries and combined with helical real-space reconstruction (He and Scheres, 2017) 132 to validate the symmetry parameters. This revealed that the 410 Å and 430 Å diameter tubes are 133 134 formed by elementary helices composed of respectively 6.3 and 6.6 units per turn with a small pitch of 9 Å for the 410 Å diameter and 18 Å for the 430 Å diameter. The latter displays an 135 136 additional C2 symmetry around the helical axis, explaining the doubling of the pitch (Figures S1C 137 and D). Although, the asymmetric units along the elementary helix are not biochemically connected, they translate into filaments with extended interaction surfaces between subsequent 138 asymmetric units. The symmetry parameters of the filaments are relatively similar for both 139 diameters (Rise/Twist of 8.227 Å /16.877° and 8.641 Å /17.701° for the 410 Å and 430 Å 140 diameters, respectively), and therefore likely represent the preferred polymerization mode of the 141 repeating unit, the CHMP2A-CHMP3 heterodimers (Figures S1C and D). 142

Six filaments (2*3 in the case of the C2 symmetric helix) form left-handed six-start helices with helical pitches of ~175 Å for both diameters (**Figures 1A and B**). The 430 Å filament is composed of 21.33 units per turn (**Figures 1A and B**) and the 410 Å filament contains 20.33 units per turn (**Figures 1C and D**), indicating a helical repeat after three turns and similar inter-filament interactions along the helical axis. Comparison of the 430 and 410 Å demonstrates that removal of one heterodimer reduces the tube diameter by 20 Å.

The three-dimensional (3D) helical reconstruction shows an overall resolution of 3.6 Å for the 410 Å and 3.3 Å for the 430 Å diameter (**Figure 1; Figure S2 and Figure S3A**) with local resolutions ranging from 3.3 Å to 4.6 Å for the 430 Å diameter and 3.6 to 5.2 Å for the 410 Å diameter (**Figure S3B and C**). The map of the 430 Å diameter was employed (**Figure 1E**) to build the atomic model of the repeating unit of the filament, formed by the CHMP2A-CHMP3 heterodimer revealing both protomers in the open ESCRT-III conformation (**Figure 1F and Figure S3D, Table S1**).

Comparison of the closed (Bajorek et al., 2009; Muziol et al., 2006) and open CHMP3 156 conformations showed the conformational transitions upon CHMP3 activation, which involves 157 extension of the helical hairpin (residues P12-A101) that is identical in both conformations (r.m.s.d. 158 of 1.082 Å) by a linker and helix 3 (L117). The following short connection forms an elbow and 159 translates helix 4 by ~10 Å positioning helix 4 in a 140° angle with respect to the hairpin axis. Helix 160 161 4 is composed of the closed conformation helix 4 and most of the disordered linker connecting to helix 5 via a 90° kink at positions M151 to D152 (Figure 2A). The remaining CHMP3 residues 170 162 to 220 are flexible and disordered in the structure. Both CHMP3 and CHMP2A open conformations 163

are similar as superposition of their Cα atoms revealed an r.m.s.d. of 0.934 Å (Figure S4A),
 suggesting that CHMP2A can fold into the same closed conformation structure as CHMP3.

The repeating unit of the filament is the CHMP2A-CHMP3 heterodimer formed by parallel 166 167 interaction of their hairpins with the CHMP2A hairpin tip shifted by six helical turns with respect to 168 CHMP3 (Figure 2B). The heterodimer interaction covers 2026 Å² of CHMP2A and 1997 Å² of 169 CHMP3 surfaces involving 55 and 51 interface residues, respectively. The structure of the 170 heterodimer was further confirmed by mutagenesis. Introducing pairs of cysteine demonstrated 171 that CHMP2A D57C together with CHMP3 S75C and CHMP2A N18C together with CHMP3 V110C (Figure S5A) assembled into disulfide-linked heterodimers upon polymerization 172 into tube-like structures as shown by SDS-PAGE analyses and negative staining EM (Figure S5B 173 174 and C). Furthermore, mutagenesis of CHMP2A-CHMP3 interface residues (Figure S5D) prevented polymerization as expected (Figure S5E). The principle of the heterodimer hairpin 175 stacking is employed to assemble the filament, which is further stabilized by lateral interactions of 176 CHMP3i elbow helix 4 with CHMP2Ai, CHMP3i⁺¹ and CHMP2Ai⁺¹. In addition, CHMP3i helix 5 177 interacts with the tip of the CHMP3i⁺² hairpin (Figure 2B), as observed in the closed conformation 178 179 (Figure 2A) (Bajorek et al., 2009; Muziol et al., 2006). Similar to CHMP3, CHMP2A helix 4 exerts the same domain exchange interactions (Figure 2B). Cα superposition of the open CHMP3 180 conformation revealed the closest match with the S. cerevisiae Snf7 protomer (Figure S4B) and 181 considerable differences with CHMP1B, Vipp1, PspA and Vps24 (Gupta et al., 2021; Huber et al., 182 2020; Junglas et al., 2021; Liu et al., 2021; McCullough et al., 2015) (Figure S4C-F). Notably, 183 different hairpin interactions and orientations of the helical arms upon polymerization determine 184 185 the filament geometry that leads to positively curved membrane interaction by CHMP1B, Vipp1 and PspA (Gupta et al., 2021; Junglas et al., 2021; Liu et al., 2021; McCullough et al., 2015) 186 187 underlining the extensive structural plasticity of ESCRT-III proteins.

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189 CHMP2A-CHMP3 polymer interaction with membrane

190 The CHMP2A-CHMP3 polymer is tightly associated with the lipid bilayer (Figure S1E) and both CHMP2A and CHMP3 expose the same regions to the membrane. The polymerization mode 191 positions the N-terminal regions of both CHMP2A and CHMP3 at the membrane interface. 192 193 Although CHMP3 residues 1-10 and CHMP2A residues 1-7 are disordered, they are both oriented by conserved prolines towards the lipid bilayer (Figure 3A) consistent with previous suggestions 194 195 that short amphipathic N-terminal helices insert into the bilayer (Bodon et al., 2011; Buchkovich et 196 al., 2013). The main membrane interaction surfaces locate to the elbow formed by helices 3 and 197 4 (residues K104 to R131) exposing six basic residues of CHMP2A (K104, K108, R115, K118,

K124, R131) and five basic residues of CHMP3 (K106, K112, K119, K132, K136) prone to interact 198 199 with negative charges of the membrane (Figure 3B). The electrostatic potential map shows in 200 addition to the stretch of basic surfaces some negative and non-charged regions of the outer 201 polymer surface (Figure 3C). Most of the basic residues are conserved in S. cerevisiae Vps2 and Vps24 (Figure S6A). Notably, alanine mutagenesis of some CHMP3 basic residues within the 202 203 membrane interaction surface did not interfere with CHMP2A-CHMP3 polymerization in vitro 204 (Figure S7A) nor did they affect the dominant negative effect of C-terminally truncated CHMP3 205 on VLP release (Figure S7B), indicating that membrane binding is complex and not only electrostatic, consistent with plasma membrane localization of the CHMP3 mutant (Figure S7C). 206

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208 Inter-filament interactions

Conserved basic helix 1 residues of CHMP2A and CHMP3 (Figure S6A) are at the filament 209 interface opposed by a stretch of conserved acidic residues within helices 4 and 5 (Figure S6A) 210 of neighboring filaments (Figure 3D), which indicate electrostatic inter-filament interactions 211 (Figure 3C). Mutation of the helix 1 basic cluster within either CHMP2A or CHMP3 prevented 212 213 polymer formation in vitro (Figure S7D), indicating that the basic charge of helix 1 is important for filament polymerization, which is in line with mutagenesis of a similar cluster of basic residues 214 215 within helix 1 of CHMP3 abolishing its dominant negative effect on HIV-1 budding (Muziol et al., 2006). To further test the electrostatic inter-filament interactions, we exposed the helical tubular 216 CHMP2A-CHMP3 polymers to high ionic strength. This led to the partial unwinding of the filaments 217 producing single and multi-stranded filaments (Figure S8A-E) in agreement with the presence of 218 219 single and multi-start helices upon CHMP2A-CHMP3 polymerization in vitro (Effantin et al., 2013). We suggest that these electrostatic interactions between filaments enable filament sliding upon 220 221 VPS4-catalyzed remodeling. The acidic cluster in helices 4 and 5 is conserved in CHMP4A, B, C, 222 CHMP5 and CHMP6 (Figure S6B) indicating potential similar involvement in inter-filament interaction for the formation of mixed filaments. Notably, the acidic cluster is not conserved in 223 224 CHMP1A and B, which stabilizes positively curved membranes via basic charges present on the inside of the protein tube-like polymer (McCullough et al., 2015). We therefore suggest that the 225 acidic cluster is a hallmark of ESCRT-III stabilizing negatively curved membrane structures. 226

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228 VPS4 remodels and cleaves CHMP2A-CHMP3 membrane tubes

229 We next tested whether VPS4B can remodel the CHMP2A-CHMP3 membrane coated tubes as

we have shown before for CHMP2A-CHMP3 tubes without membrane (Maity et al., 2019). When

231 we incubated CHMP2A-CHMP3 and VPS4B containing membrane tubes (Figure S9A) with ATP

and Mg²⁺, complete disassembly of the tubes was observed (Figure S9B and C). In order to image 232 tube remodeling by fluorescence microscopy, VPS4B and caged ATP were incorporated into the 233 tubes wrapped with fluorescently labelled membrane. Imaging of tubes containing either only 234 caged ATP (Figures 4A and B; movie 1; Figures S10A and B; movie 2) or only VPS4B (Figures 235 4C and D; movie 3) demonstrated that photolysis used to uncage ATP did not affect the tube 236 237 structure. However, tubes containing both caged ATP and VPS4B revealed constriction and tube 238 cleavage upon ATP activation at different sites starting at 30s leading to complete disassembly 239 within 270s (Figures 4E and F; movie 4) or starting at 69s (Figures S10C and D; movie 5) or 240 50 s (**movie 6**).

