Tethering by Uso1 is dispensable: The Uso1 monomeric globular head domain interacts with SNAREs to maintain viability.

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Summary

38 Uso1/p115 and RAB1 tether ER-derived vesicles to the Golgi. Uso1/p115 contains a 39 globular-head-domain (GHD), a coiled-coil (CC) mediating dimerization/tethering and a 40 C-terminal region (CTR) interacting with golgins. Uso1/p115 is recruited to vesicles by 41 RAB1. Paradoxically, genetic studies placed Uso1 acting upstream of, or in conjunction 42 with RAB1 (Sapperstein et al., 1996). We selected two missense mutations in uso1 resulting in E6K and G540S substitutions in the GHD permitting growth of otherwise 43 44 inviable rab1-deficient Aspergillus nidulans. Remarkably, the double mutant suppresses 45 the complete absence of RAB1. Full-length Uso1 and CTRA proteins are dimeric and the 46 GHD lacking the CC/CTR is monomeric irrespective of whether they carry or not 47 E6K/G540S. Microscopy showed recurrence of Uso1 on puncta (60 sec half-life) 48 colocalizing with RAB1 and less so with early Golgi markers Sed5 and GeaA/Gea1/Gea2. Localization of Uso1 but not of Uso1^{E6K/G540S} to puncta is abolished 49 by compromising RAB1 function, indicating that E6K/G540S creates interactions 50 51 bypassing RAB1. By S-tag-coprecipitation we demonstrate that Uso1 is an associate of 52 the Sed5/Bos1/Bet1/Sec22 SNARE complex zippering vesicles with the Golgi, with 53 Uso1^{E6K/G540S} showing stronger association. Bos1 and Bet1 bind the Uso1 GHD directly, 54 but Bet1 is a strong E6K/G540S-independent binder, whereas Bos1 is weaker but 55 becomes as strong as Bet1 when the GHD carries E6K/G540S. AlphaFold2 predicts that 56 G540S actually increases binding of GHD to the Bos1 Habc domain. In contrast, E6K seemingly increases membrane targeting of an N-terminal amphipathic α -helix, 57 58 explaining phenotypic additivity. Overexpression of E6K/G540S and wild-type GHD complemented uso1_Δ. Thus, a GHD monomer provides the essential Uso1 functions, 59 60 demonstrating that long-range tethering activity is dispensable. Therefore, when 61 enhanced by E6K/G540S, Uso1 binding to Bos1/Bet1 required to regulate SNAREs bypasses both the contribution of RAB1 to Uso1 recruitment and the reported role of 62 63 RAB1 in SNARE complex formation (Lupashin and Waters, 1997), suggesting that the 64 latter is consequence of the former.

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Introduction

68 Vesicular traffic at the ER/Golgi interface is the cornerstone of the secretory pathway 69 (Barlowe and Miller, 2013; Weigel et al., 2021). In current models, in which traffic across 70 the Golgi is driven by cisternal maturation (Day et al., 2013; Pantazopoulou and Glick, 71 2019), COPII vesicles generated at specialized domains of the ER fuse homotypically 72 and heterotypically to form and feed the earliest Golgi cisternae (Rexach et al., 1994). 73 As straightforward as this step might seem, it involves a sophisticated circuitry of 74 regulation. Actual fusion is in part mediated by compartmental-specific sets of four-75 membered SNARE protein complexes (SNARE bundles) (Malsam and Sollner, 2011; 76 Pelham, 2001; Rizo and Sudhof, 2012). Most SNARES are type II single TMD proteins, 77 whose N-terminal cytosolic domain contains nearly all the polypeptide, excepting a few 78 lumenal residues. Like any other transmembrane proteins, SNAREs are synthesized in 79 the ER. This implies that they have to travel to compartments of the cell as distant as the 80 plasma membrane in a conformation that precludes them of catalyzing what would be a 81 calamitous fusion of non-cognate donor and acceptor compartments. Achieving the 82 strictest specificity is particularly challenging in the ER-to-Golgi stage that, as the first 83 step in the secretory pathway, represents an obligate point of transit for each and every 84 transmembrane SNARE. Therefore, the only SNARES acting in this first step are the Qa 85 Sed5, the Qb Bos1, the Qc Bet1 and the R-SNARE Sec22, which form the bundle 86 mediating fusion of carriers that coalesce into cisternae (McNew et al., 2000).

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Given the central role played by the secretory pathway in the physiology of every 88 89 eukaryotic cell, it is unsurprising that this step involves regulatory factors which are 90 essential for cell survival. One is the SM (Sec1, Munc-18) protein Sly1, which promotes 91 SNARE bundle formation (Bracher and Weissenhorn, 2002; Peng and Gallwitz, 2002; 92 Thomas et al., 2019) Another is the TRAPPIII complex, which interacts with the external 93 coat of COPII carriers and acts as a guanine nucleotide exchange factor (GEF) for RAB1 94 (Bracher and Weissenhorn, 2002; Cai et al., 2007; Galindo et al., 2021; Joiner et al., 95 2021; Lord et al., 2011; Peng and Gallwitz, 2002; Pinar and Peñalva, 2020; Riedel et al., 96 2017; Thomas et al., 2018; Thomas et al., 2019). This small GTPase is a key player that 97 transiently recruits protein effectors from the cytosol to donor and acceptor membranes 98 (Sogaard et al., 1994) and regulates SNARE assembly through an as yet undefined 99 mechanism (Lupashin and Waters, 1997; Sapperstein et al., 1996). One RAB1 effector 100 is a fungal protein denoted Uso1, whose highly conserved metazoan homologue is p115. 101 These are homodimers with a globular N-terminal head and a long C-terminal coiled-coil 102 region characteristic of tethering proteins, which bring donor and acceptor membranes 103 into the distance at which v- and t-SNAREs can engage into the productive trans-SNARE

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104 complex that mediates membrane fusion (Cao et al., 1998; Nakajima et al., 1991;
105 Sapperstein et al., 1996; Sapperstein et al., 1995; Seog et al., 1994; Yamakawa et al.,
106 1996).

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108 Despite most Golgi tethers are functionally redundant, Uso1 is unique in that it is an 109 essential protein. uso1-1, a S. cerevisiae amber mutation truncating most, but not all the 110 coiled-coil region, is viable, yet further upstream truncation removing the complete 111 coiled-coil is lethal, which was taken as evidence that tethering is the essential function 112 of Uso1 (Seog et al., 1994). In addition, as the coiled-coil is predicted to mediate 113 dimerization, it is broadly accepted that Uso1 is 'just' an essential homodimer that tethers 114 vesicles to the acceptor membrane. However, genetic evidence stubbornly indicates that 115 Uso1 plays additional functions related with SNAREs. For example, Sapperstein et al 116 showed that SNAREs function downstream of Uso1 (Sapperstein et al., 1996). Notably, 117 the view that Uso1 is a mere RAB1 effector was challenged by the observation that 118 overexpressing Ypt1 (yeast RAB1) rescues lethality of $uso1\Delta$, whereas the reciprocal is 119 not true, indicating that Uso1 acts upstream of or in conjunction with RAB1 (Sapperstein 120 et al., 1996).

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122 Our laboratory is interested in deciphering the domains of action of RAB GTPases in the 123 genetic and cell biological model organism Aspergillus nidulans (Pinar and Peñalva, 124 2021). We have previously gained mechanistic insight into the activation of RAB11 by 125 TRAPPII by exploiting a forward genetic screen for mutations bypassing, at the restrictive 126 temperature, the essential role of the key TRAPPII subunit Trs120 (Pinar et al., 2019; 127 Pinar et al., 2015; Pinar and Peñalva, 2020). In this type of screen, a strain carrying a ts 128 mutation in the gene-of-interest is mutagenized and strains bypassing lethality at the 129 restrictive temperature are identified and characterized molecularly. A well-130 characterized, conditionally lethal rab1 mutation is available (Pinar et al., 2013), enabling 131 us to investigate pathways collaborating with RAB1 in anterograde traffic. We isolated 132 two uso1 missense mutations causing substitutions in the globular head domain (GHD). 133 When combined together, these rescued the lethality resulting from $rab1\Delta$ and promoted 134 the localization of the protein to early Golgi cisternae by increasing Uso1 binding to the 135 cytosolic region of the Qa SNARE Bos1 and, potentially, by improving the interaction of 136 an N-terminal amphipathic α -helix with membranes. Importantly, we show that 137 endogenous expression of a protein consisting solely of the double mutant GHD, or 138 overexpression of double mutant or wild-type GHD, rescue the lethality resulting from 139 $uso1\Delta$, even though the GHD is monomeric. Our results show that one essential role of 140 RAB1 is recruiting Uso1 to membranes, and that the essential role of Uso1 is not

141 tethering membranes, but rather regulating the formation of the cognate SNARE bundle,

142 indicating that Uso1 is a component of the SNARE fusion machinery.

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Results

144 Missense mutations in *uso1* rescue the lethality resulting from $rab1\Delta$

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146 rab1^{A136D} (hereby rab1^{ts}) mutants do not grow at 37°C. However, when we plated UVmutagenized conidiospores of the rab1^{ts} mutant at this temperature, we obtained 147 148 colonies showing different degrees of growth, presumably carrying mutations rescuing 149 the lethality resulting from *rab1*^{ts}. One was chosen for further characterization. By sexual 150 crosses and parasexual genetics this strain was shown to carry a single suppressor mutation, denoted *su1rab1*^{ts}, that co-segregated with chromosome VIII. Meiotic mapping 151 narrowed su1rab1^{ts} to the vicinity (2 cM) of hisC. 40 kb centromere distal from hisC lies 152 153 AN0706 (Figure 1A) encoding Aspergillus nidulans Uso1, a conserved effector of RAB1. 154 Sanger sequencing revealed the presence of a G16A transition (denoted E6K) resulting 155 in Glu6Lys substitution in the uso1 gene of su1rab1^{ts}.

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To determine if the remaining suppressor strains were allelic to *su1rab1*^{ts}, we sequenced 157 158 uso1 from a further 13 isolates (Figure 1B). Of these, four were rab1^{ts} pseudo-revertants 159 that had acquired a functionally acceptable mutation in the altered codon, and eight 160 carried uso1 E6K, suggesting that the screen was close to saturation. However, one 161 mutation was found to be a different missense allele, su85rab1ts (denoted G540S) 162 resulting in Gly540Ser substitution. Single mutant strains carrying these uso1 mutations 163 showed no growth defect, indicating that E6K and G540S were unlikely to result in loss-164 of-function, and suggesting instead that mutant strains had acquired features that made 165 them largely independent of RAB1. These findings were unexpected, because in 166 Saccharomyces cerevisiae overexpression of Uso1 does not rescue the lethality of ypt1 Δ 167 mutants (Ypt1 is the yeast RAB1 homologue)(Sapperstein et al., 1996).

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To demonstrate that $uso1^{E6K}$ and $uso1^{G540S}$ were causative of the suppression, we 169 reconstructed them by homologous recombination. These reverse-genetic alleles 170 rescued viability of rab1^{ts} at 37°C to a similar extent as su1rab1^{ts} and su85rab1^{ts} (Figure 171 1C). *uso1*^{G540S} was the strongest suppressor, such that *rab1*^{ts} *uso1*^{G540S} double mutants 172 173 grew nearly as the wt at 37°C. Nevertheless, the two alleles showed additivity, and a triple mutant carrying uso1^{E6K}, uso1^{G540S} and rab1^{ts} grew at 42°C, unlike either single 174 175 mutant (Figure 1C). These data, together with the genetic mapping above, established that uso1^{E6K} and uso1^{G540S} are responsible for the suppression phenotype, 176

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178 RAB1 recruits Uso1/p115 to uncoated COPII vesicles and early Golgi cisternae (Allan et al., 2000). Therefore, uso1^{E6K/G540S} might, by increasing the affinity of Uso1 for RAB1, 179 compensate for the reduction in the amount of the GTPase resulting from rab1^{ts}. 180 However, Figure 1D shows that both uso1^{E6K} and uso1^{G540S} rescue the lethality resulting 181 from the complete ablation of $rab1\Delta$ at 30°C, with the strongest $uso1^{G540S}$ suppressor 182 183 rescuing viability even at 37°C, and the double mutant rescuing rab1∆ even at 42°C (Figure 1D). In contrast, $uso1^{E6K/G540S}$ did not rescue the lethality resulting from $arf1\Delta$, nor 184 185 from sed5 Δ or sly1 Δ , the syntaxin and the SM protein which are crucial for the formation 186 of the ER/Golgi SNARE bundle (Figure 1—figure supplement 1), indicating that Uso1 187 plays a role acting downstream of RAB1 and upstream of or in conjunction with the 188 SNARE machinery. This role is essential for survival (Figure 1—figure supplement 1).

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190 E6K affects a previously undetected N-terminal helix, whereas G540S is located 191 in a loop near the end of the armadillo domain.

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193 1103-residue A. nidulans Uso1 is similar in size to 961-residue p115 (bovine) and notably 194 shorter than S. cerevisiae Uso1p (1790 residues) (Yamakawa et al., 1996). Thus far, 195 atomic structures of Uso1/p115 are limited to the 600-700 residue GHD, which consists 196 of a highly conserved α -catenin-like armadillo-fold (An et al., 2009; Heo et al., 2020; 197 Striegl et al., 2009). In silico analyses robustly predict that the approximately C-terminal 198 half of Uso1/p115 consists of a coiled-coil that mediates tethering and dimerization, but 199 this region has not been characterized beyond low resolution EM studies (Yamakawa et 200 al., 1996). Neither crystal structures nor predictions provided information about the N-201 terminal extension in which Glu6Lys lies.

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203 Thus, we used AlphaFold2, imposing or not the condition that the protein is a dimer (see 204 below). Figure 2A shows the monomer and dimer models with the highest confidence 205 scores (see Figure 2—figure supplement 1). Like their relatives, Uso1 from A. nidulans 206 contains an N-terminal GHD including a previously unnoticed short α -helix in which Glu6 207 affected by E6K lies. This N-terminal extension is followed by ~34 α -helices arranged into 12 tandem repetitions of armadillo repeats (ARM1-ARM12; residues 17 through 208 209 564), each containing three right-handed α -helices except for the first two repeats. 210 Altogether, the armadillo repeats resemble the shape of a jai alai basket. Downstream 211 of the GHD, AlphaFold2 predicts a long extended coiled-coil (CC) between residues 674 212 through 1082, which would mediate dimerization (see below) (Figure 2A). The CC ends

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at a conserved C-terminal region (CTR) (Figure 2A and B), which includes an also Cterminal segment rich in acidic residues. In Uso1, twelve out of the last seventeen amino
acids are Asp/Glu (Figure 2A and B). The CC and CTR regions will be collectively
denoted the CCD domain.

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218 Even though the GHD contains Glu6 and Gly540 in the N-terminal helix and at the 219 beginning of armadillo α -helix 29, respectively, intramolecular or intermolecular (in the 220 context of a homodimer, see below) distances between these residues are long, arguing 221 against the possibility that they bind a common target as components of the same 222 interaction surface (Figure 2C). Indeed, the synthetic positive effect of the mutations 223 would be consistent with their rescuing viability through different mechanisms. The 224 previously unnoticed short α -helix predicted by AlphaFold between Phe2 and Lys12 is 225 amphipathic (Figure 2D). Glu6 lies on the polar side of this helix, such that Glu6Lys 226 increases its overall positive charge (three of the four polar residues are Lys or Arg). As 227 discussed below, it is tempting to speculate that this helix facilitates membrane 228 recruitment.

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230 Coiled-coil mediated dimerization of Uso1: the globular head is monomeric

231 It has been proposed that p115 alternates between closed and open 232 conformations to hide or expose a RAB1 binding site present in the CCD (Beard et al., 233 2005). This would be mediated by intramolecular interactions between the globular 234 domain and the C-terminal acidic region, which would be disrupted by the competitive 235 binding of golgins GM130 and giantin to the latter. We addressed whether E6K/G540S 236 promotes a conformational change in Uso1, or, alternatively, a change in the 237 oligomerization status of the protein, by analytical ultracentrifugation. We designed 238 seven constructs carrying a C-terminal His tag (Figure 3). Two corresponded to the full-239 length protein with or without E6K and G540S substitutions. The second pair included 240 wild-type and doubly-substituted versions of C-terminally truncated Uso1 lacking the CTR (Uso1 Δ CTR and Uso1^{E6K/G540S} Δ CTR). The third corresponded to wild-type and 241 doubly-substituted versions of the globular domain, denoted Uso1 GHD and Uso1 242 GHD^{E6K/G540S}. The seventh construct corresponded to the CCD domain (i.e. CC + CTR) 243 244 and was denoted Uso1 CCD. All seven proteins were expressed in bacteria, purified by 245 nickel-affinity and size-exclusion chromatography and analyzed by sedimentation 246 velocity ultracentrifugation. These experiments revealed that all protein preparations 247 were essentially homogeneous, and thus they were used to determine the corresponding 248 Svedberg coefficients. In addition, by dynamic light scattering we determined the

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translational diffusion coefficients of the constructs. With these values we deduced themolecular mass of the different proteins using Svedberg's equation.

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252 Wild-type and E6K/G540S full-length Uso1s showed the same sedimentation 253 coefficients, demonstrating that the mutations do not induce a large conformational 254 change that would have been reflected in changes in sedimentation velocity due to 255 differences in frictional forces. Molecular masses deduced from the Svedberg equation 256 indicated that these full-length proteins are homodimers, in agreement with previous 257 literature (Figure 3). Ablation of the conserved C-terminal region (CTR) did not result in 258 any significant change in the sedimentation coefficient (Figure 3, panels 3 and 4 vs. 1 and 2), irrespective of the presence or absence of the substitutions, discarding the model 259 260 in which the CTR would interact with the GHD to maintain a hypothetical closed 261 conformation (Beard et al., 2005). In addition, the molecular masses of the ΔCTR 262 proteins correspond to a dimer, implying that the acidic region is not involved in dimerization either. 263

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265 Notably the GHD, whether wild-type or mutant, behaved as a monomer (Figure 3, panels 266 5 and 6), which has important implications described below. In contrast, the coiled-coil domain, with a predicted molecular mass of 52 kDa, behaves as a dimer of ca. 100 kDa 267 268 (Figure 3, panel 7). The sedimentation coefficient of the CCD is markedly slower than 269 that of the 70 kDa monomeric GHD, suggesting an elongated shape. These 270 observations, together with the dimeric nature of the construct lacking the CTR, showed 271 that dimerization is mediated by the CCD. The absence of 443 residues corresponding 272 to the CCD plus CTR domains in the GHD construct and the monomeric nature of the 273 latter compared to full-length Uso1 dimer did not result in a commensurate decrease in 274 sedimentation coefficient, which changed from 4.8 S to 3.7 S in the wild-type (note that 275 the change in Mr goes from 246 kDa in full-length Uso1 to only 68 kDa of the 276 GHD)(Figure 3). These data strongly support AlphaFold2 predictions depicting Uso1 as 277 a dimer with a globular head and an extended coiled-coil that would retard sedimentation 278 of the protein very substantially.

