Initial ciliary assembly in *Chlamydomonas* requires Arp2/3 complex-dependent endocytosis
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11 ABSTRACT

12 Ciliary assembly, trafficking, and regulation are dependent on microtubules, but the 13 mechanisms of ciliary assembly also require the actin cytoskeleton. Here, we dissect subcellular

roles of actin in ciliogenesis by focusing on actin networks nucleated by the Arp2/3 complex in

the powerful ciliary model, *Chlamydomonas*. We find the Arp2/3 complex is required for the

16 initial stages of ciliary assembly when protein and membrane are in high demand, but cannot

17 yet be supplied from the Golgi complex. We provide evidence for Arp2/3 complex-dependent

18 clathrin-mediated endocytosis of ciliary proteins, an increase in endocytic activity upon induction

19 of ciliary growth, and relocalization of plasma membrane proteins to newly formed cilia. Our

- 20 data support a new model of ciliary protein and membrane trafficking during early ciliogenesis
- 21 whereby proteins previously targeted to the plasma membrane are reclaimed by Arp2/3

22 complex-dependent clathrin-mediated endocytosis for initial ciliary assembly.

24 INTRODUCTION

The cilium of the unicellular, green alga *Chlamydomonas reinhardtii* has long been used as a model due to its structural and mechanistic conservation relative to the cilia of mammalian cells. Cilia consist primarily of microtubules that extend from the surface of the cell and are ensheathed in plasma membrane. Their assembly relies on microtubule dynamics and trafficking of protein and membrane (Nachury, Seeley, and Jin 2010), as well as intraflagellar transport (IFT), a motor-based transport system that moves tubulin and other cargo from the base of the cilium to the tip and back again (Pedersen and Rosenbaum 2008).

32 Although cilia are composed of microtubules and depend on them for assembly, the 33 mechanisms governing ciliary maintenance and assembly extend to other cytoskeletal 34 components, namely actin. In fact, the microtubule organizing center of the cell, the centrosome. 35 from which cilia are nucleated has been found to function as an actin organizer (Farina et al. 2016; Inoue et al. 2019). In mammalian cells, cortical actin disruption results in increased ciliary 36 37 length and percentage of ciliated cells (Kim et al. 2010; Park et al. 2008). Further, in mammalian 38 cells, when ciliogenesis is triggered by serum starvation, preciliary vesicles are trafficked to the 39 centriole where they fuse to form a ciliary vesicle around the budding cilium. In the intracellular 40 pathway of ciliogenesis, it has been shown that when Arp2/3 complex-branched actin is lost. 41 depletion of preciliary vesicles at the centriole occurs due to problems with vesicle fusion. 42 suggesting a role for branched actin in intracellular ciliogenesis (Wu, Chen, and Tang 2018). 43 Further, actin itself has even been found within cilia, suggesting that actin is a key protein 44 involved in ciliary maintenance and assembly (Kiesel et al. 2020).

45 Chlamydomonas cells are ideal for tackling the question of actin-dependent ciliary 46 trafficking due to their lack of a cortical actin network and their ability to undergo consistent and 47 robust ciliogenesis without requiring serum starvation. In Chlamydomonas, disruption of actin 48 networks with Cytochalasin D (CytoD) resulted in shorter steady-state cilia (W. L. Dentler and 49 Adams 1992) and disruption with Latrunculin B (LatB), which sequesters monomers leading to 50 eventual filament depolymerization, resulted in shortened cilia and impaired regeneration 51 (Avasthi et al. 2014; Jack et al. 2019). Chlamydomonas actin networks are required for 52 accumulation of IFT machinery at the base of cilia and for entry of IFT material into cilia (Avasthi 53 et al. 2014), as well as for trafficking of post-Golgi vesicles to cilia, the synthesis of ciliary 54 proteins, and the organization of the gating region at the base of cilia (Jack et al. 2019). Many 55 key advances in our understanding of the relationship between cilia and actin have been 56 discovered using Chlamydomonas, which is proving to be a useful model for studying the actin 57 cvtoskeleton and its ciliary functions.