Tube cleavage was further confirmed by high-speed AFM (HS-AFM) imaging. First, 241 242 CHMP2A-CHMP3 tubes with and without membrane were imaged (Figures S11A-C). A comparative height histogram showed an increase in tube height of ~8 nm for the membrane 243 coated tubes (Figure S11D), as expected for an unilamellar membrane coating. Next, membrane-244 coated tubes loaded with 10 µM caged ATP with and without UV exposure (Figure S11E; movie 245 7) were recorded by HS-AFM. In the absence of VPS4B, photolysis of the caged ATP did not 246 247 induce changes in tube morphology, consistent with the observations using fluorescence microscopy. Further, no changes in tube morphology were observed without UV exposure for an 248 extended period of time of CHMP2A-CHMP3 tubes coated with membrane and loaded with 249 VPS4B and caged ATP (Figure 5A, movie 8). However, upon UV exposure, constriction and 250 cleavage of the membrane coated tubes was observed (Figure 5B; movie 9). Kymographs along 251 the tube cross section (Figure 5C) and the evolution of the height at the constriction sites over 252 253 time (Figure 5D) reveal that complete cleavage of the membrane tube occurs within a time period of ~ 200 s of UV exposure. We conclude that VPS4B can constrict the CHMP2A-CHMP3 filaments 254 255 bound to membranes that leads to membrane cleavage, reminiscent of membrane fission.

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257 Discussion

258 The structure of the CHMP2A-CHMP3 heteropolymer demonstrates how ESCRT-III filaments polymerize into rigid structures that can stabilize and/or shape negatively curved 259 membrane necks with diameters of approximately 50 nm. Such membrane structures are present 260 at many ESCRT-catalyzed processes including vesicle and virus budding or at later stages during 261 cytokinetic midbody constriction (McCullough et al., 2018; Vietri et al., 2020). Although the 262 263 structural principles of the open ESCRT-III conformation are highly conserved between CHMP2A/CHMP3 and CHMP1B (McCullough et al., 2015), Snf7/Shrub (CHMP4) (McMillan et al., 264 2016; Tang et al., 2015) and bacterial PspA and Vipp1 (Gupta et al., 2021; Junglas et al., 2021; 265

Liu et al., 2021), differences in helical hairpin stacking and orientations of the helical arms dictates 266 the geometry of the filaments that stabilize positively curved membrane or negatively curved 267 membrane as in case of CHMP2A-CHMP3 polymerization. Furthermore, assembly modes of 268 ESCRT-III monomers can vary as shown for CHMP2A, CHMP3 and CHMP4B, which can yet 269 270 adapt another filament geometry that stabilizes positively curved membranes (Bertin et al., 2020; 271 Moser von Filseck et al., 2020). Thus the plastic nature of ESCRT-III protein conformations can 272 lead to variable ESCRT-III filament geometries that can adapt a wide range of curvatures to 273 accommodate ESCRT function in different membrane remodeling processes (Pfitzner et al., 274 2021).

A striking feature of the CHMP2A-CHMP3 polymer is the narrow range of tube diameters, 275 276 indicative of a late recruitment during the constriction process. During yeast MVB biogenesis, Vps24/Vps2 (CHMP3/CHMP2) is indeed recruited last prior to complete ESCRT-III disassembly 277 and probably completion of fission (Babst et al., 2002; Saksena et al., 2009; Teis et al., 2008). 278 Likewise, CHMP4 isoforms recruit CHMP3 and CHMP2A to HIV-1 budding sites (Johnson et al., 279 2018; Morita et al., 2011; Prescher et al., 2015). The narrow range of diameters of the CHMP2A-280 281 CHMP3 polymer structure suggests that recruitment of CHMP3 and CHMP2A molds membrane necks into approximately 40 to 50 nm diameters, which sets the stage for further constriction. 282 Comparison of the 410 and 430 Å wide structures shows that removal of only one CHMP2A-283 CHMP3 heterodimer per filament turn reduces the tube diameter by 20 Å, indicating how a 284 285 stepwise removal of heterodimers can successively induce membrane constriction.

286 Another feature of the tubular polymer is that it can assemble from single or multi-stranded 287 filaments (Effantin et al., 2013) the latter being the preferred assembly in vitro. Interaction between filaments is driven by complementary positive and negative charges. However, surprisingly, high 288 ionic strength did not disassemble the tube consistently into individual filaments but partially 289 290 preserved the multi-strand architecture in vitro. The 17.5 nm width of the six-stranded helix present in the structure fits the 17 nm wide helices imaged at the midbody (Guizetti et al., 2011), 291 292 suggesting that such spirals contain six ESCRT-III filaments. Since inter-filament interactions are electrostatic, different ESCRT-III filaments may contribute to the formation of mixed multi-stranded 293 filaments (Mierzwa et al., 2017; Pfitzner et al., 2020). In line, acidic residues within helix 4 and 294 basic residues within helix 1 have been implicated in S. cerevisiae Snf7-Vps24 interaction 295 (Banjade et al., 2019). Because basic and acidic charges within these regions are conserved in 296 297 CHMP6, CHMP4A, B, C, CHMP2B and CHMP5, filaments thereof may also form side by side via homo- or hetero-filament assembly. The loose electrostatic inter-filament interactions likely 298 299 facilitate sliding of filaments upon ESCRT-III filament remodeling by VPS4, which catalyzes

filament constriction prior to complete disassembly (Maity et al., 2019). Notably, dynamic VPS4 mediated turnover of ESCRT-III has been proposed in different membrane remodeling processes
 (Adell et al., 2014; Adell et al., 2017; Mierzwa et al., 2017).

303 CHMP2A-CHMP3 polymers have been suggested to interact with negatively charged 304 membranes (De Franceschi et al., 2018; Lin et al., 2005; Whitley et al., 2003), which is confirmed 305 by the cluster of basic residues within the membrane interaction surface. Furthermore, the 306 structure indicates that short N-terminal hydrophobic motifs, implicated in ESCRT-III function 307 (Bodon et al., 2011) (Buchkovich et al., 2013) are positioned to insert into the membrane. Although this N-terminal motif is helical in a filament structure assembled by an intermediate Vps24 308 (CHMP3) conformation (Huber et al., 2020), the corresponding helices of CHMP2A and CHMP3 309 310 are not visible in the membrane-bound structure suggesting that the putative amphipathic helices 311 can adopt different membrane insertion angles. Polymer interaction with membrane is tight, which excludes other membrane proteins thereby serving as a diffusion barrier (De Franceschi et al., 312 2018). 313

CHMP3 is dispensable for HIV-1 budding (Morita et al., 2011) and S. cerevisiae Vps24 can 314 315 be substituted by Vps2 overexpression restoring partial endosomal cargo sorting (Banjade et al., 2021). Both processes depend on CHMP2A and Vps2 interaction with CHMP4 or Snf7 (Teis et 316 al., 2008) (Morita et al., 2011). The structure suggests that CHMP3 can be structurally replaced 317 318 by CHMP2A in the polymer, indicating that CHMP2A filaments on their own may form similar 319 helices. Although CHMP2A can polymerize into circular filaments with approximate diameters of 320 40 nm that often coil up, no regular CHMP2A tube-like structures have been yet imaged in vitro 321 (Effantin et al., 2013).

322 VPS4B remodels CHMP2A-CHMP3 helical tubular structures in vitro (Caillat et al., 2015; 323 Lata et al., 2008b) inducing filament constriction and cleavage that generates dome-like end caps 324 prior to complete disassembly, which was proposed to drive membrane fission (Fabrikant et al., 2009; Maity et al., 2019). Here, we show that VPS4B constricts and cleaves CHMP2A-CHMP3 325 326 membrane-coated tubes via membrane fission likely via the formation of dome-like end-caps (Maity et al., 2019). Cleavage of membrane tubes pulled from GUVs has been reported previously 327 by employing a minimal system composed of S. cerevisiae Snf7, Vps24, Vps2 and Vps4 328 (Schoneberg et al., 2018), while another model proposed sequential recruitment of S. cerevisiae 329 Snf7, followed by Vps2-Vps24, Vps2-Did2 and Did2-Ist1 for final constriction (Pfitzner et al., 2020). 330 Our data suggest that CHMP2A-CHMP3 filaments constitute together with VPS4 a minimal 331 ESCRT-III membrane fission machinery that can constrict membrane necks with 40 to 50 nm large 332 diameters to the point of fission. It is yet unclear how many helical turns are required for 333

constriction *in vivo*, although more than one, as estimated from imaging (Adell et al., 2017), would

allow filament sliding powered by ATP-driven forces that can drive filament-induced membrane

336 constriction and cleavage. Finally, catalyzing membrane fission with a minimal machinery is well

in line with ancestral ESCRT-III function (Caspi and Dekker, 2018; Ithurbide et al., 2022).