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As with full-length Uso1, the double substitution did not alter the sedimentation coefficient of the GHD (Figure 3, panels 5 and 6). To buttress the conclusion that the GHD is a monomer irrespective of the presence or absence of the mutations, we performed sedimentation velocity experiments using different protein concentrations ranging from 0.5 to 5 μ M (Figure 3—figure supplement 1, A and B). In all cases the GHD behaved as a monomer. Sedimentation profiles of Uso1 GHD lacking the His-tag showed

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a similar behavior, establishing that the monomeric state of the mutant is not due to the
tag at the C-terminal position hindering dimerization (Figure 3—figure supplement 1,C).
Therefore, sedimentation experiments did not detect any change in tertiary or quaternary
structures between wild-type and mutant GHD, which is important for the interpretation
of genetic data that will be discussed below.

In summary, (i) Uso1 is a dimer; (ii) The C-terminal acidic region is dispensable for dimerization and does not mediate an equilibrium between closed and open conformations; (iii) The globular domain of Uso1 is a monomer; (iv) The coiled-coil domain of Uso1 is a dimer; (v) the double E6K G540S substitution does not promote any conformational shift in Uso1, nor does it result in a change in the oligomerization state of the protein.

297 The punctate pattern of localization of USO1-GFP is dependent on RAB1

The membranous compartments of the Golgi are not generally stacked in fungi, permitting the resolution of cisternae, which appear as punctate structures in different steps of maturation, by wide-field fluorescence microscopy (Losev et al., 2006; Matsuura-Tokita et al., 2006; Pantazopoulou and Peñalva, 2011; Pinar et al., 2013; Wooding and Pelham, 1998). While Uso1 is predicted to localize to the Golgi, studies of its localization in fungi are limited (Cruz-García et al., 2014; Sánchez-León et al., 2015). Therefore, we tagged the *A. nidulans uso1* gene endogenously with GFP

305 Figure 4A and video 1 depicting a software-shadowed 3D reconstruction of a 306 Uso1-GFP hypha, as well as consecutive sections of deconvolved z-stacks in Figure 4B 307 show that Uso1-GFP localizes to puncta polarized towards the tip, often undergoing 308 short-distance movements (see Figure 4C and Figure 4—figure supplement 1). These 309 puncta are smaller and more abundant than those reported for other markers of the 310 Golgi, which suggested that they might represent domains rather than complete 311 cisternae. Notably, 3D (x, y, t) movies revealed that Uso1 puncta are transient, 312 recurrently appearing and disappearing with time (Figure 4C). That this recurrence did 313 not reflect that the puncta go in-and-out of focus was established with 4D (x, y, z, t) 314 movies, which revealed a similar behavior of Uso1 irrespective of whether 3D or 4D 315 microscopy was used (video 2) Uso1 foci with confidence, we constructed movies with 316 middle planes only (i.e. 3D x, y, t series). After careful adjustment of live imaging 317 conditions, we achieved a 2 fps time resolution with relatively low bleaching for time 318 series consisting of 400 photograms (video 3). These conditions sufficed to track Uso1 319 puncta over time using kymographs traced across linear ROIs covering the complete 320 width of the hyphae (Figure 4C). However, as the abundance of Uso1 puncta made 321 automated analysis of Uso1 maturation events troublesome, we tracked them manually

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with the aid of 3D (x,y,t) representations generated with Imaris software combined with direct observation of photograms in movies (Figure 4C and Figure 4—figure supplement 1). The boxed event magnified in Figure 4E (see video 4) illustrates a prototypical example. The right Figure 4E montage shows frames corresponding to this event for comparison. We analyzed n = 60 events, which gave an estimation of the average halflife of Uso1 residing in puncta of 60 sec +/- 25.26 S.D. (Figure 4D).

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329 Nakano and co-workers have proposed that the transfer of lipids and proteins between 330 ER exit sites (ERES) and the early Golgi occurs through a kiss-and-run mechanism 331 (Kurokawa et al., 2014). Because Uso1-GFP punctate structures resemble, in size and 332 abundance, ER exit sites labelled with COPII components, we studied Uso1-GFP cells 333 co-expressing Sec13 endogenously labeled with mCherry (Bravo-Plaza et al., 2019). 334 The maximal intensity projection (MIP) shown on Figure 5A, and video 5 show that the 335 two markers are closely associated, but only in a few instances they showed 336 colocalization. These examples did not represent simple overlap, as they were found to 337 colocalize in the Z dimension using orthogonal views or montages (Figure 5B and C). 338 These observations have not been pursued further with time-resolved sequences, but at 339 the very least we can conclude that the reporters are closely associated in space. In view 340 of this, we determined that Uso1 structures originate downstream of COPII-mediated ER 341 exit. Therefore, we investigated, using sarA6, a temperature-sensitive allele of the gene 342 encoding A. nidulans SAR1 (Hernández-González et al., 2014), whether the punctate 343 Uso1 structures are dependent on this master GTPase regulating COPII biogenesis. 344 Figure 5D shows that this is indeed the case; the number of Uso1-GFP puncta was 345 significantly reduced relative to the wt when cells were shifted from 28°C to 37°C, 346 indicating that Uso1 populates a membrane compartment with Golgi identity.

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348 To determine the 'sub-Golgi' localization of Uso1 puncta, we filmed Uso1-GFP along with 349 different Golgi markers (Figure 6). Uso1-GFP showed no overlap (Pearson's coefficient 350 0.17 ± 0.06 S.D., n = 16 cells) with cisternae labeled with mCherry-Sec7, the late Golgi 351 ARF1 GEF that is a prototypic marker of the TGN (Arst et al., 2014; Day et al., 2018; 352 Galindo et al., 2016; Halaby and Fromme, 2018; Losev et al., 2006; McDonold and 353 Fromme, 2014; Pantazopoulou, 2016; Pantazopoulou and Glick, 2019; Richardson et 354 al., 2016; Richardson et al., 2012) (Figure 6; video 6). In contrast, visual observation of 355 cells expressing mCh-Sed5 and Uso1-GFP revealed substantial, yet incomplete, overlap 356 of the reporters (Figure 6), reflected in a Pearson's coefficient of 0.44 \pm 0.04 S.D., n =357 15 cells (Figure 6). The Qa syntaxin Sed5 drives fusion of COPII vesicles with early Golgi 358 cisternae, with Qb, Qc and R-SNAREs Bet1, Bos1 and Sec22, and mediates intra-Golgi 359 trafficking, with Qb, Qc and R-SNARES Sft1, Gos1 and Ykt6), respectively (Banfield et 360 al., 1995; McNew et al., 2000; Parlati et al., 2002; Pelham, 1999; Wooding and Pelham, 361 1998). These data suggest that Uso1 localizes to a subset of early Golgi cisternae/membranes containing Sed5. GeaA^{Gea1,2} is the only A. nidulans homologue of 362 the S. cerevisiae early Golgi ARF1 GEFs Gea1 and Gea2 dwelling at the early Golgi 363 364 (Arst et al., 2014; Gustafson and Fromme, 2017; Muccini et al., 2022; Pantazopoulou, 2016; Park et al., 2005; Wright et al., 2014). Overlapping of Uso1 with GeaA^{Gea1,2} was 365 366 more conspicuous than with Sed5 (Figure 6), which was reflected in an increased 367 Pearson's coefficient to 0.52 ± 0.06 S.D., n = 16 cells. Of note, mammalian GeaA (GBF1) 368 and Uso1 (p115) interact (Garcia-Mata and Sztul, 2003). Taken together, these data 369 indicate that Uso1 localizes to Golgi cisternae in early stages of maturation.

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371 In both mammalian cells and in yeasts Uso1/p115 has been shown to be recruited to 372 early Golgi membranes by RAB1, which is activated by the TRAPPIII GEF on COPII 373 vesicles after they bud from the ER (Allan et al., 2000; Cai et al., 2007; Lord et al., 2011; 374 Yuan et al., 2017). Therefore, we imaged Uso1 and RAB1, which revealed that indeed 375 Uso1 colocalized with RAB1 (Pearson's 0.61 \pm 0.07 S.D., n = 20 cells) (Figure 6). 376 Altogether, the above microscopy data strongly suggest that Uso1 is transiently recruited 377 to early Golgi membrane domains enriched in RAB1, agreeing with the accepted view 378 that RAB1 acts by recruiting Uso1.

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380 Uso1 delocalization after RAB1 impairment rescued by E6K G540S.

381 We next tested if the subcellular localization of Uso1 is dependent on RAB1, and 382 if this dependency can be bypassed by E6K/G540S. To this end we first showed that in 383 a RAB1⁺ background endogenously tagged wild-type and E6K/G540S Uso1-GFP have 384 the same punctate localization pattern (Figure 7A), and that both supported vigorous wt 385 growth (Figure 7B lanes 1, 3, 5 and 7), indicating that the tagged proteins are functional. Next, we introduced in these strains rab1^{ts} (Jedd et al., 1995; Pinar et al., 2013) by 386 387 crossing. This allele, which completely prevents growth at 37°C, is a hypomorph at 388 permissive (25-30°C) temperatures, which permitted testing RAB1 dependence under 389 standard microscopy conditions (28°C). Figure 7A shows that wt Uso1-GFP was largely 390 delocalized to the cytosol by rab1^{ts}. This establishes that Uso1 localization to membranes 391 is subordinated to RAB1. Notably, the wt punctate pattern of Uso1 localization was 392 restored by the Uso1 E6K/G540S mutant substitution, correlating with correction of the 393 synthetic growth defect (Figure 7B, compare lanes 2, 4, 6 and 8). All these data indicated 394 that the localization of Uso1 is compromised when RAB1 function is impaired, and that

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the E6K/G540S substitutions augment Uso1 affinity for a membrane anchor(s)
independent of RAB1. They additionally suggest that the principal physiological role of
RAB1 is ensuring the proper localization of Uso1.

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Delocalization of Uso1 in the *rab1^{ts}* background was not solely dependent on RAB1.
Wild-type *uso1-GFP* displayed a synthetic negative interaction with *rab1^{ts}* (Figure 7B,
compare lanes 2 and 4 at 30°C), suggesting that the presence of GFP in the C-terminus
interferes with a RAB1-independent mechanism that facilitates its recruitment to
membranes (see below)

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405 Genetic evidence that a network involving the CTR and the Grh1/Bug1 golgin

406 contributes to the recruitment of Uso1 to membranes.

To follow up the above observation, we focused on golgins. In mammalian cells, the Cterminal region of p115 interacts with GM130, a golgin which is recruited to the early
Golgi by GRASP65 (Beard et al., 2005). The equivalent proteins in budding yeast are
denoted Bug1 and Grh1 (Behnia et al., 2007)(Figure 8A).

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412 GRASP65 contains two C-terminal PDZ [post synaptic density protein (PSD95), 413 Drosophila disc large tumor suppressor (Dlg1), and zonula occludens-1 protein (zo-1)] 414 domains, which bind an also C-terminal peptide in GM130 (Hu et al., 2015).. Similar to 415 its metazoan counterparts, Aspergillus Grh1 contains an N-terminal α -helix and two PDZ 416 domains, in this case followed by ~130 disordered residues (Figure 8B, Figure 8—figure 417 supplement 1A and C). We modelled the Grh1-Bug1 interaction using AlphaFold2. 418 Residues 666-675 of the C-terminal peptide of Bug1 bind to a hydrophobic cleft located 419 between PDZ1 and PDZ2. Bug1 Leu668 and 670 coordinate their side chains with 420 residues from both PDZ domains (e.g. Phe45 and Trp44 in PDZ1 and Trp171 and Val179 421 in PDZ2). A second interaction involves Bug1 C-terminal residues 683-690 fitting within 422 a second groove in PDZ1, such that the four C-terminal residues form a β -strand 423 extending the β -sheet of N-terminal PDZ1 domain (Figure 8B, Figure 8—figure 424 supplement 1B). Further genetic evidence that a network of interactions similar to that 425 acting in yeast and mammalian cells operates in the A. nidulans ER-Golgi interface was 426 obtained by constructing strains with combinations of gene-replaced alleles. These 427 consisted of $uso1 \triangle CTR$ encoding Uso1 lacking the C-terminal region (residues 1-1041) and containing or not E6K G540S, rab1^{ts}, and deletion alleles of the Aspergillus BUG1 428 (AN7680) and *GRH1* (AN11248) genes (Figure 8A). Combining *rab1*^{ts} with *uso1*∆*CTR* 429 430 resulted in a synthetic negative interaction at 30°C akin to that seen with uso1-GFP

431 (lanes 2 and 4 in Figure 8C). That the E6K/G540S double substitution rescued this 432 negative interaction strongly indicates that the CTR cooperates with RAB1 in the 433 recruitment of Uso1 to membranes (Figure 8C, lanes 2, 4, 6 and 8). We note that the 434 control wild-type strain used in these experiments contains a construct completely 435 analogous to the mutant allele, ruling out that the genetic manipulation (for example the 436 introduction, linked to the uso1 locus, of a selection marker, potentially chromatin-437 disruptive) is causative of the observed phenotype. Another trivial explanation that we 438 ruled out by Western-blot analysis was that the deletion off the 62 C-terminal residues in 439 $uso1 \triangle CTR$ resulted in increased degradation, which it did not (Figure 8D).

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441 If interactions involving the CTR of p115 were conserved in fungi, BUG1 (GM130 442 equivalent) and GRH1 (GRASP65) should also show a synthetic negative interaction 443 with rab1^{ts}. Figure 8E (lanes 1,3 and 5) shows that neither bug1 Δ nor grh1 Δ affects 444 growth. However, both deletion alleles showed a strong synthetic negative interaction 445 with *rab1*^{ts} (Figure 8E, lanes 1,3 and 5). Remarkably, the synthetic negative phenotype 446 was rescued by the presence of E6K/G540S substitutions in Uso1, further suggesting 447 that they promote Uso1 recruitment to its locale of action, compensating for the loss of 448 the Bug1-Uso1 CTR interaction. We conclude that interaction involving the CTR of Uso1 449 and the BUG1/GRH1 complex cooperates with RAB1-mediated mechanisms to recruit 450 Uso1 to membranes.

451 The globular head domain (GHD) of Uso1 carrying the double E6K/G540S

452 substitution supports cell viability

453 Uso1 has traditionally been considered the archetype of a coiled-coil tether 454 recruiting ER-derived vesicles to the cis-Golgi. In S. cerevisiae, uso1-1 (Sapperstein et 455 al., 1996), is a C-terminally truncating, conditionally-lethal ts allele whose encoded 456 protein still retains 20% of the coiled-coil region. In contrast, uso1-12 and uso1-13 457 removing the complete coiled-coil region are lethal (Seog et al., 1994). Thus, we asked 458 whether the double E6K/540S substitution bypasses the requirement of the coiled-coil region. To this end, we constructed, by gene replacement, a uso1^{GHD} allele expressing 459 460 a protein truncated immediately after the GHD, lacking residues 660-1103. By 461 heterokaryon rescue, we demonstrated that this allele is lethal (Figure 9A). 462 Unexpectedly, given that the *rab1*^{ts} suppressor substitutions lie outside the coiled-coil 463 region, the equivalent allele containing E6K/G540S sufficed for the fungus to survive at 464 30°C. This result has two key implications: that the coiled-coil of Uso1 is not essential for 465 survival and as the GHD of Uso1 is a monomer in solution, it follows that dimerization of 466 Uso1 is not essential either (see also below).

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468 In view of these unexpected results we wondered whether the structure of the GHD 469 synthesized in bacteria differed from the physiological form in Aspergillus, such that the 470 GHD were a dimer in vivo. To address this possibility, we ran an Aspergillus cell-free 471 extract expressing HA3-tagged Uso1 GHD through a Sepharose column. As control, we 472 ran in parallel a sample of bacterially-expressed His-tagged GHD used in sedimentation 473 velocity experiments. Western blot analysis of the fractions (Figure 9B) demonstrated 474 that both proteins eluted at the same position, corresponding to that expected for a 475 globular protein with the size of the GHD. As controls we ran in the same column a similar 476 pair of proteins corresponding to full length Uso1. The bacterially-expressed and the 477 native Uso1 proteins also co-eluted (Figure 9B), but this time at a position corresponding 478 to a highly elongated dimer, consistent with sedimentation velocity experiments. Thus, 479 in vivo, the GHD, expressed from its own promoter, is a monomer, and it is sufficient to 480 sustain viability if it carries the double mutant substitution that bypasses the requirement 481 for RAB1. These data strongly argue against tethering being the essential physiological 482 role that Uso1 plays.

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484 We next investigated why the E6K/G540S GHD suffices for viability only at 30°C. Often 485 thermo-sensitivity results from protein instability, which is enhanced at high 486 temperatures. Western blot analysis of the allele-replaced strain expressing E6K G540S 487 GHD as the only source of Uso1 revealed that levels of the truncated Uso1 mutant were 488 minuscule relative to the wt or to the equivalent ΔCTR allele (Figure 9C). We reasoned 489 that increasing expression would result in E6K/G540S GHD supporting growth over a 490 wider range of temperatures. Thus, we drove its expression with the promoter of the 491 inulinase inuA gene, which is inducible by the presence of sucrose in the medium and 492 almost completely shut off on glucose (Hernández-González et al., 2018; Peñalva et al., 493 2020). Initially we tested wild-type and E6K/G540S GHD in a uso1+ background. This 494 had no phenotypic consequences despite the fact that western blots confirmed that that 495 the truncated proteins were being overexpressed (Figure 9D; Figure 9E, lanes 1, 3, 5, 7 496 and 10;). Then we proceeded to delete the resident USO1 gene in the wild-type and 497 mutant GHD overexpressing strains. As expected, neither of the resulting pair of strains 498 was able to grow on medium with glucose as the only carbon source (Figure 9E, lanes 499 2 and 6). Notably, the strain expressing E6K/G540S GHD as sole Uso1 source grew 500 essentially as the wild-type and 30°C and, although debilitated, was viable at 37°C, 501 showing a substantial improvement of the growth capacity displayed by the gene-502 replaced mutant (Figure 9, lanes 7,8 and 9). Thus, if expressed at sufficiently high levels, 503 E6K/G540S GHD maintains viability at the optimal growth temperature.

504

505 Unexpectedly, the wild-type GHD also rescued the viability of the $uso1\Delta$ mutant when 506 this was cultured with sucrose as carbon source at 30°C and 37°C. In fact, at 30°C, the 507 uso1 inuAp::GHD strain grew like the wt (Figure 9E, lanes 3 and 4)(note that these 508 experiments were carried out in a RAB1+ background), suggesting that increased 509 binding to a Golgi receptor facilitated by mass action compensated for the loss of the 510 coiled-coil region and associated dimerization. The E6K/G540S GHD would have gained 511 affinity for this receptor, explaining why the doubly substituted GHD suppressed mis-512 localization of Uso1-GFP when RAB1 is compromised, even when its steady-state levels 513 were very low. Forced expression, combined with a potentially increased binding affinity 514 of E6K/G540S GHD to such a hypothetical receptor might be toxic.