58 The actin cytoskeleton of Chlamydomonas contains two actin genes: IDA5, a 59 conventional actin with 91% sequence identity to human β -actin; and *NAP1*, a divergent actin 60 that shares only 63% of its sequence with human β -actin (Hirono et al. 2003; Kato-Minoura et al. 1998). We consider NAP1 to be an actin-like protein as opposed to an actin related protein 61 (ARP) because it has a higher sequency identity to actin than to conventional ARPs, and 62 63 because it is able to functionally compensate for the conventional filamentous actin (Jack et al. 64 2019; M. Onishi et al. 2018; M. Onishi, Pringle, and Cross 2016; Masayuki Onishi et al. 2019). Under normal, vegetative conditions, the conventional IDA5 is the primary actin expressed, but 65 66 when cells are treated with LatB. the LatB-insensitive NAP1 is upreculated (M. Onishi et al. 2018; M. Onishi, Pringle, and Cross 2016; Hirono et al. 2003). This separability of the two actins 67 68 has led to the discovery that they can compensate for each other in ciliary maintenance and 69 assembly (Jack et al. 2019). Studies of the role of actin in ciliary assembly have used global 70 disruption by knocking out either one of the filamentous actins or acutely knocking out both, yet 71 actin networks have diverse compositions and topologies that lead to specific subfunctions 72 within cells.

Actin networks rely on the actin binding proteins that contribute to the formation,
 arrangement, and function of the network. One such actin binding protein is the Arp2/3 complex,

75 which nucleates branched or dendritic actin networks by nucleating a daughter filament off the side of an existing mother filament. The dendritic networks nucleated by the Arp2/3 complex are 76 77 primarily found to be responsible for functions that involve membrane remodeling, for example 78 lamellipodia and endocytosis (Campellone and Welch 2010). The Arp2/3 complex from most 79 eukarvotes consists of seven subunits: Arp2, Arp3, and ARPC1-5 (Supplemental Figure 1) and 80 each subunit plays a specific role of varying importance in the nucleation process. ARPC2 and ARPC4 form the core of the complex and the primary contacts with the mother filament, Arp2 81 82 and Arp3 serve as the first subunits of the daughter filament, and ARPC1 and ARPC3 play a 83 role in nucleation but are not critical for branch formation (Robinson et al. 2001; Gournier et al. 84 2001). Each of these subunits are found in Chlamydomonas but have varying degrees of 85 sequence homology compared with conventional Arp2/3 complexes (Supplemental Figure 1). Interestingly, the ARPC5 subunit has yet to be found in *Chlamydomonas*. ARPC5 is thought to 86 87 be important for the association of ARPC1 to the complex, but a mammalian complex lacking 88 ARPC5 and ARPC1 maintains some nucleating and branching activity and is able to cross-link 89 actin normally (Gournier et al. 2001). 90 Here, using the chemical inhibitor CK-666 to inhibit the nucleating function of the Arp2/3

90 Here, using the chemical inhibitor CK-666 to inhibit the nucleating function of the Arp2/3 91 complex (Hetrick et al. 2013) and a genetic mutant of a critical Arp2/3 complex member, ARPC4 92 (Cheng et al. 2017; Li et al. 2019), we take a more delicate approach to investigating the role of 93 actin in ciliary assembly by separating different actin networks into their subfunctions based on 94 topology. Specifically, we probe the involvement of actin networks nucleated by the Arp2/3 95 complex in ciliary maintenance and assembly. This approach in these cells has allowed us to 96 propose a new model implicating a subset of filamentous actin in redistribution of membrane 97 and proteins for the initial stages of ciliogenesis.