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339 Methods

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341 **Expression and purification**

CHMP2AAC containing residues 1 to 161 was subcloned in a pMAL-c5X vector with an additional 342 TEV site at the amino terminal end and expressed as a MBP fusion protein in the C41 (DE3) E. 343 coli bacterial strain (Lucigen). Expression was induced for 1h at 37°C. Bacteria were lysed by 344 sonication in a buffer containing 50 mM HEPES pH 7.5, 300 mM NaCl, 1 mM DTT, 5 mM EDTA 345 and protease inhibitors. Cleared lysate was applied onto an amylose resin (New England Biolabs), 346 washed with buffer A (25mM HEPES 7.5, 150 mM NaCl, 1mM DTT), then with Buffer B (25 mM 347 HEPES pH 7.5, 1 M NaCl, 1 M KCl, 1mM DTT) followed by a last wash with Buffer A. Finally, 348 349 protein was eluted with buffer C (25 mM HEPES 7.5, 150 mM NaCl, 10 mM maltose). The most concentrated fraction was directly applied to size exclusion chromatography (SEC) Superdex 200 350 column (GE Healthcare) in buffer D (25 mM HEPES pH 7.5, 150 mM NaCl). 351

Full-length CHMP3 was subcloned in a pProEX-HTb vector (Life Technologies, Thermo Fisher) 352 and expressed in BL21Gold (DE3) E. coli bacterial strain (Agilent). Expression was induced for 3h 353 at 37°C and purified as described (Lata et al., 2008b) with minor modifications. Bacteria were 354 355 lysed by sonication in buffer E (25 mM HEPES pH 7.5, 150 mM NaCl, 10mM imidazole) containing protease inhibitors and the cleared lysate was applied onto a Ni²⁺-chelating sepharose (Cytivia), 356 washed extensively with lysis buffer E, and subsequently with buffer F (25 mM HEPES pH 7.5, 357 358 300 mM NaCl, 300 mM KCl, 20 mM imidazole) and buffer G (25 mM HEPES pH 7.5, 300 mM NaCl, 300 mM KCl, 50 mM imidazole). Finally, CHMP3 was eluted with buffer H (25 mM HEPES 359 360 7.5, 150 mM NaCl, 350 mM imidazole) and cleaved overnight at 4°C with Tobacco Etch Virus (TEV) protease at 1:100 (w/w) ratio in the presence of 10 mM β -mercaptoethanol. Cleaved protein 361 was then applied on a second Ni²⁺-chelating sepharose in order to remove TEV, the His-tag and 362 uncleaved protein. The final step included size exclusion chromatography (SEC) on a Superdex 363 75 column (GE Healthcare) in buffer D. CHMP3 concentrated at 300 µM was frozen for further 364 365 use.

366 CHMP2A∆C-mutants containing residues 9 to 161 and CHMP3-mutants containing residues 9 to
 367 183 were synthesized (ThermoFisher), subcloned in a pETM40 vector (PEPcore facility-EMBL

Heidelberg) and the pProEX-HTb (ThermoFisher) vector, respectively. Mutants were expressed
 and purified as described above for wild type sequences.

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371 CHMP2A-CHMP3 membrane tube generation

372 For helical tube formation as described previously (Lata et al., 2008b), 10 µM CHMP2A∆C was 373 mixed with 20 µM full-length CHMP3 and incubated for 48-72h at 4°C. After incubation, tubes 374 were harvested by centrifugation at 20,000g for 30min and the pellet containing CHMP2AAC-375 CHMP3 tubes was resuspended in buffer D. In order to wrap the CHMP2AAC-CHMP3 tubes with a lipid bilayer, the following lipid film was produced containing 70% Egg phosphatidyl choline (Egg 376 PC), 10% dioleoyl glycero phosphoserine (DOPS), 10% dioleoyl glycerol phosphoethanolamine 377 (DOPE), 10% brain phosphatidylinositol-4,5-bisphosphate (PI(4,5)P2) and 2µL of dioleoyl-sn-378 glycero-3-phosphoethanolamine-N-(lissamine rhodamine B sulfonyl) (LISS-Rhodamin PE) (all 379 Avanti Polar lipids). The lipid film was resuspended in water at a final concentration of 5 mg/mL. 380 The CHMP2A∆C-CHMP3 tubes (25 µL) were mixed with 25 µL of 2% CHAPS, 25 µL of lipids and 381 0.1 mg/mL of TEV protease (to remove the MBP from CHMP2AAC) and incubated at room 382 383 temperature for 2h. To remove free lipids/micelles and CHAPS, the tubes were dialyzed twice for 48-72h against buffer I (25mM Tris pH 7.4, 25 mM NaCl, 1 mM β-mercaptoethanol and 0.5 g of 384 Bio-Beads (Biorad). After dialysis, CHMP2A∆C-CHMP3 tubes wrapped with bilayer were 385 incubated with Bio-Beads overnight at 4°C and removed by centrifugation. The guality of the 386 bilayer wrapped CHMP2A∆C-CHMP3 tubes was assessed by negative staining EM prior to cryo-387 388 EM data collection, fluorescence microscopy imaging and HS-AFM analysis.

To test remodeling by VPS4, CHMP2A-CHMP3 tubes were incubated with 10 to 20 µM VPS4B, 5 mM MgCl₂ and 5 or 10 mM caged ATP (#A1048 Invitrogen) prior to membrane wrapping, following the protocol described above. Because deposition of the membrane onto the CHMP2A-CHMP3 protein coat requires extensive dialysis, the final VPS4 concentration present within the tubes can be only estimated from SDS-PAGE; however, the final concentration of caged ATP inserted into the tubes cannot be determined.

395

396 CHMP2A-CHMP3-VPS4 membrane tube imaging

Epifluorescence video microscopy of CHMP2A-CHMP3 membrane tubes containing VPS4B and
caged ATP was performed using an Olympus IX83 optical microscope equipped with a UPFLN
100X O-2PH/1.3 objective and an ORCA-Flash4.0 Digital sCMOS camera (Hamamatsu). A 5 μL
aliquot of ESCRT-III tube suspension was spread on a slide, covered with a glass coverslip (#1)
and sealed with twinsil speed 22 (Picodent, ref 13001002) for imaging. Caged-ATP was uncaged

using a 10s 10% 365-nm LED illumination (Figures 4A-F and movies S1-4) or using at each time
point a 100ms 30% 365-nm LED illumination (Figure S10 and movies S5 and S6). ESCRT tubes
were fluorescently imaged using a 550nm LED (10% with an exposure time of 100ms) at 1 frame/s.
Images were acquired using the Volocity software package. Images were analyzed, adjusted, and
cropped using ImageJ software.

407

408 **HS-AFM analysis**

409 The AFM images were acquired in amplitude modulation tapping mode in liquid, using high-speed atomic force microscopes (RIBM, Japan) (Maity et al., 2019; Maity et al., 2020). The HS-AFM 410 imaging was performed using USC-F1.2-k0.15 cantilevers (NanoWorld, Switzerland), an 411 412 ultrashort cantilever with a nominal spring constant of 0.15 N/m and a resonance frequency ≈ 0.5 MHz. All HS-AFM recordings were done at room temperature and in buffer D. Uncaging of the 413 caged ATP was performed by directly irradiating 365 nm UV light at the AFM sample stage using 414 an optical fiber. The membrane coated tubes were immobilized at the surface using streptavidin 415 on top of a lipid bilayer (DOPC) on mica containing 0.01% biotinylated lipid (Keya et al., 2017). 416 417 HS-AFM images were analyzed using Igor Pro, and ImageJ with additional home written plugins (Maity et al., 2022). Height measurements were performed on raw images after tilt correction. 418

419

420 Dominant negative effect of CHMP3(1-150) wild type and mutants

421 CHMP3 residues 1-150 wt or mutants were synthesized (ThermoFisher) and cloned into the pEGFP-N1 vector using restriction sites Xho1- HindIII. To determine the effect of GFP-CHMP3 (1-422 423 150) on virus like particle (VLP) production upon HIV-1 Gag expression, 293T cells were seeded into 10mm dishes and transfected 24 hr later using a Jetprime (Polyplus) technique. The cultures 424 425 were co-transfected with 0.5 µg of Rev-independent HIV-1 Gag construct and with 2 µg of either pcDNA or wild type and mutant versions of GFP-CHMP3(1-150). Twenty-four hours post 426 transfection, VLPs released into the culture medium were pelleted through sucrose. HIV-1 Gag 427 428 proteins in VLPs and cell lysates were detected by Western blotting with a mouse anti-p24 antibody (183-H12-5C). For life cell imaging cells were seeded in glass bottomed µ-dishes and 429 co-transfected with 0.8 µg of Rev-independent HIV-1 Gag construct, 0.2 µg of Gag-mCherry 430 (Jouvenet et al., 2008) and with 1 µg of either wild type or mutant GFP-CHMP3(1-150). ESCRT-431 III and Gag protein localization was analyzed by spinning disc confocal microscopy 24 hr post 432 433 transfection in HeLA cells.