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Uso1 is an associate of the early Golgi SNARE machinery, with the double substitution E6K/G540S increasing this association

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518 What is the nature of this hypothetical receptor? To address this question, we screened 519 for interactors of Uso1 among proteins acting at the same functional level (consumption 520 of COPII vesicles by the early Golgi) using a modified version of the S-tag co-521 precipitation approach that we used to characterize of TRAPP complexes (Pinar et al., 522 2019)(Figure 10A). We constructed strains expressing wild-type or mutant Uso1, tagged 523 endogenously with the S-tag and, as negative unrelated control, BapH (an effector of 524 RAB11 acting in late steps of the secretory pathway (Pinar and Peñalva, 2017). Then, 525 derivatives of these three strains co-expressing each of the candidate Uso1 GHD 526 targets, tagged with HA3 (also endogenously), were constructed. The resulting panel 527 (Figure 10B) was screened for HA-tagged proteins co-precipitating more efficiently with 528 the E6K/G540S version of Uso1 than with the wild-type, and satisfying the criterium of 529 not co-purifying with BapH. To this end cell-free extracts of these strains were incubated 530 with S-agarose beads that were recovered by centrifugation. Proteins associating with 531 the S-baits were revealed by anti-HA western blotting.

532

533 That not every protein specifically co-purified with Uso1 baits was demonstrated by the 534 results obtained with the COG component COG2, which did not associate with any of 535 the three baits (Figure 10C, 10). In contrast, β -COP was a promiscuous non-specific 536 interactor pulled down by all three baits (Figure 10C, 11). Notably, the screen identified 537 the Golgi syntaxin Sed5 within the specific Uso1 associates (Figure 10C, 1), an 538 association reported previously by others for both p115 and fungal Uso1 (Allan et al., 539 2000; Sapperstein et al., 1996). That the PM SNARE Sso1 did not interact at all with any

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of the S-baits demonstrated that Uso1 does not bind promiscuously to syntaxins (Figure
10C, 5). Importantly, Sed5 was brought down more efficiently by E6K/G540S Uso1, and
not at all by BapH, even though levels of this unrelated bait, as assessed by silver
staining of pull-downs, were markedly higher than those of either Uso1 version.
Therefore, Sed5 (or its associates) might represent a potential anchor bound by
E6K/G540S Uso1 with increased affinity to compensate for the lack of RAB1-mediated
recruitment.

547

548 This analysis was extended to other members of the SNARE bundle forming in the 549 ER/Golgi interface with the Qa Sed5: the Qb Bos1, the Qc Bet1 and the R-SNARE Sec22 550 (Parlati et al., 2002; Pelham, 1999; Tsui et al., 2001). Sec22 was slightly enriched in the 551 E6K/G540S pull-down relative to wild-type Uso1 (Figure 10C, 4). The results with Bos1 552 and Bet1 were most noteworthy (Figure 10C2 and 3). Both were markedly increased in 553 the E6K/G540S pull-downs, with Bet1 increased most. The AAA ATPase Sec18 554 disassembling cis-SNARE complexes also bound Uso1 and was slightly enriched in the 555 sample of E6K/G540S associates, as was the Uso1-interacting Golgin Bug1, but not its 556 membrane anchor Grh1 (Figure 10, C6-8). We conclude that Uso1 is a component of 557 the SNARE machinery and that this association is augmented by E6K/G540S.

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559 S-tag-coprecipitations in Figure 10 clearly singled out Bet1 (Qc) and Bos1(Qb) as the 560 preys that were most strongly enriched with the mutant E6K/G540S bait relative to wild-561 type, and therefore with the highest probability of being direct interactors bound with 562 greater affinity by the doubly substituted Uso1 mutant.

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564 During these analyses we also addressed the physiological role of golgins, which were 565 expected to be dispensable for growth, given their functional redundancy (Gillingham, 566 2018; Gillingham and Munro, 2016; Muschalik and Munro, 2018). We examined the role 567 of three golgins acting at the early Golgi: The Grh1 (AN11248)/Bug1(AN7680) complex, 568 discussed above, Cov1 (the product of AN0762) and Rud3 (AN10186). Consistent with 569 their roles, AlphaFold2 predicts that they form long coiled-coils carrying C-terminal 570 membrane anchors, characteristics of golgins. Coy1 contains a C-terminal TMD anchor 571 and an adjacent CLASP domain that, according to AlphaFold2, consists of α -helices (Figure 10-figure supplement 1). Rud3 is a dimer consisting of a long coiled-coil with a 572 573 ARF1-binding GRIP domain composed of four short helices, near its C-terminus (Figure 574 10—figure supplemental 1). Ablation of grh1, bug1, coy1 or rud3 did not prevent growth 575 of the corresponding mutants, with only $coy1\Delta$ displaying a minor growth phenotype 576 (Figure 10—figure supplement 2). Next, we deleted the corresponding genes in a rab1 Δ

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uso1^{E6K G540S} and tested whether any of the golgins was required for viability rescue. Grh1 577 578 and Bug1 were partially required, as their absence precluded rescue at 42° (Figure 10-579 figure supplement 2). In contrast, RUD3 was not required. With regard to COY1, the fact that combining $coy1\Delta$ and $rab1\Delta$ uso1^{E6K G540S} results in lethality at 30°C, 37°C and 42°C 580 581 precluded conclusions on the role of COY1 in suppression. However, we can conclude 582 that proper assembly of A. nidulans Golgi cisternae is supported by a redundant set of 583 tethers (Behnia et al., 2007), of which in Coy1 appears to be the least redundant of those 584 tested. Coy1 has been implicated in retrograde traffic within the Golgi itself (Anderson et 585 al., 2017)

586

587 Golgi SNAREs bind directly to the Uso1 GHD; effects of Uso1 E6K/G540S

588 Work by others implicated Uso1 in the assembly of the early Golgi SNARE bundle 589 (Sapperstein et al., 1996). Thus, prompted by co-association experiments, we predicted 590 that Uso1 would bind directly Bet1, Bos1 and perhaps other SNAREs implicated in the 591 biogenesis of the early Golgi. We anticipated that binding to Bet1 and Bos1 would be 592 insufficient to recruit Uso1 to membranes in the absence of RAB1, but that once 593 reinforced by E6K/G540S, Uso1 would not require RAB1 for its recruitment. To test this 594 possibility, we searched for direct and E6K/G540S-enhanced interactions between the 595 GHD and SNAREs with pull down assays carried out with purified SNARE-GST fusion 596 proteins as baits and Uso1-His6 constructs as preys.

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598 Full-length Uso1 bound, weakly, to the Sec22 R-SNARE and to the Qb Bos1, and very 599 efficiently [with *circa* 70% of the prey being pulled down (Figure 11A, B)] to the Qc Bet1. 600 In contrast, Uso1 did not bind the Qa syntaxins tested, Sed5 and Sso1 (Figure 11A). The 601 absence of interaction between Sed5 and Uso1, be it the wild-type or the E6K/G540S 602 mutant version, strongly indicated that the association detected with S-tag pull-downs 603 between Uso1 and Sed5 is bridged by other protein(s). This absence of binding cannot 604 be attributed to Sed5-GST being incompetent for binding because Sed5-GST was 605 competent in pulling-down highly efficiently its cognate SM protein Sly1, an interaction 606 that did not occur with Sso1-GST (Figure 11C). Notably, the presence of the E6K/G540S 607 double substitution in Uso1 (indicated with ** for simplicity on Fig 11) increased five times 608 the amount of protein retained by the Qb Bos1 bait (Figure 11A and B), whereas 609 interaction with Bet1 did not change (Figure 11A and B). The double substitution in Uso1 610 did not promote interaction with Sed5 either. Thus, under normal circumstances Uso1 is 611 able to bind directly to three of the four SNAREs in the ER/Golgi interface, with binding 612 to Bet1 being the strongest. The double E6K/G540S substitution increases binding to

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Bos1 very markedly and specifically, bringing it to up to the levels of Bet1 without affecting, for example, binding to Bet1 or Sec22. Consistently, the GHD is sufficient to mediate interaction with Bet1 and Bos1, as well as, if E6K G540S-substituted, the increased binding of Bos1 to Uso1 (Figure 11D and E). The GHD did not interact with Sec22, suggesting either that this R-SNARE is recruited by other parts of the protein or that binding is dimerization dependent.

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620 Besides the Sed5/Bos1/Bet1/Sec22 combination (), across cisternal maturation Sed5 621 forms SNARE bundles in Golgi compartments located downstream of Uso1 domains 622 (Pelham, 1999). In fungi, membrane fusion in the medial Golgi involves the Sed5 623 partners Gos1 (Qb) and Sft1 (Qc) substituting for Bos1 and Bet1, respectively, but 624 neither Gos1 nor Sft1 bound wild-type or E6K/G540 Uso1 GHD (Figure 11F), 625 demonstrating that interaction of Uso1 with Bos1 and Bet1 is highly specific. Therefore, 626 it seems fair to conclude that increased binding for a SNARE receptor underlies the 627 mechanism by which mutant Uso1 bypasses the need for RAB1 in the ER/Golgi 628 interface.

629

630 In summary, Uso1 is an essential protein acting in the ER/Golgi interface, and we report 631 here several important findings. We show that (i) the Golgi GTPase RAB1, which is 632 essential for viability, becomes dispensable if there is an alternative method to recruit 633 Uso1 to Golgi membranes; (ii) the coiled-coil region of Uso1 is dispensable to sustain 634 viability, implying that the tethering role of the protein is not essential either, consistent 635 with the redundant roles of other Golgin tethers; (iii) the Uso1 GHD is essential for 636 viability; (iv) the Uso1 GHD monomer, if present at suitably high levels, is sufficient to 637 maintain viability; (v) that the Uso1 GHD is a direct and specific binder of the Qb SNARE 638 Bos1, and a strong binder of the Qc Bet1; (vi) the mutation bypassing the need for RAB1 639 markedly increases the affinity of the GHD for Bos1, indicating that rab1 viability rescue 640 by E6K/G540S Uso1 occurs because SNARE anchoring provides an alternative mode 641 of recruitment to Golgi membranes to that provided physiologically by RAB1.

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643 Mechanistic insights guided by AlphaFold2 predictions

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To gain further insight into the mechanisms by which the double E6K/G540S bypasses
RAB1 we exploited AlphaFold2 to model the Uso1^{GHD} domain alone, or together with
each of the individual SNAREs, with both Bos1 and Bet1 simultaneously, and with RAB1.

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649 Consistent with biochemical data, in the predicted model Bet1 and Bos1 interact with a medial and a C-terminal region, respectively, of Uso1^{GHD}. (Figure 12; Figure 12-650 651 supplement 1 and —supplement 2). Bet1 docks against a region of the GHD that is not 652 affected by the mutations. In contrast, Gly540 and its environs dock against the N-653 terminal, triple α -helical Habc domain of Bos1. In all likelihood Gly540Ser mediates the 654 increased binding of mutant Uso1GHD to Bos1. AlphaFold2 predicts that Uso1GHD 655 interacts with Bos1 through a surface composed by the N-terminal part of the first Bos1 656 Habc α -helix and the loop between α -helices 2 and 3 (Figure 12A and B). The binding 657 surface in Uso1^{GHD} involves the second α -helices of the ARM10 and ARM11 repeats, 658 (α -helices 26 and 29), and the loop connecting the first two α -helices of ARM10 (Figure 659 12B). In AlphaFold2 models, Uso1 Gly504 (wild-type) is located at the beginning of Uso1^{GHD} α -helix 29. at the heart of the interaction surface, contributing to the Uso1-Bos1 660 661 interaction by coordinating the amide group of the Uso1 backbone with the Bos1 Glu58 662 carboxylate to create a hydrogen bond (Figure 12C). According to the most confident 663 prediction, Gly504Ser results in the hydroxymethyl side chain protruding into a small 664 pocket rimmed by the side chains of Bos1 Leu59 and Ile60, such that the Uso1 Ser504 665 hydroxyl group hydrogen-bonds the amide group of Bos1 Glu58 (Figure 12C). Besides creating a new hydrogen bond, Gly540Ser would increase the surface of interaction 666 slightly, from ~825.2 Å² to ~831.1 Å², implying that these alterations, together with a 667 668 minor shift in the environs of Gly540 detected by models (which might facilitate additional 669 interactions), would strengthen the binding of the Uso1^{GHD} to the Bos1 surface.

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671 The mechanism by which Glu6Lys contributes to increase the recruitment of Uso1 to 672 Golgi membranes appears to be different. This glutamate is located in a region with low 673 pLDDT score which shows different conformations depending on the model (Figure 2D, 674 boxed), suggesting that this region is difficult to predict due to flexibility. However, 675 models concur in the prediction of an amphipatic N-terminal α -helix containing Glu6, 676 whose substitution by Lvs (as in E6K) reinforces the positive charge of this α -helix, which 677 would be inserted into the vesicle membrane, potentially contributing to Uso1 recruitment 678 to COPII vesicles

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In the case of the strong Uso1^{GHD}-Bet1 interaction (Figure 12—figure supplement 1), the
 N-terminal region of Bet1 consisting of *circa* 80 amino acids is disordered, and the
 pLDDT score of the different models is understandably low. However, all structural
 models depicting Bet1 interacting with Uso1^{GHD} (e.g. in the context of the whole SNARE

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684 complex, Bet1 alone or the isolated Bet1 N-terminal) consistently show a region where 685 the pLDDT is higher. This region forms a kink in this N-terminal part of the Bet1 that 686 protrudes into the surface created by the α -helices 13 and 16 of ARM4 and ARM5 (Figure 687 12-figure supplement 1). Surface representations show that this section of the Bet1 688 polypeptide covers ~ 1300 Å² of the GHD, docking against the same side of the 689 boomerang-shaped solenoid as Bos1, which is consistent with the orientation that these 690 SNAREs should take during the formation of the SNARE pin (Figure 12-figure 691 supplement 2)

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693 We next modelled RAB1 binding to the GHD in the absence and presence of Uso1 694 binders Bos1 and Bet1. AlphaFold2 predicts that the GHD interacts with RAB1 through 695 a binding surface formed by the ARM3-5. repeats. On the other hand, the interactive 696 region of RAB1 conforms to canons, as Uso1 α -helices 8 and 9 (ARM3) interact with the 697 Switch II region while α -helices 11 (ARM-4) and 14 (ARM-5) interact with the Switch I) 698 (Figure 12-figure supplement 2, A and B). Importantly, the model indicates that within 699 the Uso1-GHD jai alai basket, RAB1 binds to the opposite (convex) side of the Bet1 700 interacting area at the concave side, and away from the C-terminal helix where the Bos1 701 Habc domain predictably binds, which would allow Uso1 to bind these three interactors 702 simultaneously (Figure 12-figure supplement 2C). In addition, the predicted models 703 supported two highly suggestive but as yet speculative implications. One is that the 704 position of the RAB1 hypervariable domain, which anchors the GTPase to the membrane 705 through is prenylated C-terminus, is compatible whith the hypothetical membrane 706 binding of the N-terminal Uso1 amphipatic helix (Figure 12-figure supplement 2D); The 707 second is that Uso1-RAB1 would be in an orientation that facilitates the docking of RAB1-708 loaded ER-derived vesicles with an acceptor membrane where SNARE zippering occurs 709 (see discussion); this orientation implies that the donor membrane (the position of the N-710 terminal Uso1 helix) and the acceptor membrane (the TMDs of the SNAREs) is 711 compatible with the Uso1 CTR contributing to tethering through interaction with 712 Bug1/Grh1.

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RAB1 regulates transport at the ER/Golgi interface. Using an unbiased forward genetic
screen to identify subordinated genes accounting for its essential role, we isolated two
extragenic mutations resulting in single-residue substitutions in the RAB1 effector Uso1,

Discussion

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719 usually regarded as a tether. The single-residue substitutions lie at opposite ends of the 720 jai alai basket-shaped GHD; both are individually able to rescue viability of rab1 d strains 721 at 30°C and, when combined, even at 42°C. Subcellular localization experiments hinted 722 at the mechanism by which the double mutation rescues $rab1\Delta$ lethality. Uso1 plays its 723 physiological role on an early Golgi compartment, where it largely colocalizes with RAB1, 724 with relocation to the cytosol in a RAB1-deficient background. Under normal 725 circumstances, the Uso1 CTR acts in concert with RAB1 to recruit the protein to the 726 Golgi. Genetic evidence (Figure 8) showed that this contribution requires the Golgi-727 localized tether composed of the membrane anchor Grh1 and its associated golgin 728 BUG1 (Behnia et al., 2007), homologues of human GRASP65 and GM130, respectively. 729 In the absence of RAB1, engagement of the CTR with BUG1 is insufficient to stabilize 730 wild-type Uso1 on membranes. However, the double E6K/G540S substitution relocalizes 731 Uso1 to Golgi structures, suggesting that mutant Uso1 GHD had gained affinity for 732 another element, thereby compensating for the loss of RAB1.

733

734 By S-tag co-precipitation experiments we identified proteins associating with wild-type 735 and E6K/G540S Uso1 baits. Prominent among these were the four SNAREs, Sed5, 736 Bos1, Bet1 and Sec22, mediating fusion events at the ER/ Golgi interface, and the 737 SNARE regulator Sec18, indicating that Uso1 is a component of the SNARE fusion 738 machinery. This was suggested by previous studies with S. cerevisiae showing that 739 overexpression of Bet1, Bos1 and Sec22 suppresses uso1-1 and that of Sec22 and Bet1 740 weakly suppresses $uso1\Delta$ (Sapperstein et al., 1996). Moreover, in mammalian cells 741 crosslinking studies with p115 identified Sed5, membrin (Bos1) and mBet1 as its weak 742 interactors (Allan et al., 2000), suggesting that contributing to the ER/Golgi SNARE 743 machinery is a conserved feature of Uso1/p115 family members. Of the above four A. 744 nidulans SNAREs, Bet1 and Bos1, which co-precipitated weakly with the wild-type, were 745 dramatically enriched with the mutant. Pull-down assays showed that full-length 746 Aspergillus Uso1 interacts weakly with Sec22 and strongly with Bet1, irrespective of 747 whether the bait was wild-type or E6K/G540S, indicating that they are direct 748 physiological interactors. In sharp contrast, the interaction of E6K/G540S Uso1 with the 749 Qb Bos1 was markedly augmented, strongly suggesting that this increase contributes 750 substantially to bypass RAB1. Both the high levels of binding to Bet1, and the marked 751 increase in binding to Bos1 resulting from E6K/G540S were tracked down to the GHD, 752 which is sufficient to bind these SNAREs.