98 99

100 **RESULTS**

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102 Loss of Arp2/3 complex function inhibits normal regeneration and maintenance of cilia:

103 To answer questions involving the role of Arp2/3 complex-mediated actin networks in ciliary assembly, we primarily used two tools. First, we used the chemical inhibitor CK-666 104 105 which blocks the nucleating ability of the Arp2/3 complex. Second, we obtained a null mutant of 106 the critical Arp2/3 complex member, ARPC4 (Cheng et al. 2017; Li et al. 2019) from the 107 Chlamydomonas Resource Center. This arpc4 mutant was confirmed via PCR (Supplemental **Figure 2A**). We further evaluated this mutant by creating a genetic rescue where a V5-tagged 108 109 ARPC4 construct is expressed in *arpc4* mutant cells. *arpc4*:ARPC4-V5. This was confirmed via 110 PCR, western blot, and immunofluorescence (Supplemental Figure 2).

111 We first investigated the requirement for the Arp2/3 complex in maintenance of cilia by 112 treating cells with varying concentrations of CK-666 or the inactive control CK-689 (250uM) for 2 113 hours and measuring the effect on steady state ciliary length. Consistent with previous results (Avasthi et al. 2014), we found that treating cells with CK-666 decreased ciliary length, 114 115 suggesting that the Arp2/3 complex is required for maintaining cilia (Figure 1A). We saw no 116 changes in ciliary length with the inactive CK-689 (Figure 1A) or when arpc4 mutant cells lacking a functional Arp2/3 complex were treated with CK-666 (Supplemental Figure 3-4). 117 118 Untreated arpc4 mutant cells did however recapitulate the CK-666 result by showing a 119 decreased ciliary length when compared with wild-type cells (Figure 1B). This defect in ciliary 120 length was not present in the rescue, arpc4:ARPC4-V5 (Figure 1B). Overall, these results 121 demonstrate through both chemical and genetic perturbation that the Arp2/3 complex is required 122 for normal ciliary length maintenance. 123 Next, we probed the involvement of Arp2/3 complex-nucleated actin in the more

124 complicated process of ciliary assembly where there is a high demand for protein and

membrane both from pools already existing in the cell and from synthesis (Wingfield et al. 2017;

126 Nachury, Seeley, and Jin 2010; Rohatgi and Snell 2010; Jack et al. 2019; Diener, Lupetti, and Rosenbaum 2015). Cells were deciliated by low pH shock and then allowed to synchronously 127 128 regenerate cilia after being returned to normal pH (Paul A. Lefebvre 1995). We found that cells 129 lacking a functional Arp2/3 complex were slow to regenerate their cilia, and two-thirds of cells did not regrow cilia at all (Figure 1C). This phenotype could be rescued by expression of 130 131 ARPC4-V5 in the *arpc4* mutant (Figure 1D). Importantly, the most severe defect in assembly 132 appeared to be in the initial steps when existing protein and membrane are being incorporated 133 into cilia.

134 The striking decrease in ciliary assembly is puzzling because the loss of Arp2/3 complex 135 function, and therefore only a subset of actin filaments, results in a more dramatic phenotype 136 than that of the nap1 mutants treated with LatB, which are lacking all filamentous actins (Jack et al. 2019). However, in the arpc4 mutant cells, a functional Arp2/3 complex never exists, and 137 138 therefore, cells never have Arp2/3 complex-mediated actin networks. In nap1 mutant cells 139 treated with LatB, the treatment begins shortly after deciliation resulting in an acute perturbation. 140 Further, LatB functions by sequestering actin monomers to promote filament disassembly, and 141 thus the effects may not be immediate (Spector et al. 1989). Therefore, it is likely that there is a 142 brief window where actin filaments can assert their initial role in ciliary regeneration before being 143 depolymerized. To avoid this, we began the LatB treatment in *nap1* mutants 30 minutes before 144 deciliation. This pre-treatment allows us to observe what happens when actin is not present 145 immediately after deciliation. (Figure 1E-F). In this case, we see slightly decreased ciliary length 146 consistent with the acute treatment but dramatically decreased percent ciliation, which is 147 consistent with the arpc4 mutant results.