434

435 Cryo-EM sample preparation and data collection

Cryo-electron microscopy. 3.5 µL of sample were applied to glow discharged (45s 30 mA) 1.2/1.3 436 Ultrafoil holey grids (Quantifoil Micro Tools GmbH, Germany) and they were plunged frozen in 437 liquid ethane with a Vitrobot Mark IV (Thermo Fisher Scientific) (100% humidity, temperature 20°C, 438 6 s blot time, blot force 0). The grids were pre-screened on the 200kV Glacios electron microscope 439 440 (Thermo Fischer Scientific) at the IBS (Grenoble) and data were collected at the beamline CM01 441 of the ESRF (Grenoble, France) (Kandiah et al., 2019) on a Titan Krios G3 (Thermo Fischer 442 Scientific) at 300 kV equipped with an energy filter (Bioquantum LS/967, Gatan Inc, USA) (slit width of 20 eV). 5028 movies were recorded automatically on a K2 summit direct detector (Gatan 443 Inc., USA) with EPU (Thermo Fisher Scientific) for a total exposure time of 5 s and 200 ms per 444 445 frame resulting in 25 frame movies with a total dose of $\sim 24 \text{ e}^{-}/\text{Å}^{2}$. The magnification was 130,000x (1.052 Å/pixel at the camera level). The defocus of the images was adjusted between -0.5 and 446 -1.5 µm in 0.2 µm steps. For the high ionic strength unwinding of the CHMP2A-CHMPA filament 447 the same grid and freezing conditions have been used as described above and images have been 448 recorded on the Glacios electron microscope using a K2 direct electron detector. 449

450

451 EM image analysis and 3D reconstructions

The workflow of the image analysis is shown in **Figure S2**. Electron beam-induced sample motion 452 on the recorded movie frames was corrected using MotionCor2 (Zheng et al., 2017) and the 453 contrast transfer function (CTF) was estimated with CTFFIND4 (Rohou and Grigorieff, 2015). 454 9,207 filaments were manually picked from 5,027 micrographs using the EMAN2 program 455 e2helixboxer.py (Tang et al., 2007). All subsequent data processing steps were carried out in 456 RELION3 (Scheres, 2012; Zivanov et al., 2018) unless mentioned otherwise. Initially, 89,122 457 overlapping segments were extracted with ~90% overlap between boxes of 768 x 768 pixels and 458 down sampled to a pixel size of 2.104 for initial classification steps. Several rounds of 2D 459 classification resulted in class averages that could be classified into 5 main different groups based 460 on the filament diameter, without membrane (380 Å diameter (7.4%), 410 Å diameter (51.4%), 461 430 Å diameter (31.2%), 470 Å diameter (1.9%) and 490 Å diameter (0.3%)). In order to 462 compensate for potential mis-assignment of diameters to the segments due to inaccuracies in 2D 463 464 classification, we assigned to each entire tube a diameter based on the class assignment of the 465 corresponding segments. If more than 80% of segments of a particular tube were belonging to classes assigned to a particular diameter, this entire tube would be assigned this diameter for the 466 467 subsequent steps. After re-extraction of segments with ~95% overlap, another round of 2D 468 classification was performed for each diameter group. Most populated 2D classes with filament

diameter of 430 Å and 410 Å were chosen for further processing and analysis. For determination 469 of helical symmetry, the sum of power spectra from a smaller subset (1,904 and 3,993 segments 470 from one 2D class each for 430 Å and 410 Å respectively) was calculated for both filament groups. 471 The resulting average power spectrum (Figures S1C and D) was analyzed for estimation of 472 473 helical symmetry parameters using the web tool helixplorer (http://rico.ibs.fr/helixplorer/). Based 474 on a prior visual inspection of the PS, we made following hypotheses: the layer line with a 475 maximum seemingly on the meridian could be the helical rise or the pitch (given the large diameter 476 of the tube and possibilities that selected 2D class averages contained a number of slightly outof-plane tilted segments). Given those two hypotheses, and allowing any cyclic symmetries, we 477 explored possible helical symmetries matching the experimental PS, giving a list of 20 and 15 478 479 symmetries to test for the 430 Å and 410 Å diameter classes, respectively. Those symmetries 480 were applied the real-space 2D class-averages using SPRING program on segclassreconstruct.py (Desfosses et al., 2014) in order to generate initial models and narrowing 481 down possible symmetry solutions to 14 and 10, by discarding those giving aberrant density 482 distribution. Using those initial models, each of the remaining symmetry solutions was tested for 483 3D refinement in RELION3, and the resulting maps inspected for high-resolution features such as 484 clear secondary structures, allowing to determine the helical parameters to be 18 Å pitch, 2.72 Å 485 rise, 54.39° twist, 6.6 units/turn and C2 point symmetry for 430 Å, and 9 Å pitch, 1.43 Å rise, 57° 486 twist, 6.3 units/turn and C1 point symmetry for 410 Å diameter filaments. 487

488 In order to select a more homogeneous subset of segments, we applied 3D classification and the classes (containing 25,353 and 11,396 segments for diameters 430 and 410 Å) were chosen for 489 a final round of 3D auto-refine reconstruction that converged to a 2.74 Å rise and 54.37° twist for 490 430 Å, and 1.44 Å rise and 57.04° twist for 410 Å diameter filaments. Using soft protein-only 491 masks, the final resolutions were estimated at 3.3 Å and 3.6 Å for the 430 Å and 410 Å diameter 492 493 filaments, at the FSC (Fourier shell correlation) 0.143 cutoff criterion (Rosenthal and Henderson, 494 2003). The maps were sharpened with bfactors of -96.57 Å² (430 Å) and -101.52 Å² (410 Å). Local resolution was estimated in RELION3 (Scheres, 2012) and the density maps were rendered in 495 UCSF Chimera (Pettersen et al., 2004). The statistics of the EM map are summarized in Table 496 497 **S1**.

498

499 Atomic modelling and validation

500 The SWISS-MODEL (Waterhouse et al., 2018) server was used to create homology models of 501 human CHMP3 and CHMP2A, using the open conformation of CHMP1B (PDB ID 6TZ4) as a 502 reference model. Helices with residues 15-52, 57-117 and 120-151 for CHMP3 and 14-51, 56-116

and 119-150 for CHMP2A were initially fit into the EM density as separate rigid bodies using 503 504 Chimera and then adjusted in Coot (Emsley et al., 2010). Further, the rest of the N-terminal and C-terminal residues, as well as the connecting loops were manually built and adjusted in Coot. 505 506 The CHMP2A-CHMP3 heterodimer model was then expanded by helical symmetry in each 507 direction in order to get 10 such dimers surrounding the central dimer. Thus, a total of 11 helical 508 symmetry-related dimers were again checked in Coot, before applying the first round of real-space 509 refinement in PHENIX (Adams et al., 2010) with non-crystallographic symmetry (NCS) restraints. 510 NCS, along with SS (secondary structure) restraints were then used for a second round of realspace refinement. At last, the symmetry-related dimers were removed and the central CHMP2A-511 CHMP3 dimer was saved as the final model. The statistics of the final models were tabulated 512 513 using MolProbity (Williams et al., 2018) and are summarized in **Table S1** and map versus atomic model FSC plots were computed in PHENIX (Afonine et al., 2018). All structure figures were 514 generated with UCSF Chimera, ChimeraX (Goddard et al., 2018) and PyMOL (W. Delano; The 515 PyMOL Molecular Graphics System, Version 1.8 Schrödinger, LLC, http://www.pymol.org). 516 Sequence alignments were performed with Clustal Omega (Goujon et al., 2010) and ESPript 517 518 (Robert and Gouet, 2014).

519

520 Data Availability

521 Cryo-EM maps and models were deposited to the PDB and EMDB with the following codes: 522 membrane-bound CHMP2A-CHMP3, 430 Å diameter (PDB ID 7ZCG, EMD-14630) and 523 membrane-bound CHMP2A-CHMP3, 410 Å diameter (PDB ID 7ZCH, EMD-14631).

524

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537

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- 546
- 547 **Competing interests:** The authors declare no competing interests.
- 548
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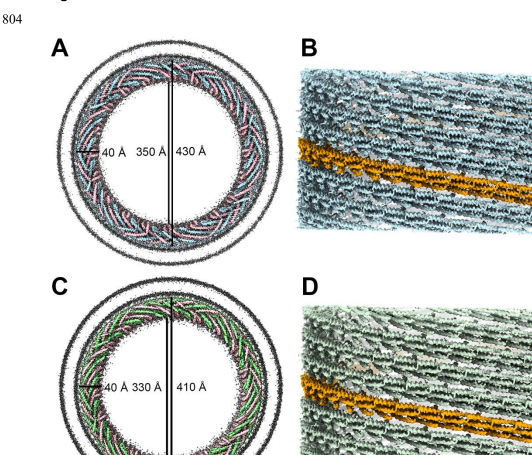
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Pitch 175 Å

29 Å



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807 Figure 1: Cryo-EM structure of CHMP2A-CHMP3 membrane-coated helical polymers

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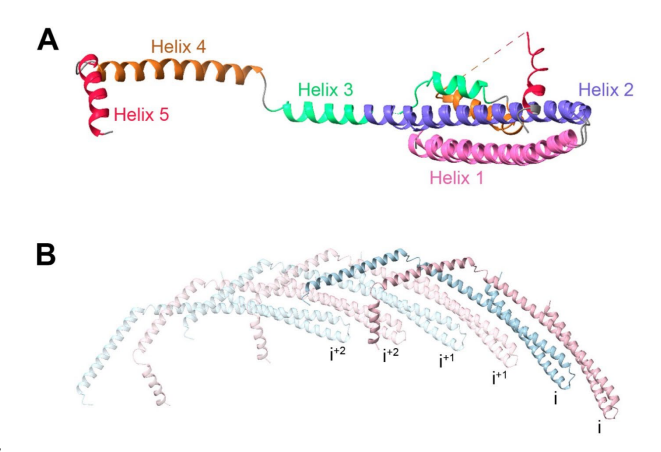
CHMP2A CHMP3

(A) Density map of the reconstructed 430 Å diameter CHMP2A-CHMP3 membrane tube with the
top view looking down the helical axis. The helical arrangement of CHMP2A (light blue) and
CHMP3 (pink) inside the bilayer membrane (dark grey) is shown. The thickness, and the inner
and outer diameter of the helical protein tube are also marked in Å.
(B) Side view of the helical polymer without the lipid membrane. One left-handed filament is

indicated in orange, and the thickness of one filament and the pitch of the helical assembly are

also marked.

- (C) Density map of the reconstructed 410 Å diameter CHMP2A-CHMP3 membrane tube with the
- top view looking down the helical axis. The helical arrangement of CHMP2A (green) and CHMP3
- (pink) inside the bilayer membrane (dark grey) is shown. The thickness, and the inner and outer
- diameter of the helical protein tube are indicated in Å.
- (D) Side view of the helical polymer without the lipid membrane. One left-handed filament is
- indicated in orange, and the thickness of one filament and the pitch of the helical assembly areindicated.
- (E) Cryo-EM density of the single repeating unit of the 430 Å diameter polymer formed by the
 heterodimer of CHMP2A (light blue) and CHMP3 (pink) is indicated.
- (F) Ribbon representation of the atomic model of CHMP2A (light blue) CHMP3 (pink)
 heterodimer.
- 826



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828 Figure 2: Atomic model and architecture of the CHMP2A-CHMP3 helical polymer.