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754 AlphaFold2 models predicting the regions of interaction of the GHD with Bos1 and Bet1 755 showed that, of the two Uso1 residue substitutions, only Gly540Ser maps to the region 756 mediating the interaction with Bos1, whereas neither affected the interacting region with 757 Bet1, agreeing with GST pull-downs. Interaction with this SNARE is predicted to involve 758 a large surface, consistent with the strong "constitutive" binding of Bet1 with the GHD. 759 AlphaFold2 also predicted that RAB1 binds to the convex face of the GHD solenoid, in a 760 position that would permit the simultaneous binding of Qb and Qc SNARES and the 761 GTPase. In addition, AlphaFold2 also detected a previously unnoticed amphipathic α-762 helix in the N-terminal region of Uso1. The Glu6Lys substitution falls within this helix, 763 making its global positive charge even greater, strongly suggesting that this substitution 764 increases Uso1 GHD binding to membranes. That the two residue substitutions act by 765 different mechanisms is coherent with their showing additivity to suppress RAB1 deficit. 766

767 The Uso1/p115 family has been implicated in the regulation of SNARE complexes (Allan 768 et al., 2000). (Shorter et al., 2002) first reported that p115 binds to SNAREs and 769 proposed that they would 'catalyze' the formation of the ER/Golgi SNARE bundle. The 770 direct interaction of p115 with unassembled mBet1 and Sec22 agrees with this role 771 (Wang et al., 2014). However, there are significant gaps in this model: Uso1 772 overexpression suppresses a partial deficit of ypt1/RAB1, but not ypt1 Δ , whereas Ypt1 773 overexpression suppresses $uso1\Delta$, indicating that Uso1 acts and in concert or upstream 774 of RAB1. [Of note, overexpression of Uso1 also suppresses bet3-1, a ts allele 775 inactivating the TRAPPIII complex, which is the RAB1 GEF (Galindo et al., 2021; Jiang 776 et al., 1998; Pinar et al., 2019; Riedel et al., 2017; Thomas et al., 2018)]. The grid of 777 reported genetic interactions strongly indicates that RAB1 regulates SNAREs (Brandon 778 et al., 2006; Lupashin and Waters, 1997; Sapperstein et al., 1996). Notably, our viability 779 rescue experiments show that E6K/G540S bypasses this non-canonical, yet essential 780 role of RAB1. As the double mutant substitution increases the recruitment of USO1 to 781 the SNARE machinery and as E6K/G540S suppresses rab10, the most parsimonic 782 interpretation of the data is that the role of RAB1 cooperating to recruit Uso1 to the 783 SNARE complexes, a cooperation that is no longer needed when recruitment is ensured 784 by other means. That p115 regulates SNAREs' assembly was proposed by (Shorter et 785 al., 2002). However, a fundamental difference with our conclusions is that they attributed 786 this role to the CCD, whereas we identify here the GHD as the positively-acting player.

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We also note that our experiments provide a mechanistic interpretation as to why overexpression of Bet1, Bos1, Sec22 and Ypt1/RAB1 rescue the viability of *uso1-1*

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removing a substantial portion of the yeast Uso1 CCD (Sapperstein et al., 1996; Seog
et al., 1994). A similar *Aspergillus* allele results in marked protein instability (Figure 9).
Thus, in all likelihood, suppression by overexpression of known direct interactors
involves stabilization of the *uso1-1* product.

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In summary, our work firmly establishes that the essential role of Uso1 resides not in its CCD domain tethering donor and acceptor membranes, but in the GHD. When expressed at sufficient levels this domain is capable of fully complementing *uso1Δ*, which is definitive evidence that neither the tethering function of Uso1 nor dimerization (the GHD is monomeric) is required for the protein to play its essential role. As the GHD binds two of the four SNAREs of the bundle, the simplest interpretation is that the essential role of Uso1 is regulating SNAREs.

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803 Ideas and Speculation

804 While the molecular details of this regulation will be addressed in future, it is tempting to 805 speculate that the GHD contributes, with Sly1, to orientate SNAREs to form a productive 806 bundle, acting as chaperones, similarly to the HOPS SM component Vps33, which 807 appears to align SNAREs in a pre-zippering stage, facilitating their assembly (Baker and 808 Hughson, 2016; Baker et al., 2015; Ren et al., 2009; Yu and Hughson, 2010; Zhang and 809 Hughson, 2021; Zhang and Yang, 2020). A second speculative interpretation is that 810 tethering occurs in two steps, with golgins acting at long distances, approximating 811 vesicles to the vicinity of SNAREs. Then SNAREs would engage RAB1-Uso1 to serve 812 as short-range tether preceding the zippering up of membranes. The > 700 Å-long Uso1 813 CCD would cooperate with Bug1 in the first step, whereas the Uso1 GHD would 814 cooperate with RAB1 and Bet1/Bos1 in the second, exploiting the fact that GHD binders 815 use surfaces located at opposite sides of the α -solenoid (Figure 12-figure supplement 816 2). Such arrangement implies that RAB1 C-terminal isoprenoids inserting into the donor 817 vesicle membrane would be circa 220 Å apart from the C-termini of the SNAREs inserted 818 in the acceptor membrane. Two-step tethering might impose one additional level of 819 specificity, preventing unproductive fusion events mediated by other SNARES circulating 820 through the ER/ interface and directing, by way of Uso1 interactions, incoming vesicles 821 to fusion-competent areas enriched in target SNAREs (Bentley et al., 2006).

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Materials and Methods

826 Aspergillus techniques

Standard *A. nidulans* media were used for growth tests, strain maintenance and conidiospore harvesting (Cove, 1966). GFP-, HA3- and S-tagged alleles were introduced by homologous recombination-mediated gene replacement, using transformation (Tilburn et al., 1983) of recipient $nkuA\Delta$ strains deficient in the nonhomologous end joining pathway (Nayak et al., 2005). Complete strain genotypes are listed in supplemental table I. These alleles were usually mobilized into the different genetic backgrounds by meiotic recombination (Todd et al., 2007).

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835 Null mutant strains were constructed by transformation-mediated gene replacement. 836 using as donor DNA cassettes made by fusion PCR (primers detailed in supplemental 837 table II) carrying appropriate selectable markers (Szewczyk et al., 2006). Integration 838 events were confirmed by PCR with external primers. When allele combinations were 839 expected to be synthetically lethal or severely debilitating, the corresponding strains 840 were constructed by sequential transformation, but the second such manipulation was 841 always carried out using pyrG^{Af} as selective marker, which favors the formation of 842 heterokaryons in which untransformed nuclei supported growth (Osmani et al., 1988). 843 Conidiospores, in which single nuclei had segregated (i.e., homokaryotic nuclei), were 844 scrapped and streaked onto plates carrying doubly-selective medium. Absence of 845 growth or appearance of microcolonies, combined with a positive PCR diagnostic of 846 heterokaryosis of the primary transformants, was taken as indication of lethality. 847 Whenever possible, colony PCR of microcolonies was always used to genotype the 848 desired genetic intervention

849

850 The following proteins were C- or N-terminally tagged endogenously, using cassettes constructed by fusion PCR (Nayak et al., 2005; Szewczyk et al., 2006): Uso1-GFP and 851 Uso1^{E6K/G540S}-GFP, Uso1-HA3 and Uso1^{E6K/G540S}-HA3; Uso1-S and Uso1^{E6K/G540S}-S; 852 853 BapH-S (Pinar and Peñalva, 2017), Sec13-mCherry (Bravo-Plaza et al., 2019; 854 Hernández-González et al., 2019), Gea1-mCherry and Sec7-mCherry (Arst et al., 2014), 855 mCherry-Sed5 (Pantazopoulou and Peñalva, 2011), mCherry-RAB1 (Pinar et al., 2013), 856 HA3-Sed5, HA3-Bet1, HA3-Bos1, HA3-Sec22, Sec18-HA3, Grh1-HA3, Bug1-HA3, 857 Cov1-HA3, COG2-HA3, and β -COP-HA3.

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Antibodies for western blotting			
Antibody	dilution	Origin	Reference
Primary			
α-HA	1:1000	rat	3F10 clone, Roche
α-His6x tag	1:10 000	mouse	#631212, Clontech
α-tubulin	1:5000	mouse	DM1A clone, Sigma
α-Uso1	1:1000	rabbit	Polyclonal antiserum, Davids Biotechnologie
Secondary (HRP-conjugated)			
α-rat IgG	1:4000	goat	#3010-05, Southern Biotech
α-mouse IgG	1:5000	goat	#A9044, Sigma
α-rabbit IgG	1:2000	donkey	#NA934, Amersham

860 Antibodies for western blotting

Antiserum against Uso1 was raised in rabbits by Davids Biotechnology. Animals were immunized with the Uso1 GHD (residues 1-659), tagged with His6x. Recombinant expression in *E. coli* and Ni²⁺ affinity purification is described below. Target antibodies were purified from raw antiserum by affinity chromatography through Hi-Trap NHS columns (#17-0716-01, Cytiva) charged with Uso1 antigen following the manufacturer's instructions. Affinity-bound antibodies were eluted with 100 mM glycine (pH 3.0), then neutralized with 2 M Tris to a pH of 7.5 and stored at -20°C.

868 inuA promoter-driven expression of Uso1 GHD in Aspergillus

869 The Uso1 GHD was expressed in a sucrose-inducible manner from an in locus 870 replacement of the inuA gene (AN11778) ORF encoding inulinase by the GHD coding 871 sequence, such that its expression was driven by the *inuA* promoter, which is induced 872 on sucrose and non-induced on glucose (Hernández-González et al., 2018). The gene 873 replacement cassette was assembled through fusion PCR of 4 different elements, listed 874 here from 5' to 3': (1) inuA promoter, (2) cDNA sequence encoding Uso1 wt or mutant 875 E6K/G540S GHD (residues 1 – 659), (3) Aspergillus fumigatus riboB gene as selection 876 marker and (4) inuA gene 3'-flanking region. A pyrG89, nkuAA::bar, riboB2 A. nidulans 877 strain was transformed with this cassette, replacing the inuA gene. The resulting strain 878 was subsequently transformed with a $uso1\Delta$ deletion cassette (A. fumigatus pyrG as 879 selection marker) to ablate endogenous of Uso1, such that the GHD was the only moiety 880 of Uso1 present.

- 881 Plasmids for protein expression in E. coli
- 882
- 883 (I) His6x-tagging constructs

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- 884 pET21b-Uso1-His6x and pET21b-Uso1(E6K/G540S)-His6x: cDNA encoding full length
- Uso1 (residues 1 -1103) was cloned into a pET21b *Ndel/Not*l linearized vector.
- 886 pET21b-Uso1ΔCTR-His6x and pET21b-Uso1(E6K/G540S)ΔCTR-His6x: lacking the C-
- 887 Terminal Region of the Coiled Coil Domain (residues 1 1040)
- 888 pET21b-Uso1GHD-His6x and pET21b-Uso1(E6K/G540S)GHD-His6x: cDNA encoding
- the Globular Head Domain of Uso1 (residues 1 659) was cloned as a *Ndel/Xhol* insert
- 890 into a pET21b *Ndel/Xhol* linearized vector.
- 891 <u>pET21b-Uso1 CCD-His6x</u>: cDNA encoding Uso1 Coiled Coil Domain (residues 660-
- 892 1103) was cloned as a *Ndel/Notl* insert into a pET21b *Ndel/Notl* linearized vector.
- 893 (II) TNT[®] expression plasmids

894 <u>pSP64-Sly1-HA</u>: this plasmid carries cDNA encoding full length Sly1 (AN2518) C895 terminally tagged with a HA3x epitope, cloned as an *Nsil/Sacl* insert into *Pstl/Sacl*896 pSP64(PolyA) vector

897 (III) GST-tagging constructs

- 898 <u>pET21b-Sed5-GST:</u> cDNA encoding Sed5/AN9526 cytoplasmic domain (residues 1–
 899 322) C-terminally tagged with GST, was cloned as a *Ndel/Sal*I insert into a pET21b
 900 *Ndel/XhoI* linearized vector.
- 901 <u>pET21b-Bos1-GST:</u> cDNA encoding Bos1/AN11900 cytoplasmic domain (residues 1–
 902 219) C-terminally tagged with GST, was cloned as a *Ndel/Sal*I insert into a pET21b
 903 *Ndel/XhoI* linearized vector.
- 904 <u>pET21b-Bet1-GST:</u> cDNA encoding Bos1/AN5127 cytoplasmic domain (residues 1–71)
 905 C-terminally tagged with GST, was cloned as a *Ndel/Sal* insert into a pET21b *Ndel/Xhol*906 linearized vector.
- 907 <u>pET21b-Sec22-GST:</u> cDNA encoding Sec22/ ASPND00903 cytoplasmic domain
 908 (residues 1–198) C-terminally tagged with GST, was cloned as a *Nhel/Sal* insert into a
 909 pET21b *Nhel/Xhol* linearized vector.
- 910 <u>pET21b-Sso1-GST:</u> cDNA encoding Sso1/AN3416 cytoplasmic domain (residues 1–
 911 271) C-terminally tagged with GST, was cloned as a *Ndel/Sacl* insert into a pET21b
 912 *Ndel/Sacl* linearized vector.
- 913 <u>pET21b-Gos1-GST:</u> cDNA encoding Gos1/AN1229 cytoplasmic domain (residues 1–
 914 208) C-terminally tagged with GST, was cloned as a *Ndel/Sal* insert into a pET21b
 915 *Ndel/Xhol* linearized vector.

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916 <u>pET21b-Sft1-GST:</u> cDNA encoding Sft1/AN10508 cytoplasmic domain (residues 1–73)

917 C-terminally tagged with GST, was cloned as a *Ndel/Sal* insert into a pET21b *Ndel/Xhol*918 linearized vector.

919 **Co-precipitation experiments with total cell extracts**

920 Preparation of Aspergillus total cell extracts was done as described, with minor 921 modifications (Pinar et al., 2019) (Pinar et al., 2019). 70 mg of lyophilized mycelium were 922 ground with a ceramic bead in a Fast Prep (settings: 20 sec, power 4). The resulting fine 923 powder was resuspended in 1.5 ml of extraction buffer [25 mM HEPES-KOH (pH 7.5), 924 200 mM KCl, 4 mM EDTA, 1% (v/v) IGEPAL CA-630 (NP-40 substitute, #I8896, Sigma), 925 1 mM DTT, 2 µM MG-132 proteasome inhibitor (#S2619, SelleckChem) and cOmplete[®] 926 ULTRA EDTA-free inhibitor cocktail (#5892953001, Roche). Approximately 0.1 ml of 0.6 927 mm glass beads were added and thoroughly mixed. This suspension was homogenized 928 with a 10 sec pulse at the Fast Prep (power 6) followed by a 10 min incubation at 4°C. 929 This homogenization step was repeated two times before clarifying the extract by 930 centrifugation at 4°C and 15,000 \times g in a microcentrifuge. Total protein concentration of 931 the extracts was determined by Bradford. Bovine Serum-Albumin BSA was then added 932 as a blocking agent to the cell extract (final concentration 1% (w/v)). Binding reactions 933 were carried out in 0.8 ml Pierce centrifuge columns (#89869, ThermoFisher): 9 mg of 934 protein were mixed with 20 µL of S-protein Agarose beads (#69704, Novagen), that had 935 been previously washed in extraction buffer with 1% (w/v) BSA. This buffer was also 936 added to complete final reaction volume of 0.6 ml. The mix was incubated for 3 h at 4°C 937 in a rotating wheel. Columns were then opened at the bottom and gently centrifuged to 938 remove the supernatant and collect the protein-bound beads. These were resuspended 939 in extraction buffer without inhibitors and incubated in rotation for 10 min at 4°C, followed 940 by two more washing steps in extraction buffer without detergent and inhibitors. To elute 941 proteins bound to the beads, 30 µl of Laemmli loading buffer [62.5 mM Tris-HCI (pH 6.0), 942 6 M urea, 2% (w/v) SDS and 5% (v/v) β -mercaptoethanol] were added and the columns 943 incubated at 90°C for 2 min. The columns were centrifuged to collect the eluate, of which 944 a 40% of the final volume were resolved in a SDS-polyacrylamide gel and then 945 transferred to nitrocellulose for α -HA (#3F10, Roche) western blotting.

946 Purification of Uso1 constructs tagged with His6x

Full-length His6-tagged Uso1, Uso1 Δ CTR, Uso1 GTD and Uso1 CCD constructs, wild type and mutant versions, were expressed in *E. coli* BL21(DE3) cells harboring pET21b-His6 derivatives and pRIL. Bacteria were cultured at 37°C in LB medium containing ampicillin and chloramphenicol until reaching an OD_{600nm} of 0.6. Then, IPTG was added to a final concentration of 0.1 mM. Cultures were shifted to 15°C and incubated for 20 h.

952 Bacterial cells were collected by centrifugation and pellets stored at -80°C. For 953 purification, frozen pellets were thawed in ice and resuspended in ice-cold bacterial cell 954 lysis buffer [20 mM sodium phosphate buffer, pH 7.4, 500 mM KCl, 30 mM imidazole, 955 5% (v/v) glycerol, 1 mM β -mercaptoethanol, 1 mM MgCl₂, 0.2 mg/ml lysozyme and 1 µg/ml of DNAse I and cOmplete[®] protease inhibitor cocktail (#11873580001, Sigma)]. 956 957 This cell suspension was mechanically lysed in a French press (1500 kg/cm²) and the 958 resulting lysate was centrifuged at 10,000 \times g and 4°C for 20 min to remove the cell 959 debris. The supernatant was then transferred to polycarbonate tubes and centrifuged at 960 $100,000 \times g$ and 4°C for 1 h in a XL-90 ultracentrifuge (Beckman Coulter) . 50 ml of 961 cleared lysate were incubated with 400 µL of Ni-Sepharose High Performance beads 962 (#17526801, Cytiva) for 2 h at 4°C. After this step, His-tagged protein-bound beads were 963 pelleted at low-speed centrifugation and washed three times in lysis buffer [20 mM sodium phosphate buffer, pH 7.4, 500 mM KCl, 5% (v/v) glycerol, 1 mM β-964 965 mercaptoethanol] with increasing concentrations of imidazole. Finally, Ni2+bound His6 966 proteins were eluted 0.5 M imidazole buffer. 5 ml of eluted protein were loaded onto a 967 HiLoad 16/600 Superdex 200 column (Cytiva) and run at 1 ml/min flow rate on an AKTA 968 HPLC system, using phosphate buffered saline PBS containing 5% (v/v) glycerol and 1 969 mM β -mercaptoethanol. Fractions containing protein were pooled, analyzed for purity by 970 SDS-PAGE followed by Coomassie staining, and finally quantified on a UV-Vis 971 spectrophotometer before being stored at -80°C.