148 This can also be observed with the inhibitor of the Arp2/3 complex, CK-666. In cells 149 treated with CK-666 immediately following deciliation, there is likely a window where the Arp2/3 150 complex can assert its role in assembly before being inhibited by CK-666. By pre-treating cells 151 with CK-666 for 1 hour before deciliation, we are able to observe what happens in the absence 152 of Arp2/3 complex function immediately following deciliation. When we do so, we see a more 153 dramatic defect in both ciliary length and percent ciliation than we do with just acute CK-666 154 treatment (Figure 1G-H), suggesting that the Arp2/3 complex is required for some very early 155 initial step of ciliary assembly that occurs even before we have a chance to treat the cells. 156

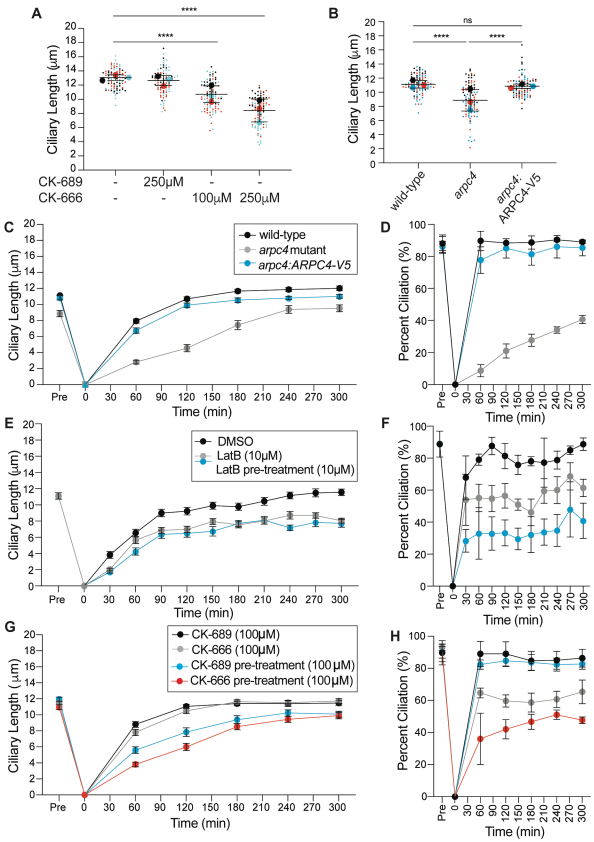




Figure 1. The Arp2/3 complex is required for normal ciliary maintenance and assembly. A) Wild-type cells containing a full active Arp2/3 complex were treated with 100µM or 250µM CK-666 or the inactive CK-689 for 2 hours.

160 Cells were then imaged using a DIC microscope and cilia were measure in ImageJ. Superplots are used to show the 161 mean of 3 separate experiments with error bars representing standard deviation. n=30 for each treatment in 3 separate 162 experiments. P<0.0001. B) Wild-type cells, arpc4 mutant cells, and arpc4 mutant cells expressing ARPC4-V5 steady 163 state cilia were also measured with no treatment. Superplots are used to show the mean of 3 separate experiments 164 with error bars representing standard deviation. n=30 for each strain for 3 separate experiments. P<0.0001. C) Wild-165 type cells and arpc4 mutant cells were deciliated using a pH shock and then cilia were allowed to regrow. The black 166 line represents wild-type, while the grey line represents the arpc4 mutant. The numbers above or below each point 167 show the percent ciliation for the wild-type and arpc4 mutant cells respectively. Means are displayed with error bars 168 representing 95% confidence interval. n=30 for each strain and each time point in 3 separate experiments. For every 169 time point except 0 min, P<0.0001 in terms of both length and percent ciliation. D) Wild-type cells, arpc4 mutant cells, 170 and arpc4 mutant cells expressing ARPC4-V5 were deciliated using a pH shock and then allowed to regrow. The black 171 line represents wild-type, while the grey line represents the arpc4 mutant and the cyan line represent the arpc4 mutant 172 expressing ARPC4-V5. Means are displayed with error bars representing 95% confidence interval. n=30 for each strain 173 and each time point in 3 separate experiments. E) nap1 mutant cells were pre-treated with 10µM LatB for 30 minutes 174 before deciliation or treated with LatB upon the return to neutral pH following deciliation. The black line represents 175 untreated cells, while the light grey line represents cells treated with LatB following deciliation and the dark grey line 176 represents cells pre-treated with LatB. Error bars represent 95% confidence interval. n=30 for 3 separate experiments. 177 For every time point P>0.0001 between DMSO and treated samples, except 30min (10µM LatB) which is ns. F) Percent 178 179 ciliation for the experiment in D. Line color is the same as D. Error bars represent standard deviation. G) Wild-type cells were pre-treated with CK-666 or the inactive CK-689 (100µM) for 1 hour befoe deciliation of treated with CK-666 or the 180 inactive CK-689 (100µM) following deciliation. Error bars represent 95% confidence interval. n=30 for 3 separate 181 experiments. H) Percent ciliation for the experiments in G. Error bars represent standard deviation. n=100 in 3 separate 182 experiments. 183