(A) Ribbon diagram of C α superposition of the closed and open CHMP3 conformations (Helix 1:

pink, Helix 2: purple, Helix 3: light green, Helix 4: brown, Helix 5: red).

(B) Three interlocked copies of CHMP2A-CHMP3 heterodimer are shown as ribbons. Helix 4 and

5 of CHMP3 (pink) interact with four subsequent protomers. Helix 4 of CHMP2A (light blue) also

- 833 makes similar interactions.
- 834
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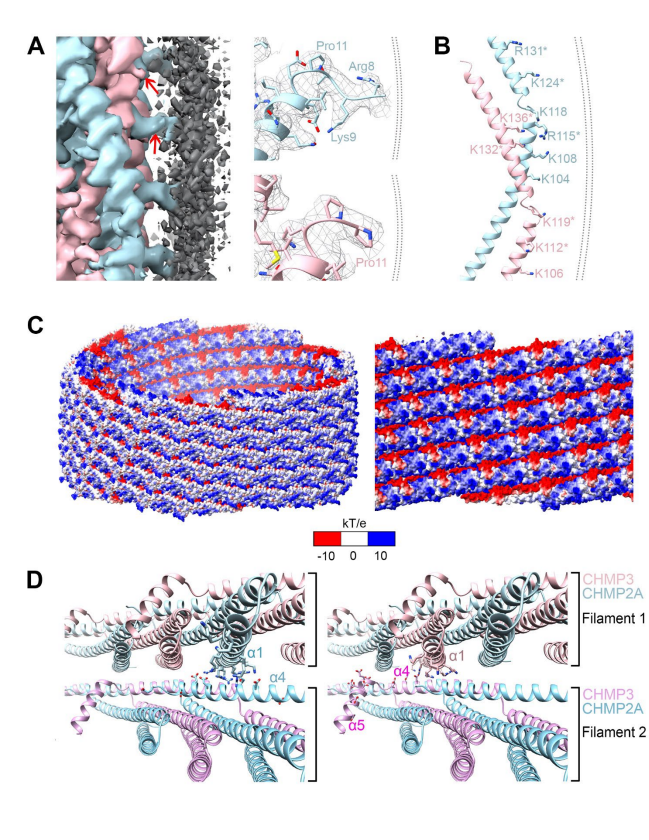


Figure 3: Membrane interaction of the CHMP2A-CHMP3 filament.

- (A) Left panel, zoomed-in view of the membrane-bound CHMP2A-CHMP3 filament, highlighting
- 840 the interface between lipid membrane (dark grey) and CHMP2A (light blue) and CHMP3 (pink).
- Red arrows are pointing to the N-termini of both CHMP2A and CHMP3 that are oriented towards
- the lipid bilayer. Right panel, the orientation of the N-termini is determined by Pro11 of CHMP2A
- 843 (light blue) and CHMP3 (pink) shown with the density map.
- (B) Ribbon diagram of the CHMP2A-CHMP3 heterodimer indicating the basic residues (sticks)
 oriented towards the membrane. Basic residues conserved in yeast Vps2 and Vps24 are marked
 by asterisks.
- (C) Electrostatic potential map of the CHMP2A-CHMP3 filament (left panel, tilted, side view), revealing the exposure of the cluster of basic charges, a small negatively charged surface and a neutral surface to the membrane. Right panel, zoomed-in view of the electrostatic surface of the
- 850 inside of the polymer showing clusters of negative charges in one filament juxtaposed to positive
- 851 charges of the neighboring filament.
- 852 (D) Close-up of the inter-filament interactions. Ribbon diagram of two neighboring filaments
- showing the basic and acidic residues of CHMP2A (left panel) and CHMP3 (right panel) implicated
- in electrostatic inter-filament interactions.
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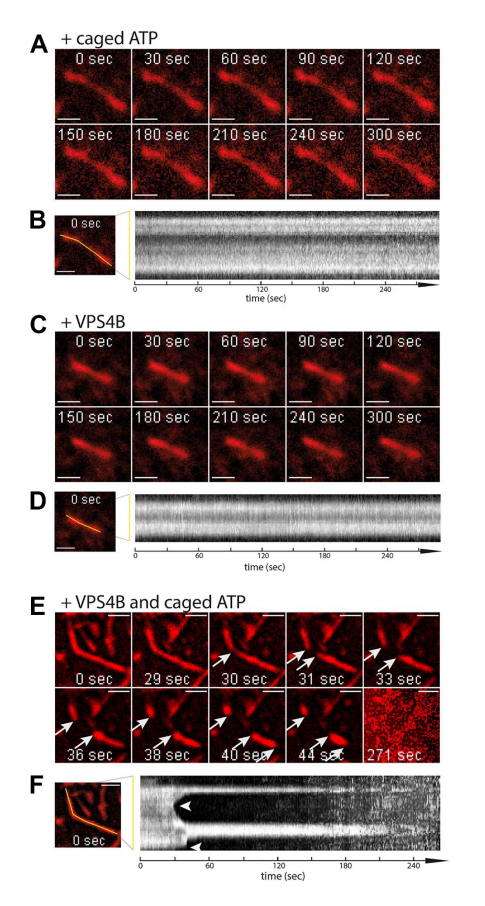
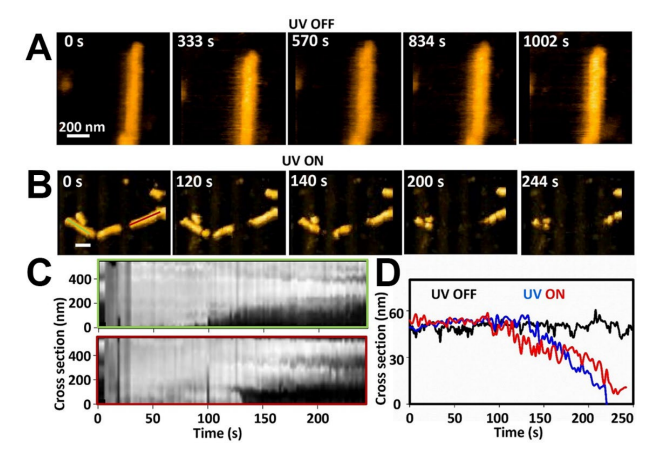


Figure 4: Imaging of VPS4B and ATP induced cleavage of CHMP2A-CHMP3 membrane

858 coated tubes.

- (A) A CHMP2A-CHMP3-caged ATP containing membrane coated tube was activated at 365 nm
- 860 (10%, 10s) to uncage ATP and imaged over 300s (movie S1). Scale bar, 1μm.
- (B) The kymograph of the tube shows that the tube stays intact over the imaging time.
- (C) A CHMP2A-CHMP3-VPS4B containing membrane coated tube was activated at 365 nm (10%,
- 10s) to uncage ATP and imaged over 300s. Scale bar, 1µm.
- (D) The kymograph of the tube shows that the tube stays intact over the imaging time. (A) to (D)
- demonstrate that imaging at 550 nm to visualize the membrane tube and ATP uncaging at 365
- nm did not change the tube structures (movie S2).
- 867 (E) Imaging of a CHMP2A-CHMP3-VPS4B-caged ATP containing membrane-coated tubes
- following ATP uncaging (365 mm, 10%, 10s) reveals constriction and cleavage of the tube at 30
- s followed by a shrinking event from both sides. Another shrinking event is observed at 40s.
- Eventually all tubes were fully disassembled 271 s (movie S3 and S4). Scale bar, 1µm.
- (F) The kymograph of the tube indicates the kinetics of cleavage and shrinking.

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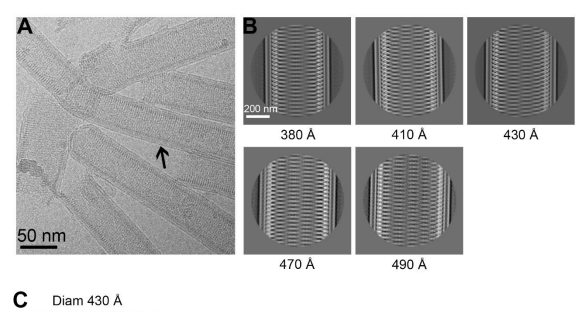
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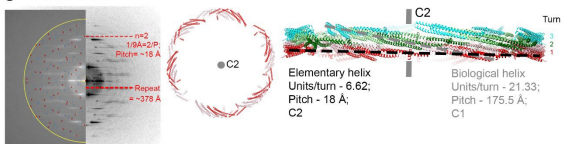
Figure 5. Constriction, cleavage, and disassembly of membrane coated CHMP2A-CHMP3
tube.

- (A) Snapshots of HS-AFM images of CHMP2A-CHMP3 tubes coated with membrane and loaded
- with 5 μM VPS4B, 10 mM caged ATP, in absence of UV irradiation. Scale bar, 200 nm.
- (B) As in A, but upon 365 nm UV irradiation. Scale bar, 200 nm.
- (C) Kymograph representation of the height vs time along the two lines in panel B (leftmost image)
- throughout the movie S10.
- (**D**) Example of height vs time profile of the membrane coated tube in absence of UV irradiation
- (in black), and in presence of UV irradiation (in blue and red).