972 Purification of SNARE constructs tagged with GST

973 cDNAs encoding the cytosolic domains of SNAREs fused to a C-terminal GST were 974 cloned into pET21b. Bacterial cultures and protein expression conditions were as 975 described above for Uso1-His6 constructs. Frozen pellets were thawed in ice and 976 resuspended in chilled bacterial cell lysis buffer [25 mM Tris-HCl (pH 7.4), 300 mM KCl, 977 5 mM MgCl₂, 1 mM DTT, 0.5 mg/ml lysozyme, 1 µg/ml of DNAse I and cOmplete[®] 978 protease inhibitor cocktail (#11873580001, Sigma)]. This cell suspension was incubated 979 for 30 min in ice before being mechanically lysed in a French press (1500 kg/cm²). The 980 lysate was incubated for a further 30 min on ice and centrifuged at 20,000 \times g and 4°C 981 for 30 min. After adding 10 mM EDTA to the clarified supernatant to stop DNAse I activity, 982 it was transferred to a 50 ml tube, mixed with 500 µL of glutathione Sepharose beads 4B 983 (#17075601, Cytiva) and rotated for 2 h at 4°C. After incubation, SNARE-GST-bound 984 beads were pelleted by gentle centrifugation and washed three times for 10 min at 4°C 985 in 25 mM Tris-HCI (pH 7.4), 500 mM KCI, 5 mM EDTA, 1 mM DTT and subsequently 986 transferred to a 0.8 ml Pierce column. Beads were washed 6 times (10 min at RT) in 200 987 µL of elution buffer [50 mM Tris-HCI (pH 8.0), 200 mM KCI, 10 mM glutathione and 1

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988 mM DTT]. These fractions were collected and pooled (~1 ml), then buffer-exchanged to
989 storage buffer (PBS, 5% (v/v) glycerol and 0.1 mM DTT) in a PD MidiTrap G-25 column.
990 Protein concentration and purity was assessed by spectrophotometry and SDS-PAGE
991 followed by Coomassie staining. Protein stocks were kept frozen at -80°C.

992 SNARE-GST pull-downs with purified Uso1-His6 constructs (Uso1, Uso1 GHD)

993 Binding reactions were performed in 0.8 ml Pierce centrifuge columns. 75 µg of purified 994 SNARE-GST were mixed with 15 µL of glutathione Sepharose 4B beads and storage 995 buffer to a final volume of 0.3 ml. Columns were rotated at 4°C for 2 h before the 996 supernatant was removed after low speed centrifugation. Subsequently, His6 preys were 997 added to a final concentration of 0.2 µM in a total volume of 0.4 ml of pull-down binding 998 buffer [25 mM HEPES-KOH (pH 7.5), 150 mM NaCl, 10% (v/v) glycerol, 0.1% (v/v) Triton 999 X-100 and 0.1 mM DTT]. Columns were rotated overnight at 4°C. Beads were collected 1000 by gentle centrifugation and washed three times for 10 min with ice-cold binding buffer, 1001 before eluting bound proteins with 30 µL of Laemmli loading buffer pre-heated at 90°C. 1002 0.5 % of the samples (eluted material or flow-through) were run onto 8% SDS-1003 polyacrylamide gels that were transferred to nitrocellulose membranes which were 1004 reacted with α -His tag antibody (#631212, Clontech). Quantitation of band intensities 1005 was done with ImageLab software (BioRad). In parallel, 4% of the elution sample volume 1006 was loaded onto 10% SDS-polyacrylamide gel and stained with coomassie dye 1007 (BlueSafe, NZY) to confirm recovery of SNARE-GST baits.

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1009 Pull-down of TNT[®]-expressed Sly1-HA3

Sly1-HA3 was synthesized with the TNT[®] SP6 Quick Coupled Transcription/Translation 1010 1011 system (#L2080, Promega), according to the instructions of the manufacturer. The 1012 reaction was primed with 1 µg of pSP64::Sly1-HA3 cDNA. 10 µL of the resulting mix 1013 were combined with 15 µL of glutathione-Sepharose beads, previously loaded with 1014 SNARE-GST baits as described above, in 0.4 ml of pull-down binding buffer, using 0.8 1015 ml Pierce columns that were rotated overnight at 4°C before beads and flow-through 1016 were recovered after gentle centrifugation. Beads were washed three times for 10 min 1017 at 4°C in pull-down binding buffer before eluting bound material with 30 µL of Laemmli 1018 loading buffer for 2 min at 90°C. 20% of the elution sample volume was analyzed by 1019 western blotting with α-HA tag antibody (#3F10, Roche) for Sly1-HA immunodetection.

1020 Size exclusion chromatography of HA-tagged cell extracts

1021 Gel filtration experiments were performed as described (Bravo-Plaza et al., 2019).

1022 Briefly, 200 µL of cell extract were loaded onto a Superose 6 10/300 column (Pharmacia)

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1023 equilibrated with running buffer [25 mM Tris-HCl (pH 7.5), 600 mM KCl, 4 mM EDTA, 1 1024 mM DTT]. Fractions of 0.5 ml were collected, from which 80 μ L were mixed with 40 μ L 1025 of Laemmli loading buffer and denatured at 90°C. 25 μ L of these samples were resolved 1026 by SDS-PAGE and analyzed by western blotting with α -HA3 tag antibody (#3F10, 1027 Roche) for western blotting. Sizing standards were myoglobin (17 kDa), BSA (67 kDa), 1028 aldolase (158 kDa) ferritin (449 kDa), thyroglobulin (669 kDa) and dextran blue (Vo).

1029 Analytical ultracentrifugation: sedimentation velocity

1030 Sedimentation velocity assays and subsequent raw data analysis were performed in the 1031 Molecular Interactions Facility of the Centro de Investigaciones Biológicas Margarita 1032 Salas. Samples (320 μ L) in PBS containing 5% (v/v) glycerol and 1 mM β -1033 mercaptoethanol were loaded into analytical ultracentrifugation cells, which were run at 1034 20°C and 48,000 rpm in a XL-I analytical ultracentrifuge (Beckman-Coulter Inc.) 1035 equipped with UV-VIS absorbance and Raleigh interference detection systems, using an 1036 An-50Ti rotor, and 12 mm Epon-charcoal standard double-sector centerpieces. 1037 Sedimentation profiles were recorded at 230 nm. Differential sedimentation coefficient 1038 distributions were calculated by least-squares boundary modelling of sedimentation 1039 velocity data using the continuous distribution c(s) Lamm equation model as 1040 implemented by SEDFIT(Schuck, 2000). Experimental Svedberg coefficient values were 1041 corrected to standard conditions ($s_{20,w}$, water, 20°C, and infinite dilution) using 1042 SEDNTERP software

1043 Dynamic Light Scattering, DLS

1044 DLS experiments were carried out in a Protein Solutions DynaPro MS/X instrument at 1045 20C using a 90° light scattering cuvette. DLS autocorrelation functions, average of at 1046 least 18 replicates, were collected with Dynamics V6 software. Analysis evidenced in 1047 most cases the presence of two diffusing species, one with faster diffusion corresponding 1048 to a discrete major species and a second with substantially slower diffusion, 1049 corresponding to higher order species, with a minor contribution to the whole population 1050 within the sample. Exceptions were the globular domain GHD that appeared as a single 1051 species, and the coiled-coil construct with a larger contribution of the higher order 1052 species. Dynamics software was also employed to export the data as text files for parallel 1053 analysis using user-written scripts and functions in MATLAB (Version 7.10, MathWorks, 1054 Natick, MA). A double exponential decay model was fit to the data via nonlinear least 1055 squares, using as starting values the translational diffusion coefficients and relative 1056 amounts of the two different species, using as starting values those from the 1057 regularization analysis and their masses (those of the discrete species as estimated by 1058 the Svedberg equation). These values were compatible with the experimental data,

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rendering a similar best-fit value of the diffusion coefficient for the major species andallowing to assess its probability distribution.

1061 Estimate of molar mass from hydrodynamic measurements.

1062 The apparent molar masses of Uso1 and its mutants were calculated via the Svedberg 1063 equation, using the *s*- and *D*-values of the major species independently measured by 1064 sedimentation velocity and DLS, respectively.

1065 Fluorescence Microscopy

1066 A. nidulans hyphae were cultured in Watch Minimal Medium WMM (Peñalva, 2005). Image acquisition equipment, microscopy culture chambers and software have been 1067 1068 detailed (Pinar et al., 2022; Pinar and Peñalva, 2020). Simultaneous visualization of 1069 green and red emission channels was achieved with a Gemini Hamamatsu beam splitter 1070 coupled to a Leica DMi8 inverted microscope. Z-Stacks were deconvolved using 1071 Huygens Professional software (version 20.04.0p5 64 bits, SVI). Images were contrasted 1072 with Metamorph (Molecular Devices). Statistical analysis was performed with GraphPad 1073 Prism 8.02 (GraphPad). Uso1-GFP time of residence in cisternae was estimated from 1074 3D movies consisting of middle planes with 400 photograms at 2 fps time resolution. 1075 Each Uso1 puncta considered in the analysis was tracked manually with 3D (x,y,t)1076 representations generated with Imaris software (Oxford Instruments) combined with 1077 direct observation of photograms in movies and kymograph representations traced 1078 across >25 px-wide linear ROI covering the full width of the hyphae.

1079 AlphaFold predictions

1080 AlphaFold2 (Jumper et al., 2021) predictions were run using versions of the program 1081 installed locally and on ColabFold (Mirdita al., 2022) et with the 1082 AlphaFold2_advanced.ipynb notebook and the MMseqs2 MSA option. In all cases, the 1083 five solutions predicted by AlphaFold2 by default were internally congruent, and we 1084 always chose the one ranked first by the software. Uso1 GHD (1-674), Bos1, Bet1 and 1085 RAB1 were initially submitted as hetero-oligomers. Subsequently, predictions were 1086 submitted as 1:1 complexes of Uso1 GHD with Bos1, Bet1 and RAB1, as described in 1087 the table below. The solutions were also very similar when comparing the different 1088 combinations displayed in the table were fed to the software, strongly supporting the 1089 validity of the results.

1090

	AlphaFold2 runs, subunits included	
Uso1 GHD	Bet1	

Uso1 GHD	Bet1	Bos1	
Uso1 GHD	Bet1	Bos1	Rab1
Uso1 GHD	Bet1 (1-49)		
Uso1 GHD (N-terminal)	Bet1 (1-49)		
Uso1 GHD	Bet1 (1-49)	Bos1 Habc (1-126)	
Uso1 GHD E6K/G540S	Bet1 (1-49)	Bos1 Habc (1-126)	
Uso1 GHD		Bos1	
Uso1 GHD		Bos1 Habc (1-126)	
Uso1 GHD dimer		Bos1 Habc (1-126)	
Uso1 GHD E6K/G540S		Bos1	
Uso1 GHD E6K/G540S		Bos1 Habc (1-126)	
Uso1			
Uso1 dimer			
Uso1 dimer E6K/G540S			
Bet1	Bos1	Sec22	Sed5
Rud3 dimer			
Coy1 dimer			
Grh1	Bug1		

1091

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- 1102 Additional information
- 1103 Competing interests
- 1104 The authors declare that there are no competing interests

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1106 Availability statement

All DNA molecules used here may reconstructed by standard techniques using primers listed in supplemental Table II. All strains listed under supplemental Table I are available for academic purposes upon reasonable request to the corresponding author. They are deposited and maintained by the corresponding laboratory in the so-denoted Madrid (MAD) collection.

- 1112
- 1113

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1115

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Agencia Estatal de Investigación, Ministerio de Ciencia e Innovación, Spain	• • • • • • • • • • • • • • • • • • • •	Ignacio Bravo-Plaza
Agencia Estatal de Investigación, Ministerio de Ciencia e Innovación, Spain	PID2021-124278OB-100	Miguel A. Peñalva & Eduardo A. Espeso, co-IPs
Comunidad de Madrid	S2017/BMD-3691	Miguel A. Peñalva
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1116

1117 Author contributions

1118 Ignacio Bravo Plaza, Conceptualization, Data curation, Formal analysis, Validation, 1119 Investigation, Visualization, Writing — review and editing; Víctor G. Tagua, Data 1120 curation, Formal analysis, Validation, Investigation; Herbert N. Arst, Jr., 1121 Data curation. Formal analysis. Validation. Investigation. Conceptualization. 1122 Supervision, Writing — review and editing; Ana M. Alonso, Data curation, Formal 1123 analysis, Validation, Investigation; Mario Pinar, Conceptualization, Data curation, 1124 Formal analysis, Validation, Investigation, Supervision, Writing — review and editing ; 1125 Begoña Monterroso, Conceptualization, Data curation, Formal analysis, Validation, 1126 Supervision, Writing — review and editing; Antonio Galindo, Conceptualization, Data 1127 curation, Formal analysis, Validation, Investigation, Visualization, Writing - review and 1128 editing and Miguel Á. Peñalva, Conceptualization, Data curation, Formal analysis, 1129 Validation, Investigation, Supervision, Funding acquisition, Visualization, Writing -1130 original draft, Writing - review and editing.

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- 1140
- 1141
- 1142
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1144	
1145	Legends to figures
1146	Figure 1. Characterization of mutations bypassing the essential role of RAB1.
1147	(A). Genetic map in the region surrounding uso1 with genetic markers used as landmarks
1148	for mapping.
1149	(B). Molecular identification of the nucleotide changes in <i>suArab1ts</i> strains
1150	(C) and (D): growth tests showing <i>rab1ts</i> - and <i>rab1</i> Δ -rescuing phenotypes, respectively,
1151	of individual mutations, and synthetic positive interaction between E6K and G540S.
1152	Strains produce either green or white conidiospores (conidiospore colors are used as
1153	genetic markers). In (C), strains were point-inoculated. In (D) conidiospores were spread
1154	on agar plates to give individual colonies.
1155	Figure 1—figure supplement 1: E6K/G540S do not rescue lethality resulting from
1156	arf1 Δ , sly1 Δ or sed5 Δ .
1157	Top, uso1 is an essential gene. Singly-nucleated conidiospores derived from a
1158	heterokaryotic strain in which one class of nuclei carries a deficient pyrG uracil
1159	biosynthetic gene whereas the second class contains a $uso1\Delta$ allele tagged with
1160	functional <i>pyrG</i> were unable to grow on medium lacking pyrimidines at any of the
1161	tested temperatures. Bottom: Similar experiments showing that unlike $rab1\Delta$
1162	strains, strains carrying lethal arf1 Δ , sly1 Δ and sed5 Δ alleles cannot be rescued by
1163	uso1 ^{E6K/G540S} . Top panel, strains with green conidiospores; bottom, strains with
1164	white conidiospores.
1165	Figure 2: Localization of the amino acid substitutions within the Uso1
1166	AlphaFold2 structure
1167	(A). AlphaFold2 cartoon representations of A. nidulans Uso1 in monomeric and dimeric
1168	forms. Confidence estimations for this and other models are detailed under extended
1169	data S2. Red, N-terminal tail; marine blue, globular head domain; gray, coiled-coil; green,
1170	limit of the CTR. The rest of the CTR is shown as surface representation.
1171	(B). Amino acid alignment of fungal sequences with mammalian p115 showing strong
1172	conservation within the CTR: ANIDU, Aspergillus nidulans; PRUBE, Penicillium rubens;
1173	TREES, Thrichoderma ressei; SSCLE, Sclerotinia scleriotorum; MORYZ, Magnaporthe
1174	oryzae; CIMM, Coccidioides immitis; BOVIN, Bos taurus.
1175	(C). Position of the Gly6Lys and Gly540Ser substitutions. Only the GHD of dimeric full-

- 1176 length Uso1 are shown. The two different chains are colored in green and yellow,
- 1177 respectively. Distances between mutated residues are displayed in armstrongs.
- 1178 (D). The N-terminal amphipathic α -helix affected by the Glu6Lys substitution.

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1179 Figure 2—figure supplement 1. AlphaFold2 predictions of Uso1

- 1180 Ribbon representation of AlphaFold 2-predicted structures of full-length Uso1 (A) 1181 and Uso1 GHD (B), color-coded by pLDDT values. Graphs at the bottom are the 1182 corresponding plots of predicted aligned error of the residues (PAE).
- 1183

1184Figure 3: Determining molecular masses and oligomerization status of the

1185 different Uso1 constructs by velocity sedimentation analysis

The different panels display the sedimentation profiles of the protein being analyzed, with % of the main species, scheme of the different constructs and their limits and pictures of Coomassie stained-gels showing the purity of the protein preparations. The table below depicts biophysical parameters of the constructs used to obtain relative molecular masses. *s*_{exp} is the experimentally determined Svedberg coefficient; *D*_{exp}, translational diffusion coefficient of the main species; *Mr*, molecular mass deduced from Svedberg equation; M1 predicted molecular mass of the monomer; n = (*Mr* /M1).

1193 Figure 3—figure supplement 1: GHD is a monomer across a range of

1194 concentrations

1195 (A) and (B). Sedimentation velocity experiments with wild-type and E6K/G540S 1196 mutant GHD, respectively, showing that they behave as monomers at 1197 concentrations up to 5 μ M.(C). Sedimentation velocity profile of E6K/G540S 1198 mutant GHD lacking the His-tag, showing that the presence of the latter does not 1199 interfere with oligomerization, and a picture of a Coomassie stained gel showing 1200 the purity of the protein preparation on the right.

1201 Figure 4: Subcellular localization of Uso1

1202 (A). Uso1-GFP localizing to punctate cytoplasmic structures, 3D shaded by software.

- (B). Sections of a deconvolved Z-stack and its corresponding MIP. Uso1-GFP in invertedgreyscale for clarity
- 1205 (C). Kymograph showing the transient recruitment of Uso1 to punctate cytoplasmic1206 structures.
- 1207 (D). Average time of residence of Uso1 in these structures. Error bars, 95% Cl.
- 1208 (E). Example of one such structures visualized with a kymograph and with the 1209 corresponding movie frames (Movie 4).
- 1210
- 1211Figure 4—figure supplement 1: Methodology for tracking the half-life of1212Uso1-GFP on punctate structures.

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1213	(1) 3D movies were acquired by streaming pictures to the computer RAM at 2 fps.
1214	Appropriate reduction of excitation light intensity permitted acquisition of 400
1215	frames without apparent phototoxicity.