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185The Arp2/3 complex is required for the incorporation of existing membrane and proteins186for ciliary assembly:

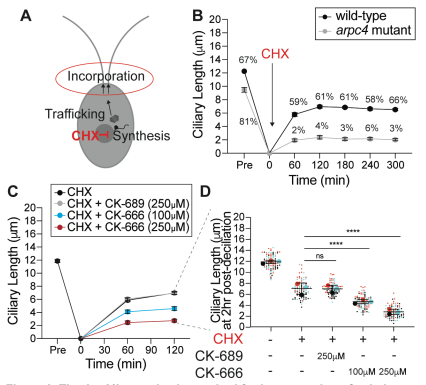
187 There are several distinct, filamentous actin-dependent steps of ciliary assembly after 188 severing, including the incorporation of existing protein and membrane and the synthesis of new protein for cilia. Using a method that labels nascent peptides, we found that loss of ARPC4 did 189 190 not prevent upregulation of translation following deciliation (Supplemental Figure 5). In this 191 experiment, we are halting translation and fluorescently labelling newly translated polypeptides. 192 Wild-type and *arpc4* mutant cells were tested using this reaction either before deciliation. 193 following deciliation and one hour of regrowth, or following deciliation and one hour of regrowth 194 in cycloheximide (CHX), which blocks protein synthesis by blocking the elongation step of 195 protein translation. Both wild-type and arpc4 mutant cells displayed an increase in cell 196 fluorescence, especially in the area around the nucleus, following deciliation, indicating an 197 increase in protein synthesis following deciliation (Supplemental Figure 5). Importantly, this 198 increase in cell fluorescence was not significantly different between wild-type and arpc4 mutant 199 cells, suggesting that the loss of Arp2/3 complex function does not prevent the upregulation of 200 protein synthesis that follows deciliation. This also suggests that the cells are aware that the 201 cilia have been severed, as they respond with increased protein synthesis.

202 Given that arpc4 mutant cells respond to deciliation with protein synthesis, another 203 possibility for the role of the Arp2/3 complex in ciliary assembly involves the first step, which 204 requires that a pool of existing proteins and membrane are incorporated into cilia in an actin-205 dependent manner (Jack et al. 2019). Further, disruption of Arp2/3 complex-mediated actin 206 networks results in slow initial ciliary assembly, when it is likely that existing protein is being 207 incorporated. We tested this by treating cells with cycloheximide (CHX), a protein synthesis 208 inhibitor, we can eliminate the contribution of two steps in the process of ciliary assembly (Figure 2A, Supplemental Figure 4) (Rosenbaum, Moulder, and Ringo 1969). Without protein 209 210 synthesis, there is no trafficking or incorporation of new proteins. Therefore, any ciliary growth 211 we see is due to the incorporation of the existing protein alone. Under normal conditions, cells 212 that are deciliated and treated with cycloheximide typically grow cilia to about half-length, or 6µm, within 2 hours (Figure 2B). In the arpc4 mutant strain treated with CK-666, cilia display 213 214 minimal growth (Figure 2B). In fact, throughout a five-hour period, only 6% of cells were able to 215 form cilia at all (Figure 2B). This suggests that the Arp2/3 complex and the actin networks

216 nucleated by the complex are indispensable for the incorporation of existing protein and 217 membrane during ciliary assembly.