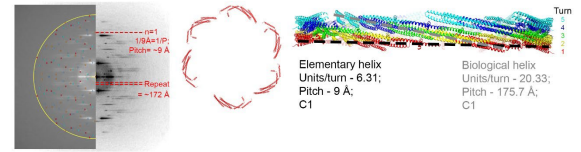
884

885	Supplemental Figures
886 887	Structural basis of CHMP2A-CHMP3 ESCRT-III polymer assembly and membrane
888	cleavage
889	
890	Kimi Azad ¹ , Delphine Guilligay ¹ , Cecile Boscheron ¹⁺ , Sourav Maity ²⁺ , Nicola De Franceschi ^{1,3+} ,
891	Guidenn Sulbaran ¹ , Gregory Effantin ¹ , Haiyan Wang ¹ , Jean-Philippe Kleman ¹ , Patricia
892	Bassereau ³ , Guy Schoehn ¹ , Wouter H Roos ² , Ambroise Desfosses ^{1*} , Winfried Weissenhorn ^{1*}
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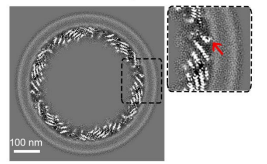




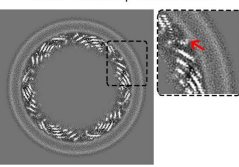
D Diam 410 Å



E Diam 430 Å Map Central Slice



Diam 410 Å Map Central Slice



901 Figure S1: Cryo-EM data processing of CHMP2A-CHMP3 membrane-coated tubes and

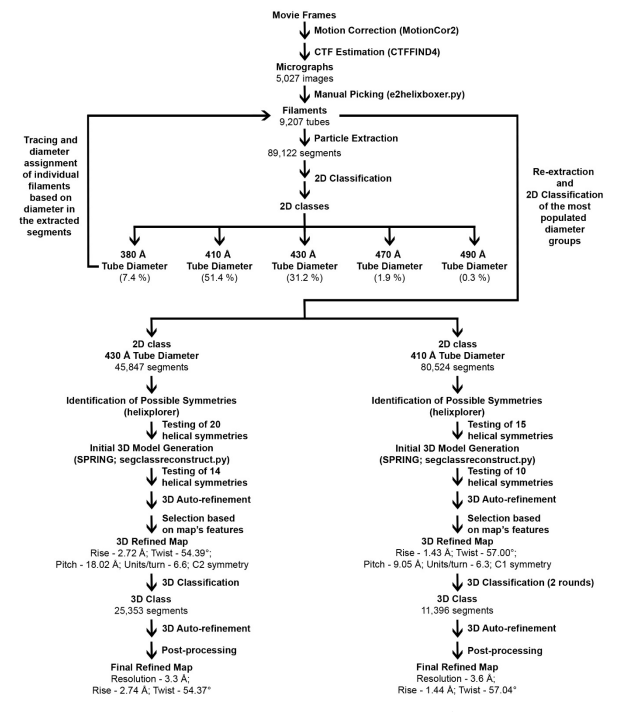
902 helical symmetry analyses

903 (A) Representative cryo-electron micrograph of CHMP2A-CHMP3 membrane tubes, with an arrow
 904 pointing to the lipid bilayer. Scale bar, 50 nm.

905 **(B)** Selected 2D class averages of manually picked datasets, arranged according to the tube diameter ranging from 380 to 490 Å as indicated.

907 **(C)**, **(D)** Helical symmetry determination and representation of the elementary and biological 908 helices for the 430 Å and 410 Å diameter tubes. Left panel, the sum of the 2D power spectra of 909 segments corresponding to one class-average show in both cases a maximum on or near the

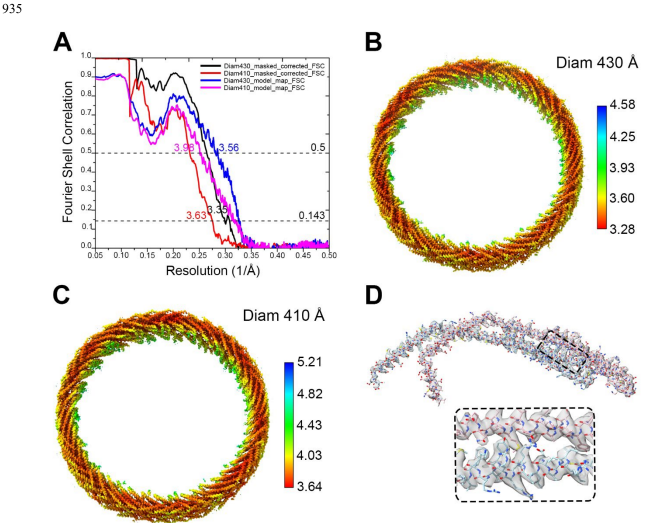
- meridian corresponding to the pitch (Bessel order n=1 for C1 helix; n=2 for C2 helix) instead of
- 911 the axial rise, due to the large helical diameter and possible inclusion of slightly out-of-plane tilted
- 912 segments in the class-average. The left half of the sum of the power spectra show the calculated
- 913 position from helixplorer of the two first maxima of each Bessel function corresponding to the
- 914 determined symmetry. Middle panel, six asymmetric units (CHMP2A-CHMP3 dimers; in red) of
- the elementary helix are represented. The symmetry related protomers of the 430 Å diameter are
- 916 colored in light red.
- 917 (C), (D) Right panel, side view of the elementary helix with turns (red, yellow, green, blue and
- aqua) indicated with a black dashed line. The grey dashed line follows one turn of the biological
 helix. The central grey line (in C) highlights the C2 symmetry axis. Symmetry parameters of both
- 920 the elementary and biological helices are indicated.
- 921 (E) Central slices looking down the helical axis of the cryo-EM 3D reconstructions of the 430 and
- 922 410 Å diameter tubes. The red arrows in the right zoom-in images indicate the density of the N-
- 923 terminal region prone to insert into the lipid bilayer.
- 924



925

Figure S2: Cryo-EM image processing workflow of 430 and 410 Å diameter tubes structure determination.

Basic image processing strategy used for helical 3D reconstruction and refinement of 430 and 410
Å diameter tubes is shown. Helical filaments were segmented and classified based on the tube
diameter. Segment subsets were subjected to symmetry determination (<u>http://rico.ibs.fr/helix</u>
<u>plorer/</u>) and initial 3D model generation in SPRING, followed by symmetry refinement and final 3D
structure refinement in RELION. A complete description of the processing workflow is provided in
'Materials and Methods' section.



936

Figure S3: 430 and 410 Å diameter tubes FSC curves, local resolution maps and atomic
 model fitting.

939 (A) FSC curves for the 430 Å (black) and 410 Å (red) diameter tube maps, with the resolutions at

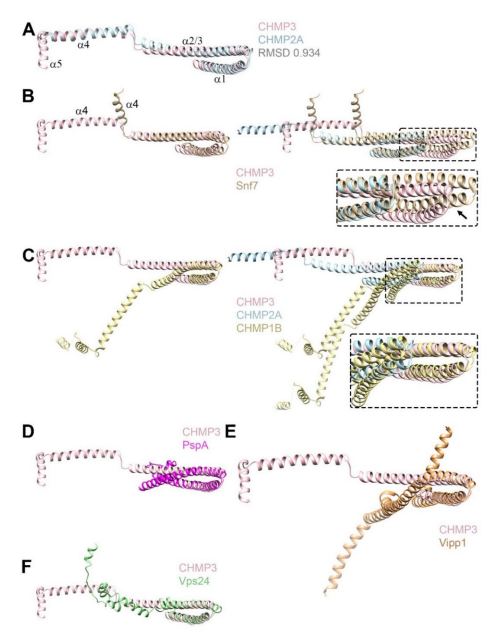
940 the FSC cut-off of 0.143 are indicated. Model versus map FSC curves, with the resolutions at the

FSC cut-off of 0.5 are indicated for the 430 Å (blue) and 410 Å (pink) diameter tube maps.

Local resolution estimates are mapped onto the 430 Å **(B)**, and 410 Å **(C)** diameter tube cryo-EM density maps and the color keys (right) highlight the local resolution values in Å.

944 **(D)** The refined atomic model of CHMP2A-CHMP3 dimer was fit into the corresponding cryo-EM 945 density map of the 430 Å diameter tube. The inset (below) represents the zoomed-in view of the

- 946 fitted model, indicating CHMP2A and CHMP3 helices and the corresponding map.
- 947
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Figure S4: Comparison of ESCRT-III open conformations highlights their versatile polymerization modes

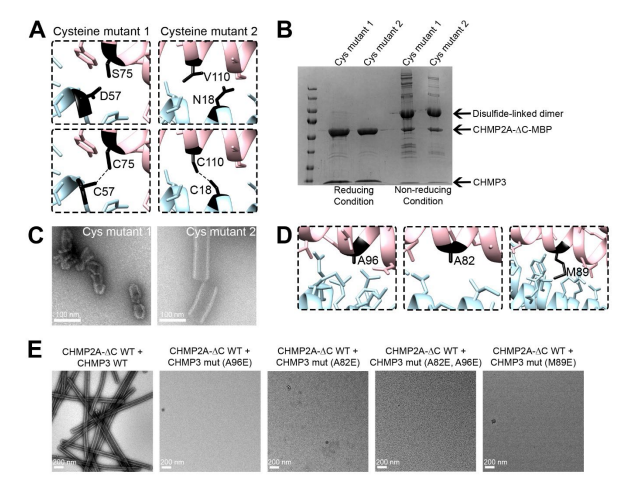
952 (A) Ribbon diagram of Cα superposition of CHMP2A and CHMP3 reveals an RMSD of 0.934 Å.

953 (B) Ribbon diagram of Cα superposition of CHMP3 with an Snf7 monomer (left panel) and with

the homodimer (crystallographic dimer) (right panel), which indicates different hairpin interaction (arrow) for the second protomer and different orientations of Snf7 helix 4.