- 1216 (2) The behavior of punctate structures over time was represented in kymographs,
- 1217 in which vertical lines represent the time of residence of Uso1 on membranes.
- 1218 (3). As vertical lines frequently overlapped, jeopardizing the quality of this analysis,
- we imported the time series into a 3D viewer as if they were (x, y, z) series. Rotation
 across the different axes facilitated unambiguous tracking of the trajectories across
 time.
- 1222 (4) The length of the trajectories was measured and converted to time units.1223 Bottom graphs display examples of time trajectories.
- 1224

1225 Figure 5: Uso1 puncta do not colocalize with ERESs

- (A). Low extent co-localization of Sec13 ERES and Uso1 structures. Z-stacks for the two
 channels were acquired simultaneously, deconvolved and represented as MIPs. Two
 rare examples of colocalization are arrowed.
- (B). Photograms of a dual channel Z-stack with a Sec13-labeled nuclear envelope
 focused in the middle plane, illustrating that while some puncta show colocalization, the
 red Sec13 signal and the green Uso1 signal do not usually overlap.
- 1232 (C). A MIP of the same z-stack showing orthogonal views with some overlapping puncta1233 (arrows).
- (D). A ts mutation in the *sarA* gene encoding the SarA^{Sar1} GTPase governing ER exit
 markedly reduces the number of Uso1-GFP puncta upon shifting cells to restrictive
 conditions. Box-and-whisker plots: Statistical comparison was made using one-way
 ANOVA with Dunn's test for multiple comparisons. Whiskers are in Tukey's style: Only
 significant differences were indicated, using asterisks.
- 1239

1240 Figure 6: Uso1 localizes to RAB1-containing Golgi cisternae

- 1241 (A). Tip cells showing Uso1 colocalization with the indicated subcellular markers. Images
- 1242 are MIPs of deconvolved Z-stacks.
- 1243 (B). Magnified images of the color-coded shaded regions of the cells shown in A.
- 1244 (C). Pearson's coefficients of the different combinations.
- 1245

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1246	Figure 7: Uso1 localization to punctate structures is dependent on RAB1
1247	(A). Complete de-localization of Uso1-GFP to the cytosol by rab1ts and relocalization by
1248	E6K/G540S.
1249	(B). Uso1-GFP and rab1ts show a synthetic negative interaction that is rescued by the
1250	E6K/G540S double substitution. Strains in lanes 7 and 8 carry the wA2 mutation resulting
1251	in white conidiospores.
1252	
1253	Figure 8. Genetic evidence showing that the CTR region of Uso1 contributes to
1254	its recruitment to membranes.
1255	(A). Top, scheme of the predicted interactions. Bottom, engineering a gene-replaced
1256	allele lacking the CTR domain by homologous recombination.
1257	(B). The Bug1 C-terminal residues fit into the groove formed between the two Grh1 PDZ
1258	domains and into the pocket of the N-terminal PDZ domain (PDZ1).
1259	(C). A gene-replaced $uso1 \triangle CTR$ allele encoding a protein truncated for the CTR domain
1260	shows a synthetic negative interaction with <i>rab1ts</i> .
1261	(D). Western blot analysis. Removal of the CTR does not result in Uso1 instability.
1262	(E). $bug1\Delta$ and $grh1\Delta$ show a synthetic negative interaction with $rab1ts$ that is rescued
1263	by the double E6K/G540S substitution in Uso1
1264	Figure 8—figure supplement 1: AlphaFold2 modelling of Grh1-Bug1.
1265	(A). Cartoon, with alpha-helices shown as cylinders, of the nearly N-terminal PDZ
1266	domains of Grh1
1267	(B). AlphaFold 2 prediction of a 1:1 Grh1-Bug1 complex, trimmed of disordered
1268	regions
1269	(C). complete AlphaFold2 model of Grh1-Bug1 with color-coded model
1270	confidences values.
1271	Figure 9: The GHD of Uso1 is sufficient to support cell viability
1272	(A). Gene-replaced uso1 ^{GHD} allele carrying the double E6K/G540S substitution is
1273	sufficient to rescue viability at 30°C, but not at higher temperatures. (B). The Uso1 GHD
1274	is a monomer in vivo. Fractions collected from Superose columns loaded with the
1275	indicated protein extracts and reference His-tagged proteins were collected and
1276	analyzed by western blotting with $lpha$ -HA and $lpha$ -His antibodies. (C) Truncating Uso1 after
1277	the GHD results in markedly reduced protein levels, as determined by $\alpha\text{-Uso1}$ GHD
1278	western blotting. The band (yellow asterisks) moving slower than Uso1 (magenta
1279	asterisk) and at nearly the same position of Uso1 Δ CTR (blue asterisk) represents cross-
1280	reacting contaminants unrelated to Uso1. The right panel shows a longer exposure for

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1281 the indicated region, to reveal the faint GHD band (green dot). (D). Overexpression of 1282 Uso1, wild-type and E6K/G540S mutant, under the control of the *inuA* promoter, which 1283 is turned off on glucose and induced on sucrose. Western blots reacted with α -Uso1 1284 GHD antiserum. (E). Overexpressed GHD, be it E6K/G540S or wild-type, as the only 1285 source of Uso1 supports viability.

1286

Figure 10: Screening the preferential association of proteins acting in the ER/Golgi interface with E6K/G540S Uso1.

- (A). S-tagged baits (Uso1, wt and E6K/G540S, and the unrelated protein BapH), 1289 1290 expressed after gene replacement, were captured with their associated polypeptides on 1291 S-protein agarose beads. Candidate associates, also expressed after gene replacement, 1292 were tagged with HA3. (B). Schematic depiction of the proteins listed in these 1293 experiments showing their sites of action.(C). Anti-HA3 western blot analysis of the 1294 indicated S-bait and HA3-prey combinations. Equal loading of Uso1 proteins was 1295 confirmed by silver staining of precipitates. Note that BapH, chosen as negative control, 1296 is expressed at much higher levels than Uso1 proteins. Each panel is a representative 1297 experiment of three experimental replicates.
- 1298 **Figure 10—figure supplement 1**.
- (A) AlphaFold2 model, with PAE plot, of the GRIP domain of A. nidulans RUD3,predicted to be a dimer.
- (B) AlphaFold2 model of Coy1, calculated as a monomer. TMH is the nearly C terminal transmembrane helix that contributes to its recruitment to membranes

Figure 10—figure supplement 2: Growth phenotypes of null mutants of genes encoding golgins.

- 1305(A). Ablation of individual golgins Bug1/Grh1 and Rud3 does not result in1306detectable growth defects. $coy1\Delta$ strains have a subtle growth phenotype.
- 1307 (B). Negative effects of $grh1\Delta$, $bug1\Delta$, $coy1\Delta$ and $rud3\Delta$ on the ability of 1308 $uso1^{E6K/G540S}$ to rescue $rab1\Delta$. Note that $coy1\Delta$ and $rab1\Delta$ $uso1^{E6K/G540S}$ are 1309 synthetically lethal. For convenience, this set of strains carried a mutation resulting 1310 in white conidiospores, as opposed to the wild-type green color.
- 1311

Figure 11: The Uso1 GHD interacts directly with Bos1 and Bet1 SNAREs acting in the ER/Golgi interface

1314 (A). Purified fusion proteins in which the cytosolic domains of the indicated SNAREs
 1315 have been fused to GST were used in pulldown experiments with His-tagged, purified

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1316 wild-type and E6K/G540S Uso1. The plasma membrane Qa syntaxin Sso1 was used as 1317 negative control. Pulled-down material was analyzed by anti-His western blotting. (B). 1318 Quantitation of the above experiment; significance was determined by unpaired t-student 1319 tests. Error bars represent S.E.M. (C). As in A, but using in vitro synthesized, HA3-tagged 1320 Sly1 as prey. Samples were analyzed by anti-HA western blotting. (D). As in A, but using 1321 wild-type and mutant GHD as preys, rather than full-length Uso1. (E). Quantitation of the 1322 experiment in D. (F). GST pull-down experiment comparing the ability of the GHD to 1323 interact with the early Golgi Qb and Qc SNAREs (Bos1 and Bet1), with that of their 1324 medial Golgi counterparts (Qb Gos1 and Qc Sft1).

1325

1326 Figure 11—figure supplement 1. AlphaFold2 prediction of the ER/Golgi SNARE

- 1327 **bundle**.
- 1328 (A) Sec5/Bos1/Bet1/Sec22 predicted SNARE bundle. (B) Quality control (pLDDT,
 1329 color coded, and PAE) of the model.

Figure 12. AlphaFold2 models provide insight into the additive mode of suppression shown by E6K and G540S.

- 1332 (A). Model of full length Uso1 bound to the ER/Golgi SNAREs Bos1 and Bet1.
- 1333 (B). Top, ribbon representation of the Bos1 N-terminal Habc domain and Uso1GHD.

1334 Bottom, Inset combining surface and ribbon depiction.

(C). Increased binding of Bos1 to G540S Uso1 appears to involve insertion of Ser540
into a pocket located in the Habc domain of the Qb SNARE. Partial view of the Bos1Uso1 GHD surface of interaction in the wild type (left) and mutant (right) models. G540
and S540 are annotated.

1339 (D). The N-terminal amphipathic α -helix of Uso1 comprising the E6K substitution lies 1340 within a flexible stretch of the protein that might facilitate its insertion into membranes. 1341 Alignment of six independent predictions, with Glu6 highlighted in red. The Uso1 GHD 1342 was modeled alone, in a complex with SNARE proteins or with Ypt1. The N-terminal a-1343 helix (boxed) adopts different positions, suggesting high flexibility.

1344 Figure 12—figure supplement 1. AlphaFold2 prediction of the Bet1-GHD

1345 interaction.

1346The putative binding surface of Bet1 and Uso1 as determined by AlphaFold2. Top1347images, cartoon of Bet1-GHD interactions, colored by pLDDT score. Alignment of1348four independent predictions involving the Bet1 N-terminal region and Uso1 GHD.1349A single model for Uso1 GHD is shown on the top representation for simplicity. In1350spite of the disordered nature of the N-terminal Bet1 region, the Bet1-Uso1 binding

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interface is consistent among models. Bottom, surface representation of the Nterminal Bet1 region (orange) in complex with the GHD. Also indicated is the Habc
domain of Bos1 (yellow) bound to the GHD.

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Figure 12—figure supplement 2. AlphaFold 2 prediction of the RAB1 binding site on the Bet1/Bos1/Uso1 GHD complex.

- 1357 (A) and (B): cartoon representation of the GHD-RAB1 complex. The model is1358 depicted as pipes and planks
- 1359 (C): Orthogonal views of the Uso1 GHD-RAB1-Bet1-Bos1Habc structural model.
- 1360The Uso1 GHD is shown as surface to emphasize the distant binding sites of the1361Bos1 Habc domain, RAB1 and the Bet1 N-terminal region
- 1362 (D): Ribbon representation of the model shown in C but including the full-length
- 1363 SNARE subunits, i.e. the GHD domain of Uso1, RAB1 and the SNARES Bet1 and
- Bos1. Proteins are in the correct orientation to connect membranes separated by
- 1365 23 nm, counting from the SNARE TMDs to the prenylated RAB1 residues.
- 1366

1367 Figure 12—figure supplement 3. Quality control assessment of AlphaFold2

- 1368 predictions for the indicated complexes.
- 1369

1370 Supplemental Table I

1371 List and complete genotypes of *A. nidulans* strains used in this work

1372 Supplemental Table II

1373 Primers used for PCR-based genetic manipulations

1374 Figure 8 source data

- 1375 Raw images for western blots in panel D and uncropped pictures with used exposures
- 1376 and regions indicated.

1377 Figure 9 source data

- 1378 For panels B, C, D; raw images for western blots and uncropped pictures with used
- 1379 exposures and regions indicated.

1380 Figure 10 source data

1381 Raw images for western blots and silver-stained gels and uncropped pictures with used1382 exposures and regions indicated.

1383	Figure 11 source data
1384 1385	Raw images for western blots and Coomassie-stained gels and uncropped pictures with used exposures and regions indicated.
1386	
1387	Rich file media
1388	Video 1: Shaded 3D reconstruction of a hypha expressing Uso1-GFP
1389	Video 2: 4D acquisition showing the dynamics of Uso1-GFP.
1390	4D (x, y, z, t) in which Z-stacks were acquired at a rate of 1 frame every 2.6 sec
1391	Video 3: Dynamics of Uso1-GFP at 2 fps
1392	3D acquisition (200 frames) showing the dynamics of Uso1-GFP. Time resolution, 2 fps
1393	Video 4: Single Uso1-GFP cisterna tracked over time
1394	Example of Uso1-GFP cisterna. The video contains 96 photograms acquired at 2fps
1395	Video 5: ·3D reconstruction of a hypha expressing fluorescently labeled Uso1-
1396	GFP and Sec13-mCh
1397	There is little colocalization between Uso1-GFP and Sec13 ERES
1398	Video 6: ·4D video (1 fpm) of a hypha expressing fluorescently labeled Uso1-GFP
1399	and Sec7-mCh
1400	Uso1 does not colocalize at all with the TGN marker Sec7

1402	
1403 1404 1405	References
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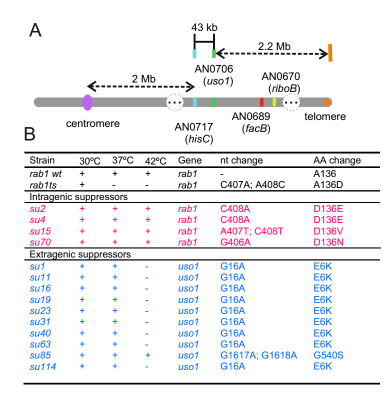
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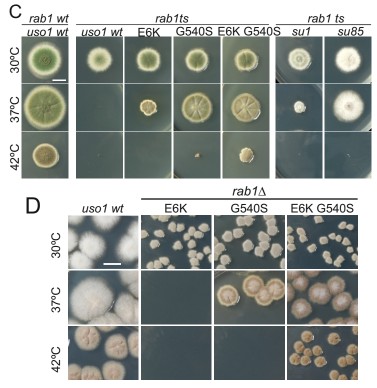


Figure 1. Characterization of mutations bypassing the essential role of RAB1

(A). Genetic map in the region surrounding *uso1* with genetic markers used as landmarks for mapping.B). Molecular identification of the nucleotide changes in *suArab1*^{ts} strains (C) and (D): growth tests showing *rab1*^{ts}- and *rab1* Δ -rescuing phenotypes, respectively, of individual mutations, and synthetic positive interaction between E6K and G540S. Strains produce either green or white conidiospores (conidiospore colors are used as genetic markers). In (C), strains were point-inoculated. In (D) conidiospores were spread on agar plates to give individual colonies.

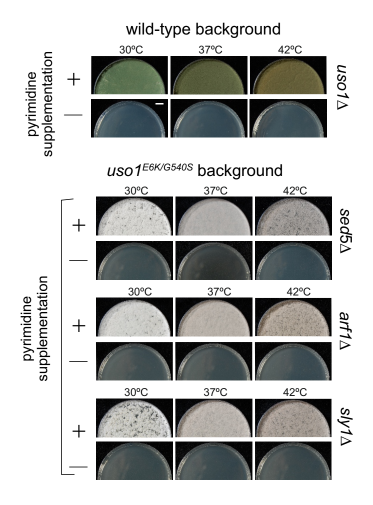


Figure 1—figure supplement 1: E6K/G540S do not rescue lethality resulting from arf1 Δ , sly1 Δ or sed5 Δ .

Top, *uso1* is an essential gene. Singly-nucleated conidiospores derived from a heterokaryotic strain in which one class of nuclei carries a deficient *pyrG* uracil biosynthetic gene whereas the second class contains a *uso1* Δ allele tagged with functional *pyrG* were unable to grow on medium lacking pyrimidines at any of the tested temperatures. Bottom: Similar experiments showing that unlike *rab1* Δ strains, strains carrying lethal *arf1* Δ *,sly1* Δ and *sed5* Δ alleles cannot be rescued by *uso1*^{E6K/G540S}. Top panel, strains with green conidiospores; bottom, strains with white conidiospores.

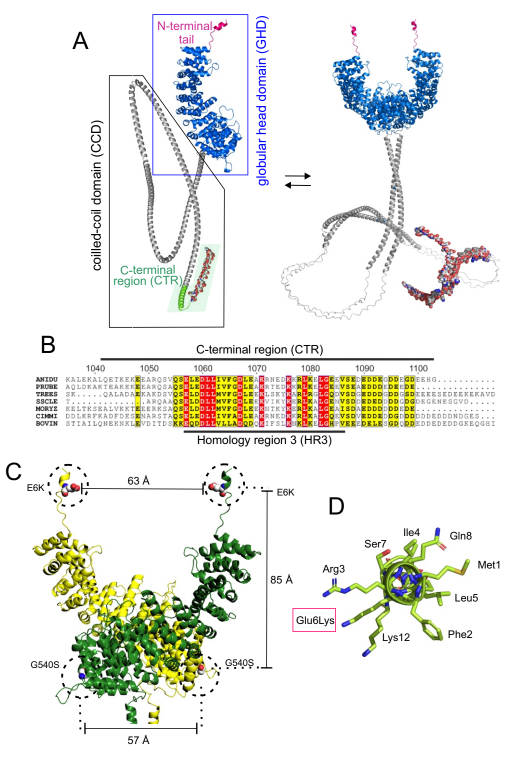
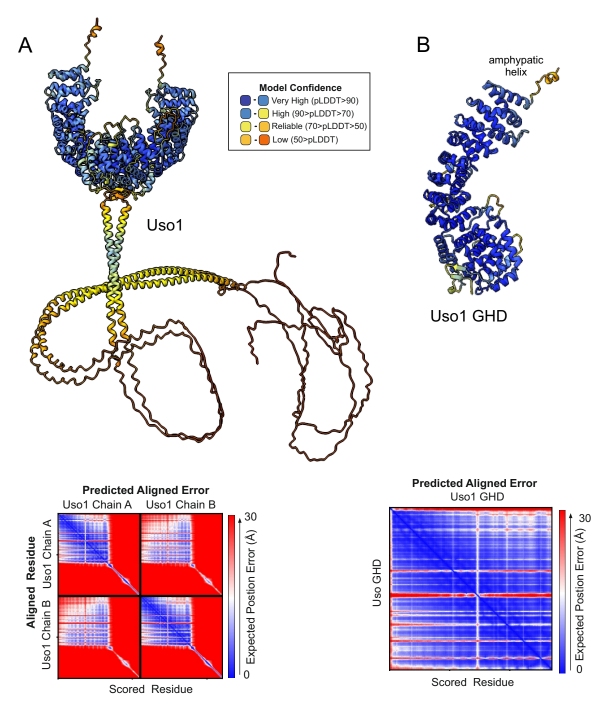


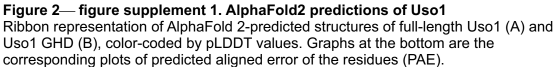
Figure 2: Localization of the amino acid substitutions within the Uso1 AlphaFold2 structure (A). AlphaFold2 cartoon representations of *A. nidulans* Uso1 in monomeric and dimeric forms. Red, N-terminal tail; marine blue, globular head domain; gray, coiled-coil; green, limit of the CTR. The rest of the CTR is shown as surface representation.

(B). Amino acid alignment of fungal sequences with mammalian p115 showing strong conservation within the CTR: ANIDU, *Aspergillus nidulans*; PRUBE, *Penicillium rubens*; TREES, *Thrichoderma ressei;* SSCLE, *Sclerotinia scleriotorum*; MORYZ, *Magnaporthe oryzae*; CIMM, *Coccidioides immitis*; BOVIN, *Bos taurus*.