218 We suspected that the *arpc4* mutant cells either lacked the normal pool of ciliary 219 precursor proteins or were unable to incorporate it. However, the inability of the genetic mutants 220 to regenerate in cycloheximide prevents us from being able to do the typical studies testing new 221 protein synthesis, precursor pool size, and new protein incorporation outlined in Jack et al. 2018 222 as they all require regeneration in cycloheximide. To get around this, we used an acute 223 perturbation through chemical inhibition in wild-type cells that have a normal ciliary precursor 224 pool (as evidenced by their ability to grow to half-length in cycloheximide). These cells were 225 deciliated and then CK-666 was added (in addition to cycloheximide) only for the regrowth, and thus it was not able to affect the size of the precursor pool. Cells treated with CK-666 and 226 227 cycloheximide could not incorporate the precursor pool we know exists in these wild-type cells 228 into cilia, while cilia of cells treated with only cycloheximide or cycloheximide and the inactive 229 control for CK-666, CK-689 were able to grow to half length (Figure 2C-D, Supplemental 230 Figure 4). This suggests that the problem with incorporation we see in cells lacking a functional 231 Arp2/3 complex lies outside of the availability of the precursor pool.

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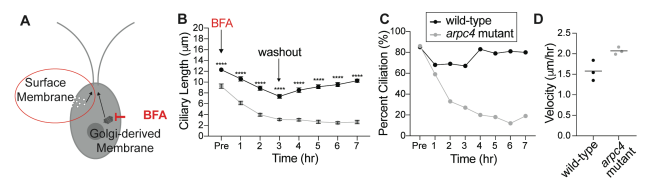
Figure 2. The Arp2/3 complex is required for incorporation of existing protein during ciliary assembly. A) Treating cells with cycloheximide inhibits protein synthesis, which means only incorporation of existing protein into the 236 237 238 239 cilia is observed. B) Wild-type cells and arpc4 mutants were deciliated and then allowed to regrow in 10µM CHX. The percentages above the lines represent the percent of cells with cilia at the indicated time points. The mean is shown with error bars representing 95% confidence interval. n=30 for each strain and each time point in 3 separate experiments. For every time point besides 0 min, P<0.0001 for both length and percent ciliation. C) Wild-type cells 240 were deciliated and then treated with a combination of 10µM cycloheximide (CHX) and CK-666 (100µM or 250µM) or 241 CK-689 (the inactive control, 250µM) at the same concentration during regrowth. The mean is shown with error bars 242 representing 95% confidence interval. n=30 for each strain and each time point in 3 separate experiments. At both 1 243 and 2 hour time points P<0.0001 for cells treated with CK-666 compared to wild-type cells, and ns for cells treated 244 with CK-689 compared to wild-type cells. D) The graph shows the 2 hour time point from C, or the length of their cilia 245 after 2 hours of treatment and regrowth. Superplots are used to show the mean of 3 separate experiments with error

bars representing standard deviation. n=30 for each treatment group 3 separate experiments. For both the 100μM
 and 250μM CK-666 treatments with CHX, P<0.0001.