956 (C) Ribbon diagram of C α superposition of CHMP3 and CHMP1B (left panel) and the

- superposition of the CHMP2A-CHMP3 heterodimer onto the CHMP1B homodimer (right panel)
 indicate the differences in helical hairpin stacking (zoom, right panel) and orientations of the C terminal helical arms (helices 3 to 5).
- 960 (**D**) Ribbon diagram of Cα superposition of CHMP3 (pink) with PspA and (**E**) with Vipp1.
- 961 (F) Ribbon diagram of Cα superposition of CHMP3 with an intermediate Vps24 conformation that
 962 forms filaments on its own.
- 963

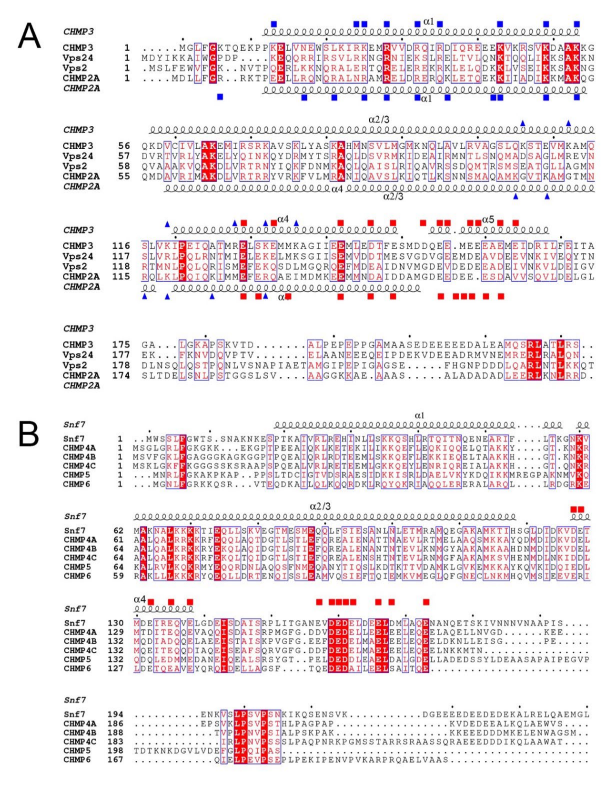


964 965

966 Figure S5: Structure-based mutagenesis of CHMP2A-CHMP3 heterodimer formation and

967 polymerization *in vitro*.

- 968 **(A)** Close-up views of the pairs of residues (black) mutated to cysteine to induce the formation of 969 disulfide-linked CHMP2A (light blue) - CHMP3 (pink) heterodimers upon polymerization.
- 970 (B) Cysteine cross-linking of the CHMP2A-CHMP3 heterodimer. Mutant CHMP2A_D57C was
- 971 incubated with CHMP3_S75C and CHMP2A_N18C with CHMP3_V110C to induce polymerization
- as reported for wild-type CHMP2A and CHMP3 (Lata et al., 2008b). SDS-PAGE analysis showing
- 973 that both CHMP2A_D57C-CHMP3_S75C and CHMP2A_N18C-CHMP3_V110C formed disulfide-
- 974 linked dimers under non-reducing SDS PAGE conditions.
- 975 (C) Negative staining electron micrographs showing regular tube formation for CHMP2A_N18C-
- OCHMP3_V110C (right), while CHMP2A_D57C-CHMP3_S75C (left) produced only shorter tubes.
 Scale bar, 100 nm.
- 978 **(D)** Close-up views of the CHMP3 interface residues A96, A82 and M89E tested for heterodimer 979 formation and polymerization.
- 980 **(E)** Negative staining electron micrographs of CHMP2A-CHMP3 wild-type and mutants 981 (CHMP3_A96E, CHMP3_A82E, CHMP3-A82E_A96E and CHMP3_M89E) assemblies as 982 indicated. Scale bar, 200 nm.
- 983



985 986

987 Figure S6: ESCRT-III sequence alignment.

(A) Sequence alignment of CHMP3 (AF219226), S. cerevisiae Vps24 (QHB09957), S. cerevisiae
 Vps2 (P36108.2) and CHMP2A (NM_198426.3). Secondary structure elements are shown for
 CHMP3 above the sequence and for CHMP2A below the sequence alignment. Blue triangles

indicate basic residues of CHMP3 (above) and CHMP2A (below) exposed at the membrane
 binding interface. Blue rectangles show basic residues and red squares conserved acidic residues

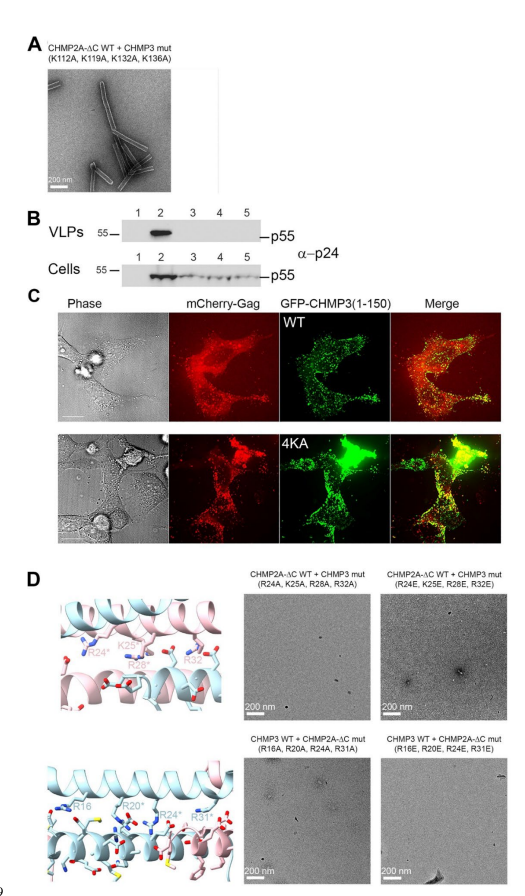
993 exposed at the interface between filaments.

(B) Sequence alignment of *S. cerevisiae* Snf7 (Z73197.1) and its secondary structure (pdb 5FD9),

995 CHMP4A (NM_014169.5), CHMP4B (NM_176812.5) CHMP4C (NM_152284), CHMP5

996 (NM_016410.6) and CHMP6 (NM_024591.5). Conserved acidic residues implicated in inter-

filament interactions in the CHMP2A-CHMP3 polymer are indicated as red squares.



1000 Figure S7: Structure-based mutagenesis of CHMP2A-CHMP3 polymer formation

1001 **(A)** Negative staining electron micrograph showing regular tube formation by CHMP2A-1002 CHMP3 K112A, K119A, K132A, K136A polymerization. Scale bar, 200 nm.

1003 (B) The substitution of four basic residues in GFP-CHMP3(1–150) 4KA (K112A, K119A, K132A,

1004 K136A) does not diminish its dominant-negative effect on HIV-1 budding. Western blot analyses

of Gag released from Gag expressing cells as Gag-VLPs (upper panel) and detection of Gag in
 total cell extracts (lower panel): lane 1, Gag expression; lane 2, Gag and GFP-VPS4A E228Q
 expression; lane 3, Gag and GFP-CHMP3(1-150) expression; lane 4, Gag and GFP-CHMP3(1 150) 4KA expression.

1009 **(C)** Representative fluorescence images of HeLa cells transfected with Gag/mCherry-Gag and

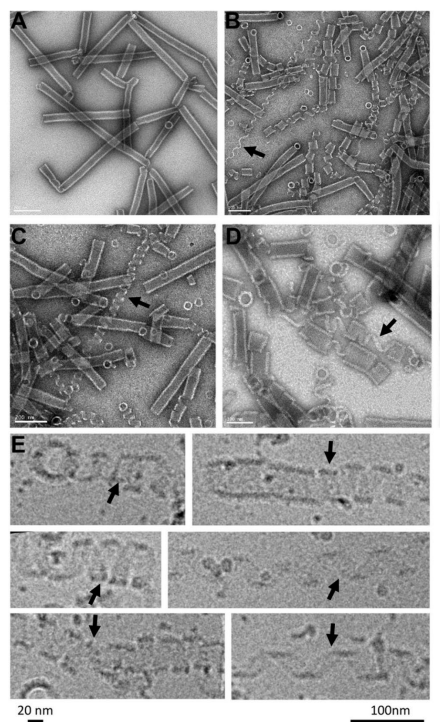
- 1010 CHMP3(1-150) or CHMP3(1-150)4KA. Cellular distribution of wild-type and mutant KA GFP-1011 CHMP3(1–150) indicates predominantly plasma membrane and intracellular localization as well 1012 as co-localization with mCherry-Gag. Scale bar, 10 µm.
- 1013 (D) Negative staining electron micrographs showing no polymer formation of CHMP3 mutants
- 1014 (upper left panel, close-up of a ribbon diagram illustrating the interface residues) R24A, K25A,

1015 R28A, R32A (upper middle panel) and R24E, K25E, R28E, R32E (upper right panel) with

1016 CHMP2A. (Lower panel) CHMP2A mutants (lower left panel, close-up of a ribbon diagram

- illustrating the interface residues) R16A, R20A, R24A, R31A (lower middle panel) and R16E,
- 1018 R20E, R24E, R31E (lower right panel) did not polymerize with CHMP3 *in vitro*. Scale bar, 200 nm.
- 1019

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1023 Figure S8: High ionic strength unwinds the CHMP2A-CHMP3 filaments.