©. Position of the Gly6Lys and Gly540Ser substitutions. Only the GHD of dimeric full-length Uso1 are shown. The two different chains are colored in green and yellow, respectively. Distances between mutated residues are displayed in armstrongs.

(D). The N-terminal amphipathic α -helix affected by the Glu6Lys substitution.





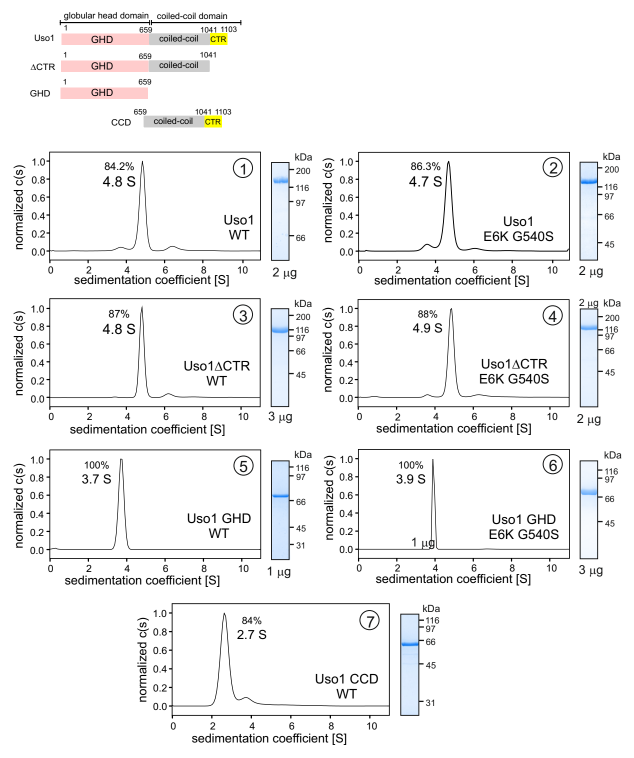


Figure 3: Determining molecular masses and oligomerization status of the different Uso1 constructs by velocity sedimentation analysis

The different panels display the sedimentation profiles of the protein being analyzed, with % of the main species, scheme of the different constructs and their limits and pictures of Coomassie stained-gels showing the purity of the protein preparations. The table below depicts biophysical parameters of the constructs used to obtain relative molecular masses. s_{exp} is the experimentally determined Svedberg coefficient; D_{exp} , translational diffusion coefficient of the main species; Mr, molecular mass deduced from Svedberg equation; M_1 predicted molecular mass of the monomer; $n = (M_r/M_1)$

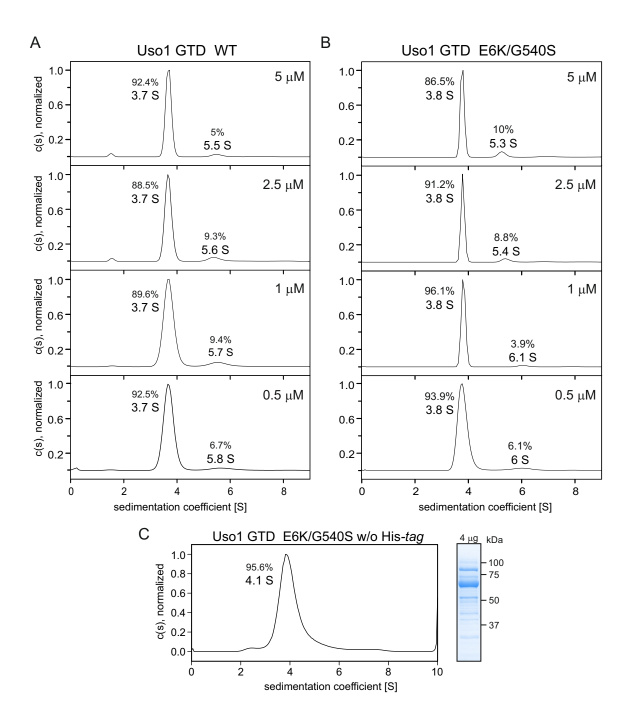


Figure 3—figure supplement 1: GHD is a monomer across a range of concentrations (A) and (B). Sedimentation velocity experiments with wild-type and E6K/G540S mutant GHD, respectively, showing that they behave as monomers at concentrations up to 5 μ M.(C). Sedimentation velocity profile of E6K/G540S mutant GHD lacking the His-tag, showing that the presence of the latter does not interfere with oligomerization, and a picture of a Coomassie stained gel showing the purity of the protein preparation on the right.

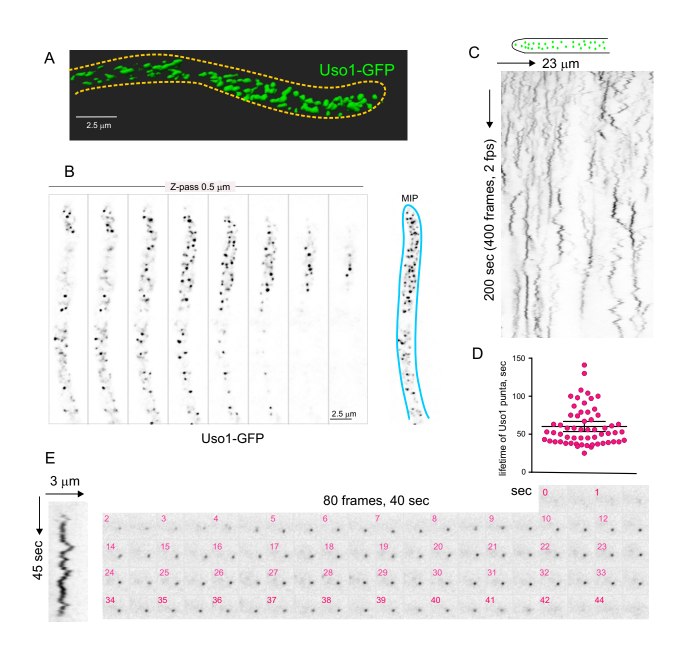


Figure 4: Subcellular localization of Uso1

(A). Uso1-GFP localizing to punctate cytoplasmic structures, 3D shaded by software.

(B). Sections of a deconvolved Z-stack and its corresponding MIP. Uso1-GFP in inverted greyscale for clarity

(C). Kymograph showing the transient recruitment of Uso1 to punctate cytoplasmic structures.

(D). Average time of residence of Uso1 in these structures. Error bars, 95% Cl.

(E). Example of one such structures visualized with a kymograph and with the corresponding movie frames (Movie 4).

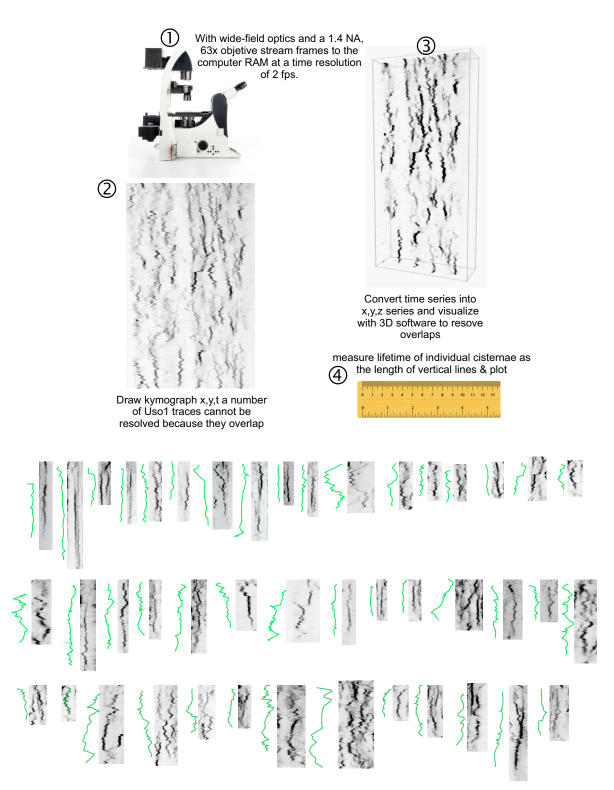


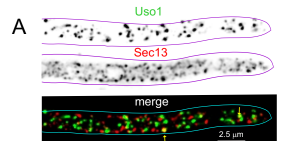
Figure 4—figure supplement 1: Methodology for tracking the half-life of Uso1-GFP on punctate structures.

1) 3D movies were acquired by streaming pictures to the computer RAM at 2 fps. Appropriate reduction of excitation light intensity permitted acquisition of 400 frames without apparent phototoxicity.

2) The behavior of punctate structures over time was represented in kymographs, in which vertical lines represent the time of residence of Uso1 on membranes.

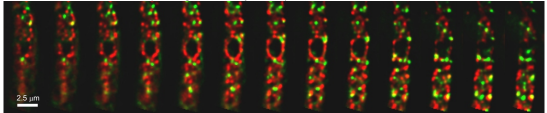
3). As vertical lines frequently overlapped, jeopardizing the quality of this analysis, we imported the time series into a 3D viewer as if they were (x, y, z) series. Rotation across the different axes facilitated unambiguous tracking of the trajectories across time.

4) The length of the trajectories was measured and converted to time units. Bottom graphs display examples of time trajectories.



В

→ z-stack: 1 photogram every 0.25 µm Sec13 Uso1



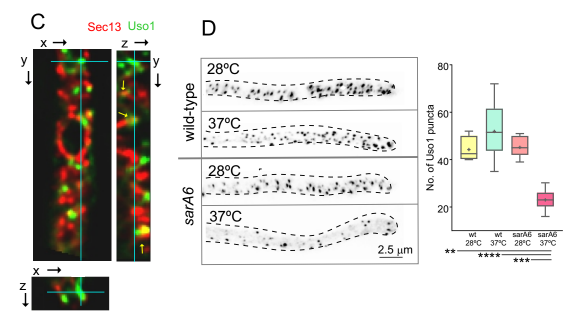


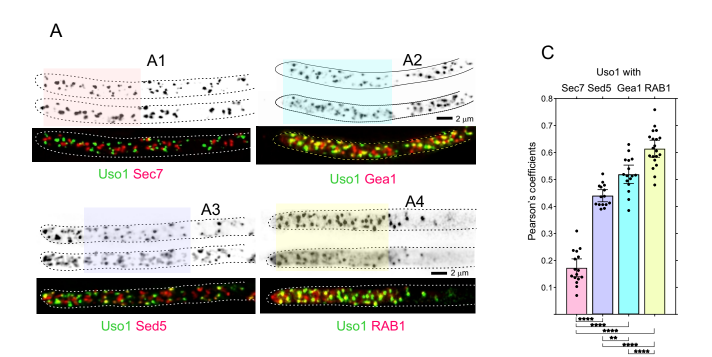
Figure 5: Uso1 puncta do not colocalize with ERESs

(A). Low extent co-localization of Sec13 ERES and Uso1 structures. Z-stacks for the two channels were acquired simultaneously, deconvolved and represented as MIPs. Two rare examples of colocalization are arrowed.

(B). Photograms of a dual channel Z-stack with a Sec13-labeled nuclear envelope focused in the middle plane, illustrating that while some puncta show colocalization, the red Sec13 signal and the green Uso1 signal do not usually overlap.

(C). A MIP of the same z-stack showing orthogonal views with some overlapping puncta (arrows).

(D). A *ts* mutation in the *sarA* gene encoding the SarA^{sar1} GTPase governing ER exit markedly reduces the number of Uso1-GFP puncta upon shifting cells to restrictive conditions. Box-and-whisker plots: Statistical comparison was made using one-way ANOVA with Dunn's test for multiple comparisons. Whiskers are in Tukey's style: Only significant differences were indicated, using asterisks.



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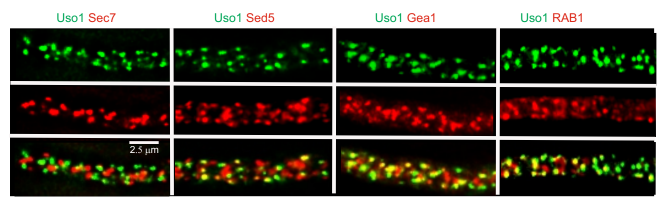


Figure 6: Uso1 localizes to RAB1-containing Golgi cisternae

(A). Tip cells showing Uso1 colocalization with the indicated subcellular markers. Images are MIPs of deconvolved Z-stacks.

(B). Magnified images of the color-coded shaded regions of the cells shown in A.

©. Pearson's coefficients of the different combinations

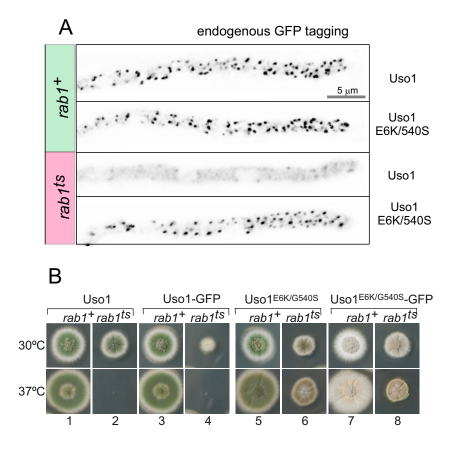


Figure 7: Uso1 localization to punctate structures is dependent on RAB1

(A). Complete de-localization of Uso1-GFP to the cytosol by *rab1ts* and relocalization by E6K/G540S.

(B). Uso1-GFP and *rab1ts* show a synthetic negative interaction that is rescued by the E6K/G540S double substitution. Strains in lanes 7 and 8 carry the *wA2* mutation resulting in white conidiospores.

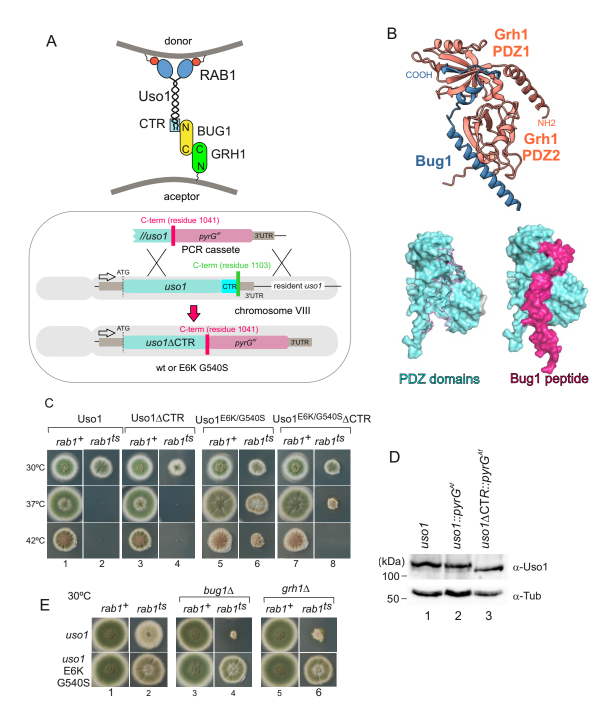


Figure 8. Genetic evidence showing that the CTR region of Uso1 contributes to its recruitment to membranes.

(A). Top, scheme of the predicted interactions. Bottom, engineering a genereplaced allele lacking the CTR domain by homologous recombination. (B). The Bug1 C-terminal residues fit into the groove formed between the two Grh1 PDZ domains and into the pocket of the N-terminal PDZ domain (PDZ1). (C). A gene-replaced $uso1 \triangle CTR$ allele encoding a protein truncated for the CTR domain shows a synthetic negative interaction with *rab1ts*.

(D). Western blot analysis. Removal of the CTR does not result in Uso1 instability. (E). $bug1\Delta$ and $grh1\Delta$ show a synthetic negative interaction with rab1ts that is rescued by the double E6K/G540S substitution in Uso1

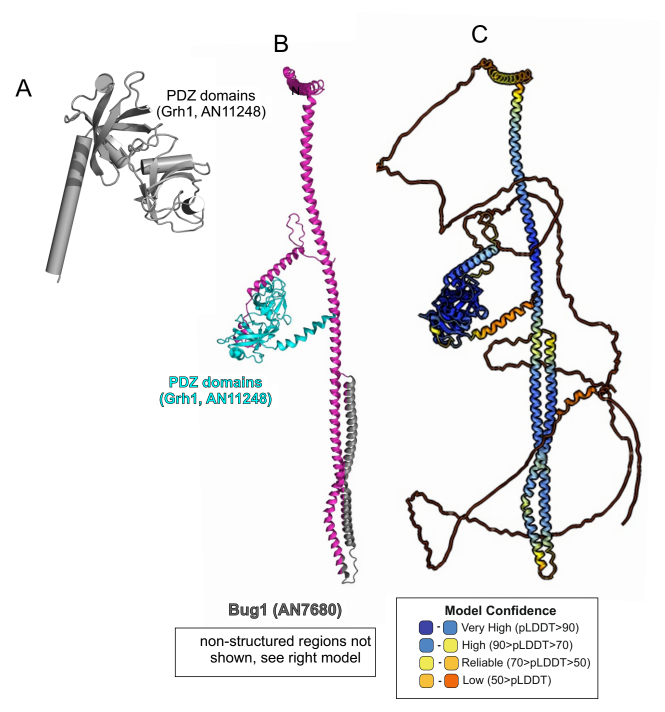


Figure 8 — figure supplement 1: AlphaFold2 modelling of Grh1-Bug1.

(A). Cartoon, with alpha-helices shown as cylinders, of the nearly N-terminal PDZ domains of Grh1

(B). AlphaFold 2 prediction of a 1:1 Grh1-Bug1 complex, trimmed of disordered regions ©. complete AlphaFold2 model of Grh1-Bug1 with color-coded model confidence values.