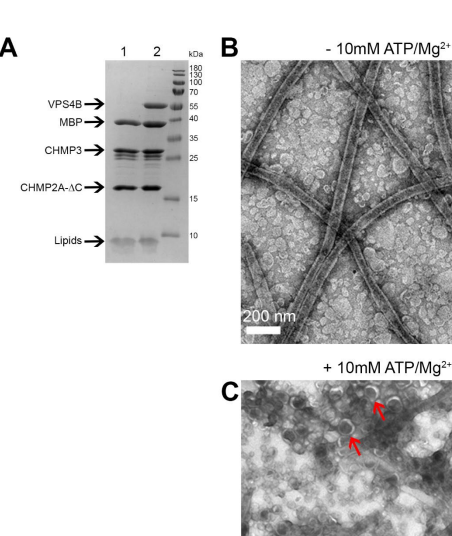
1024 **(A)** Negative staining electron micrographs showing CHMP2A-CHMP3 wild-type polymers after 1025 treatment with 1M NaCl **(B, C)** and 1M KCl **(D)**.

1026 (E) Close-up of cryo-EM images shows unwinding of ~20 nm wide filaments corresponding to the

six-start helix observed in the structure. Single and multi-stranded unwound filaments are

indicated by arrows.





1030

1031 Figure S9: Incorporation of VPS4B and ATP into CHMP2A-CHMP3 membrane tubes

200 nm

1032 induces their disassembly.

1033 **(A)** SDS-PAGE analyses of purified CHMP2A-CHMP3 polymers; lane 1, CHMP2A-CHMP3 polymers cleaved with TEV and coated with a lipid bilayer; lane 2 CHMP2A-CHMP3 polymers,

1035 TEV cleaved and incorporation of VPS4B prior to lipid bilayer coating.

1036 Negative staining electron micrographs of CHMP2A-CHMP3-VPS4B membrane-coated polymers

- ¹⁰³⁷ before (**B**) and after (**C**) incubation with ATP and Mg²⁺ Red arrows point to membrane vesicles
- 1038 resulting from tube cleavage. Scale bar, 200 nm.

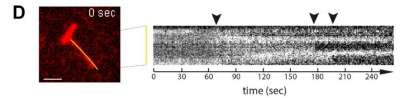


	0 sec	30 sec	60 sec	90 sec	120 sec
	150 sec	180 sec	210 sec	240 sec	300 sec
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0	30	60	90	120	150	180	210	240	270
				time	(sec)				

C + VPS4B and caged ATP

0 sec	58 sec	69 sec	80 sec	125 sec
		A		
165 sec	171 sec	177 sec	178 sec	262 sec
1	3	*		



1040

1041 Figure S10: Imaging of VPS4B and ATP induced cleavage of CHMP2A-CHMP3 membrane 1042 coated tubes.

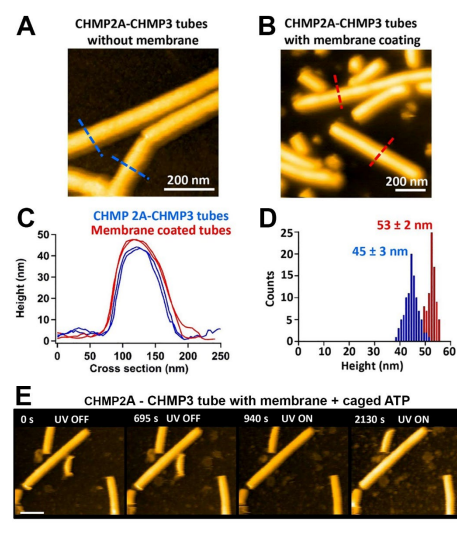
1043 **(A)** A CHMP2A-CHMP3-caged ATP membrane coated tube was activated at 365 nm (365 nm 1044 30%, 100 ms at each time point) to uncage ATP and imaged over 282s, which indicated that uncaging did not change the tube structure (movie S3).

(B) The kymograph of the tube (yellow line) shows that the tube stays intact over the imaging time.
 (C) Imaging of a CHMP2A-CHMP3-VPS4B-caged ATP membrane-coated tube following ATP uncaging (365 nm, 30%, 100 ms at each time point) reveals cleavage of the tube at several sites

1049 over the imaging time (movie S5). Scale Bar, 1µm.

1050 **(D)** The kymograph of the tube (yellow line) indicates tube cleavage (arrows) over the imaging 1051 time.

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- 1053



1054 1055

1056 Figure S11. Height distribution of CHMP2A-CHMP3 tubes with and without membrane-1057 coating.

- 1058 (A) AFM image of CHMP2A-CHMP3 tubes without membrane.
- 1059 (B) AFM image of CHMP2A-CHMP3 tubes coated with membrane.
- 1060 **(C)** Cross-section of AFM images of CHMP2A-CHMP3 in panel A (blue dotted line) and panel B (red dotted line).
- 1062 **(D)** Height histogram of CHMP2A-CHMP3 tubes with (red) and without (blue) membrane coating with respect to the surface.
- 1064 **(E)** Snapshots HS-AFM images of membrane-coated tubes loaded with 10 mM caged ATP taken
- 1065 with and without UV irradiation. The UV was turned on from 700 s onwards. Scale bar, 200 nm.
- 1066 1067

1068 **Movies**

- 1069
- 1070 Movie 1: Fluorescence microscopy imaging of CHMP2A-CHMP3 membrane coated tubes.
- 1071 A CHMP2A-CHMP3-caged ATP membrane-coated tube was activated at 365 nm for 10s to 1072 uncage ATP and imaged over 291s.
- 1073 Movie 2: Fluorescence microscopy imaging of CHMP2A-CHMP3 membrane coated tubes.
- 1074 Another dataset of CHMP2A-CHMP3-caged ATP membrane-coated tube activated at 365 nm for 1075 100 ms at each time point to uncage ATP and imaged over 300s.
- 1076 Movie 3: Fluorescence microscopy imaging of CHMP2A-CHMP3 membrane coated tubes.
- 1077 A CHMP2A-CHMP3-VPS4B membrane-coated tube was activated at 365 nm for 10s to uncage 1078 ATP and imaged over 251s.
- 1079 **Movie 4: Fluorescence microscopy imaging of CHMP2A-CHMP3 membrane coated tubes.** 1080 Imaging of a CHMP2A-CHMP3-VPS4B-caged ATP membrane-coated tube following ATP 1081 uncaging (365 nm, 10s). This demonstrates tube fission followed by a shrinking event from both 1082 sides
- 1083 Movie 5: Fluorescence microscopy imaging of CHMP2A-CHMP3 membrane coated tubes.
- Another dataset of CHMP2A-CHMP3-VPS4B-caged ATP membrane-coated tube following ATP uncaging (365 nm, 100 ms at each time point) reveals cleavage of the tube at several sites over the imaging time.
- 1087 Movie 6: Fluorescence microscopy imaging of CHMP2A-CHMP3 membrane coated tubes.
- 1088 Imaging of a CHMP2A-CHMP3-VPS4B-caged ATP membrane-coated tube following ATP
- 1089 uncaging (365 nm) showing at 36s a shrinking event from the end of a tube and at 51s a cleavage 1090 of the tube.
- 1091 **Movie 7: HS-AFM imaging of CHMP2A-CHMP3 membrane-coated tubes.** HS-AFM movie of 1092 membrane-coated CHMP2A-CHMP3 tubes loaded with 10 mM caged ATP, taken before and after 1093 UV (365 nm) irradiation. Imaging time 5 seconds/frame.
- 1094 **Movie 8: HS-AFM imaging of CHMP2A-CHMP3 membrane-coated tubes.** HS-AFM movie of 1095 membrane-coated CHMP2A-CHMP3 tubes loaded with 5 μM VPS4B and 10 mM caged ATP, 1096 taken without UV irradiation. Imaging time 3 seconds/frame.
- 1097 **Movie 9: HS-AFM imaging of CHMP2A-CHMP3 membrane-coated tubes.** HS-AFM movie of 1098 membrane-coated CHMP2A-CHMP3 tubes loaded with 5 μM VPS4B and 10 mM caged ATP, 1000 taken with 265 nm LIV/irrediction. Imaging time 2 accords/frame.
- taken with 365 nm UV irradiation. Imaging time 2 seconds/frame.
- 1100
- 1101 1102

1103 Table S1 – Cryo-EM data collection, refinement and validation statistics

	Membrane-bound CHMP2A + CHMP3 (Tube Diameter 430 Å) (EMD-14630, PDB 7ZCG)	Membrane-bound CHMP2A + CHMP3 (Tube Diameter 410 Å) (EMD-14631, PDB 7ZCH)	
Data collection and processing			
Magnification	130,000x	130,000x	
Voltage (kV)	300	300	
Electron exposure per frame (e ⁻ /Å ²)	0.96	0.96	
Number of frames	25	25	
Defocus range (µm)	0.5-1.5	0.5-1.5	
Pixel size (Å)	1.052	1.052	
Refined helical symmetry	54.37°, 2.74 Å	57.04°, 1.44 Å	
Point group symmetry	C2	C1	
Initial particle images (no.)	45,847	80,524	
Final particle images (no.)	25,353	11,396	
Map resolution (Å)	3.3	3.6	
FSC threshold	0.143	0.143	
Refinement			
Initial model used (PDB code)	de novo	de novo	
Model resolution (Å)	3.6	4.0	
FSC threshold	0.5	0.5	
Map sharpening <i>B</i> factor (Å ²)	-96.57	-101.52	
Model-to-map fit			
Correlation coefficient, CC (mask)	0.80	0.77	
Model Composition			
Chains	22	22	
Nonhydrogen atoms	27,148	27,148	
Protein residues	3,366	3,366	
<i>B</i> factor (Å ²)			
Protein	74.18	90.17	
R.m.s. deviations			
Bond lengths (Å)	0.004	0.004	
Bond angles (°)	0.758	0.762	
Validation			
MolProbity score	1.86	1.89	
Clashscore	15.23	21.23	
Rotamer outliers (%)	0.00	0.00	
Cbeta deviations (%)	0.00	0.00	
Ramachandran plot			
Favored (%)	97.02	97.68	
Allowed (%)	2.98	2.32	
Disallowed (%)	0.00	0.00	

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