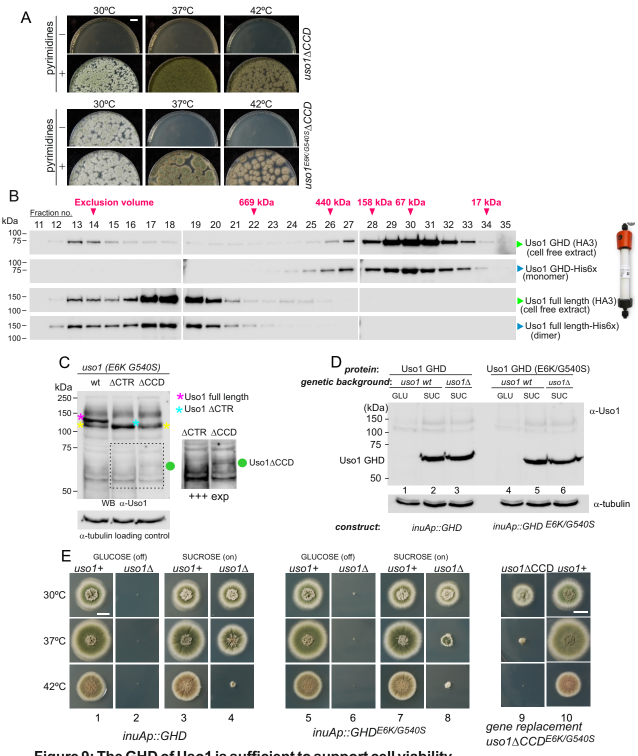


Figure 9: The GHD of Uso1 is sufficient to support cell viability (A). Gene-replaced *uso1*^{GHD} allele carrying the double E6K/G540S substitution is sufficient to rescue viability at 30°C, but not at higher temperatures. (B). The Uso1 GHD is a monomer in vivo. Fractions collected from Superose columns loaded with the indicated protein extracts and reference His-tagged proteins were collected and analyzed by western blotting with α -HA and α -His antibodies. (C) Truncating Uso1 after the GHD results in markedly reduced protein levels, as determined by α -Uso1 GHD western blotting. The band (yellow asterisks) moving slower than Uso1 (magenta asterisk) and at nearly the same position of Uso1 Δ CTR (blue asterisk) represents cross-reacting contaminants unrelated to Uso1. The right panel shows a longer exposure for the indicated region, to reveal the faint GHD band (green dot). (D). Overexpression of Uso1, wild-type and E6K/G540S mutant, under the control of the *inuA* promoter, which is turned off on glucose and induced on sucrose. Western blots reacted with α -Uso1 GHD antiserum. (E). Overexpressed GHD, be it E6K/G540S or wild-type, as the only source of Uso1 supports viability.

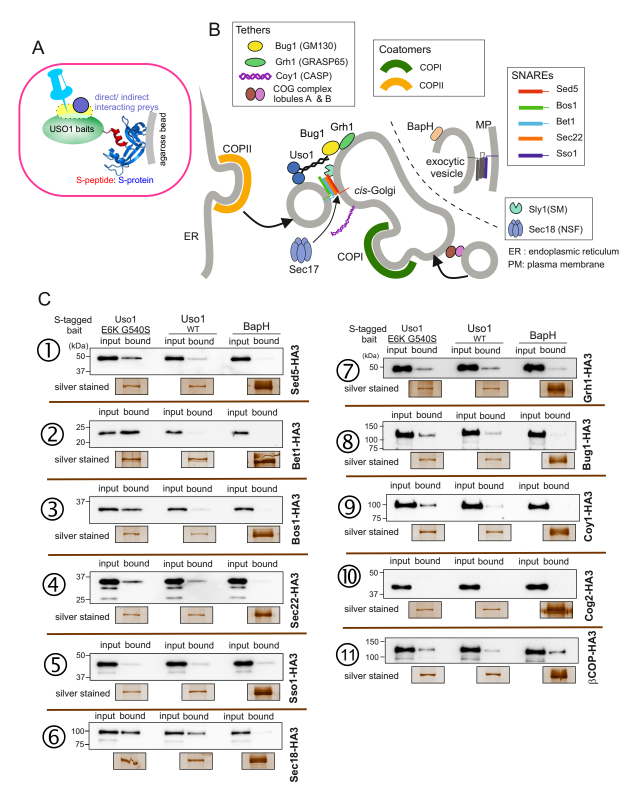


Figure 10: Screening the preferential association of proteins acting in the ER/Golgi interface with E6K/G540S Uso1.

(A). S-tagged baits (Uso1, wt and E6K/G540S, and the unrelated protein BapH), expressed after gene replacement, were captured with their associated polypeptides on S-protein agarose beads. Candidate associates, also expressed after gene replacement, were tagged with HA3. (B). Schematic depiction of the proteins listed in these experiments showing their sites of action.(C). Anti-HA3 western blot analysis of the indicated S-bait and HA3-prey combinations. Equal loading of Uso1 proteins was confirmed by silver staining of precipitates. Note that BapH, chosen as negative control, is expressed at much higher levels than Uso1 proteins. Each panel is a representative experiment of three experimental replicates.

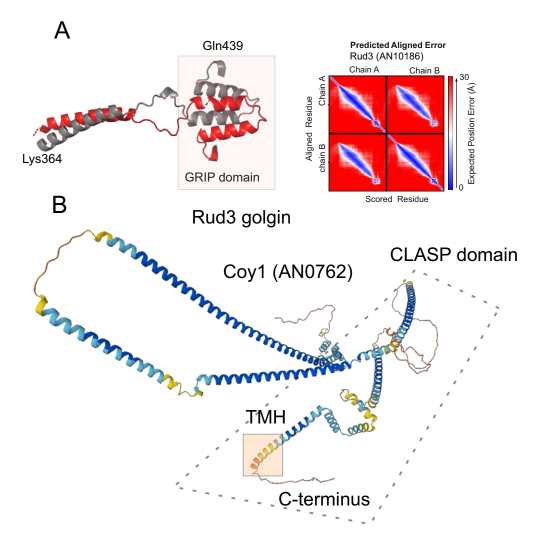


Figure 10—figure supplement 1.

/A) AlphaFold2 model, with PAE plot, of the GRIP domain of *A*. *nidulans* RUD3, predicted to be a dimer.
(B) AlphaFold2 model of Coy1, calculated as a monomer. TMH is the nearly C-terminal transmembrane helix that contributes to its recruitment to membranes

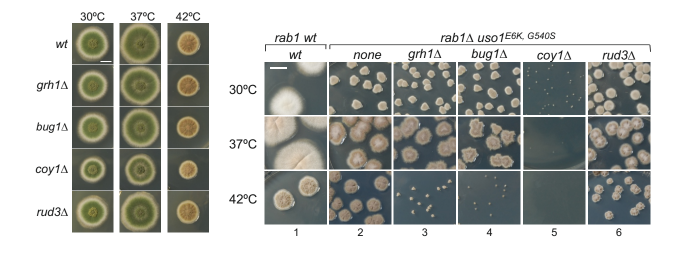
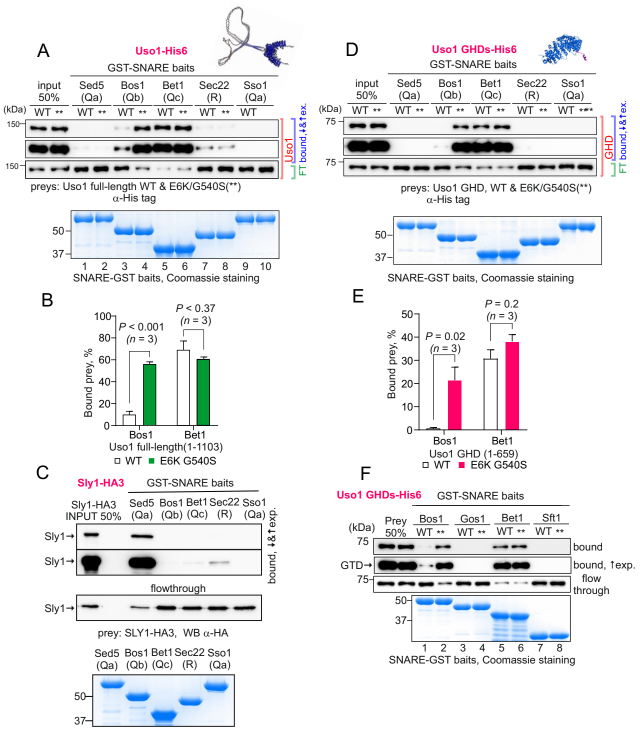


Figure 10—figure supplement 2: Growth phenotypes of null mutants of genes encoding golgins.

(A). Ablation of individual golgins Bug1/Grh1 and Rud3 does not result in detectable growth defects. $coy1\Delta$ strains have a subtle growth phenotype.

(B). Negative effects of $grh1\Delta$, $bug1\Delta$, $coy1\Delta$ and $rud3\Delta$ on the ability of $uso1^{E6K/G540S}$ to rescue $rab1\Delta$. Note that $coy1\Delta$ and $rab1\Delta$ $uso1^{E6K/G540S}$ are synthetically lethal. For convenience, this set of strains carried a wA1 mutation resulting in white conidiospores, as opposed to the wild-type green color.



SNARE-GST baits, Coomassie staining

Figure 11: The Uso1 GHD interacts directly with Bos1 and Bet1 SNAREs acting in the ER/Golgi interface

(A). Purified fusion proteins in which the cytosolic domains of the indicated SNAREs have been fused to GST were used in pulldown experiments with His-tagged, purified wild-type and E6K/G540S Uso1. The plasma membrane Qa syntaxin Sso1 was used as negative control. Pulled-down material was analyzed by anti-His western blotting. (B). Quantitation of the above experiment; significance was determined by unpaired *t*-student tests. Error bars represent S.E.M. (C). As in A, but using in vitro synthesized, HA3-tagged Sly1 as prey. Samples were analyzed by anti-HA western blotting. (D). As in A, but using wild-type and mutant GHD as preys, rather than full-length Uso1. (E). Quantitation of the experiment in D. (F). GST pull-down experiment comparing the ability of the GHD to interact with the early Golgi Qb and Qc SNAREs (Bos1 and Bet1), with that of their medial Golgi counterparts (Qb Gos1 and Qc Sft1).

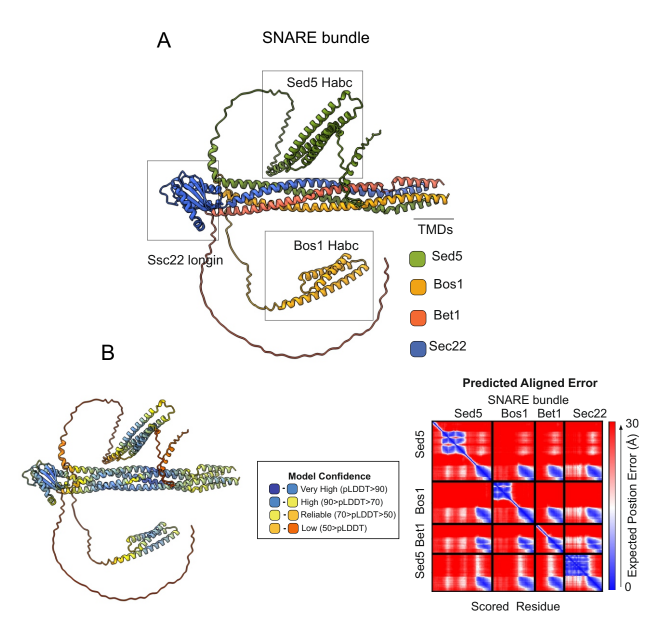


Figure 11—figure supplement 1. AlphaFold2 prediction of the ER/Golgi SNARE bundle.

(A) Sec5/Bos1/Bet1/Sec22 predicted SNARE bundle.

(B) Quality control (pLDDT, color coded, and PAE) of the model.

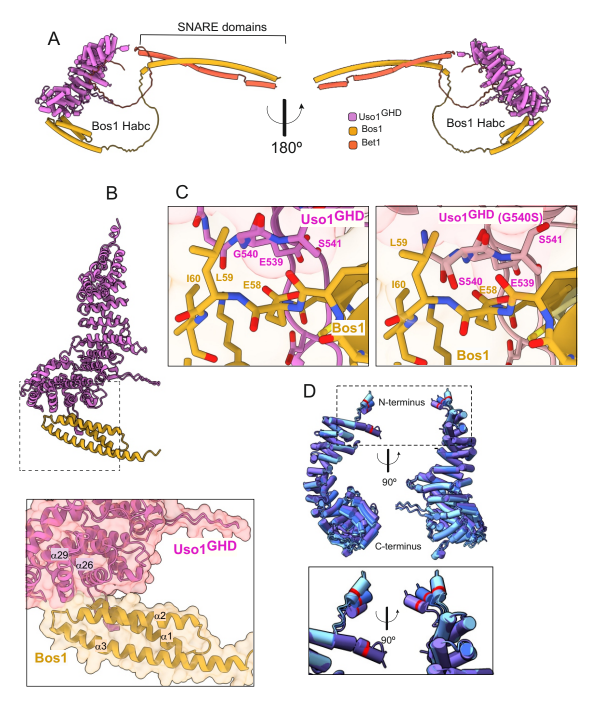


Figure 12. AlphaFold2 models provide insight into the additive mode of suppression shown by E6K and G540S.

(A). Model of full length Uso1 bound to the ER/Golgi SNAREs Bos1 and Bet1.

(B). Top, ribbon representation of the Bos1 N-terminal Habc domain and Uso1GHD. Bottom, Inset combining surface and ribbon depiction.

(C). Increased binding of Bos1 to G540S Uso1 appears to involve insertion of Ser540 into a pocket located in the Habc domain of the Qb SNARE. Partial view of the Bos1-Uso1 GHD surface of interaction in the wild type (left) and mutant (right) models. G540 and S540 are annotated.

(D). The N-terminal amphipathic α -helix of Uso1 comprising the E6K substitution lies within a flexible stretch of the protein that might facilitate its insertion into membranes. Alignment of six independent predictions, with Glu6 highlighted in red. The Uso1 GHD was modeled alone, in a complex with SNARE proteins or with Ypt1. The N-terminal α -helix (boxed) adopts different positions, suggesting high flexibility.

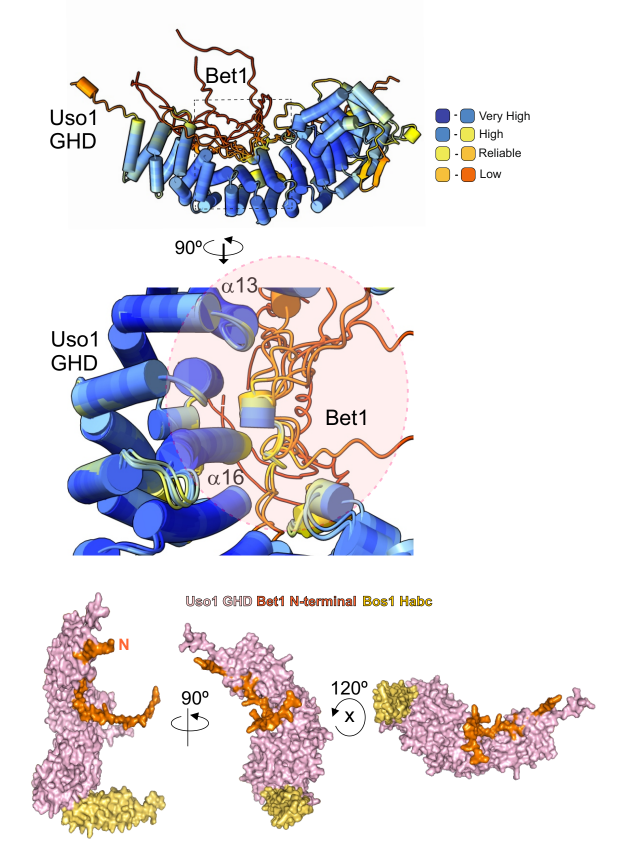


Figure 12—figure supplement 1. AlphaFold2 prediction of the Bet1-GHD interaction.

The putative binding surface of Bet1 and Uso1 as determined by AlphaFold2. Top images, cartoon of Bet1-GHD interactions, colored by pLDDT score. Alignment of four independent predictions involving the Bet1 N-terminal region and Uso1 GHD. A single model for Uso1 GHD is shown on the top representation for simplicity. In spite of the disordered nature of the N-terminal Bet1 region, the Bet1-Uso1 binding interface is consistent among models. Bottom, surface representation of the N-terminal Bet1 region (orange) in complex with the GHD. Also indicated is the Habc domain of Bos1 (yellow) bound to the GHD.

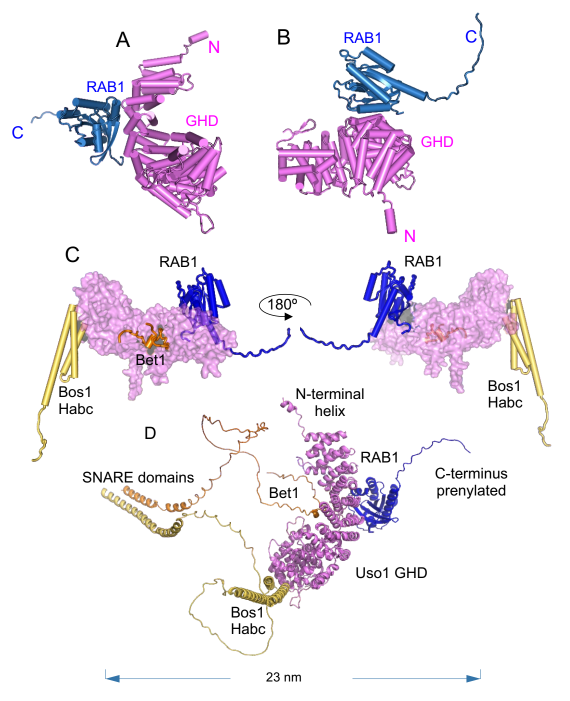


Figure 12—figure supplement 2. AlphaFold 2 prediction of the RAB1 binding site on the Bet1/Bos1/Uso1 GHD complex.

(A) and (B): cartoon representation of the GHD-RAB1 complex. The model is depicted as pipes and planks

(C): Orthogonal views of the Uso1 GHD-RAB1-Bet1-Bos1Habc structural model. The Uso1 GHD is shown as surface to emphasize the distant binding sites of the Bos1 Habc domain, RAB1 and the Bet1 N-terminal region

(D): Ribbon representation of the model shown in C but including the full-length SNARE subunits, i.e. the GHD domain of Uso1, RAB1 and the SNARES Bet1 and Bos1. Proteins are in the correct orientation to connect membranes separated by 23 nm, counting from the SNARE TMDs to the prenylated RAB1 residues.

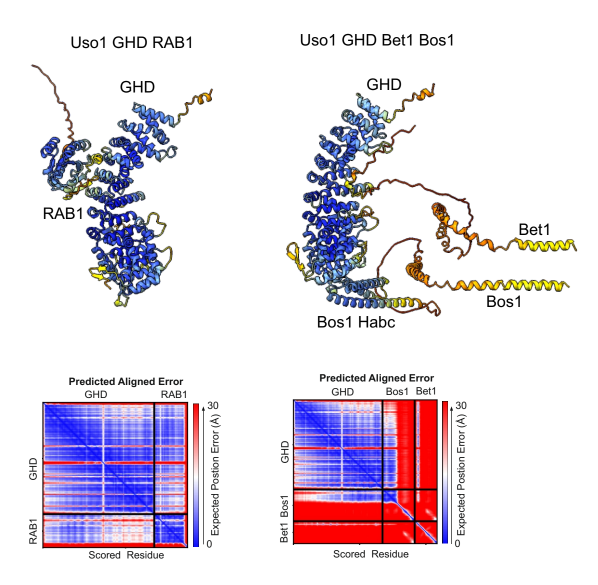


Figure 12—figure supplement 3. Quality control assessment of AlphaFold2 predictions for the indicated complexes.