249250 Cilia of arpc4 mutant cells resorb faster in the absence of the Golgi:

251 Because we see defects in ciliary assembly and maintenance when cells are likely 252 incorporating existing protein, and we know the protein needed for assembly is in excess due to 253 our acute perturbations with CK-666, we next investigated membrane delivery to cilia. This is of 254 particular interest as the Arp2/3 complex is canonically thought to be involved in membrane 255 remodeling functions. Typically, the Golgi is thought to be the main source of membrane for cilia 256 (Nachury, Seeley, and Jin 2010; Rohatgi and Snell 2010), and both ciliary membrane, 257 membrane proteins, and even axonemal proteins are transported in or attached to vesicles in 258 cytosol (Wood and Rosenbaum 2014). In Chlamydomonas, this has been demonstrated by the 259 ciliary shortening of cells treated with Brefeldin A (BFA), a drug that causes Golgi collapse by 260 interfering with ER to Golgi transport (W. Dentler 2013). To determine if the Arp2/3 complex is involved in the trafficking of new protein from the Golgi to cilia, we examined the Golgi following 261 262 deciliation using transmission electron microscopy (TEM) in arpc4 mutants (Supplemental Figure 6A). The Golgi appeared grossly normal, and in all cases had approximately the same 263 264 number of cisternae (Supplemental Figure 6A-B) and did not show an abnormal accumulation 265 of post-Golgi membrane as previously reported when perturbing all filamentous actin (Jack et al., 2019). 266

267 Alternative pathways for delivery of material to the cilia have also been found in Chlamydomonas. For example, surface proteins were biotinylated and then cells were 268 269 deciliated, meaning the membrane and proteins within cilia were lost. When cilia were allowed 270 to regrow, biotinylated proteins were found to reside within the new cilia suggesting they came from the plasma membrane (W. Dentler 2013). Therefore, we hypothesized that due to its role 271 272 in membrane remodeling, and particularly endocytosis, in other organisms, the Arp2/3 complex 273 may be part of an endocytic pathway that provides membrane and perhaps membrane proteins 274 to cilia (Figure 3A). To test if membrane could be coming from an endosomal or endocytic 275 source other than the Golgi, we treated cells with 36µM BFA to collapse the Golgi and block 276 exocytosis so cells would be forced to utilize other sources of ciliary proteins and membranes. 277 Wild-type cilia treated with BFA resorb slowly, but arpc4 mutant cells had a faster resorption rate (Figure 3B and D, Supplemental Figure 4). Further, the number of cells with cilia in the 278 279 arpc4 mutant cells dramatically decreased with BFA treatment (Figure 3C). Meanwhile, cells 280 treated with other known ciliary resorption-inducing drugs that do not specifically target Golgi 281 traffic, 3-isobutyl-1-methylxanthine (IBMX) (Pasquale and Goodenough 1987) or sodium 282 pyrophosphate (NaPPi) (P. A. Lefebvre et al. 1978) show an increased velocity of resorption in 283 the wild-type cells compared to the *arpc4* mutant cells (**Supplemental Figure 7**), suggesting the 284 faster resorption of the *arpc4* mutant cells in BFA is specific to the effects of BFA on the cell. 285 Thus, wild-type cells are more capable of maintaining cilia without membrane supply from the 286 Golgi, suggesting that there must be another source for membrane that is dependent upon the 287 Arp2/3 complex.



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Figure 3. The Arp2/3 complex is required for ciliary maintenance in the absence of intact Golgi. A) Treating cells with Brefeldin A (BFA) causes the Golgi to collapse meaning any membranes and proteins used to maintain the cilia must come from other sources. B) Cells were treated with 36µM BFA for 3 hours at which time the drug was washed out. Wild-type is represented by black, while arpc4 mutants are grey. The mean is shown with error bars representing 95% confidence interval. Error bars represent 95% confidence interval of the mean. n=30 for each time point and each strain in 3 separate experiments. **** represents P<0.0001. C) Percent ciliation of the cells in B. n=100. D) Resorption speed for wild-type cells and arpc4 mutant cells as determined by fitting a line to the first 4 time points before washout 297 and determining the slope of the line. Line represents the mean of 3 separate experiments. N=3. P=0.0314 298

300 Apical actin dots are dependent on the Arp2/3 complex:

301 Since ciliary membrane proteins can come from the Golgi or the plasma membrane and 302 arpc4 mutant cells have a more severe defect in incorporating ciliary proteins from non-Golgi 303 sources, we asked if Arp2/3 complex-mediated actin networks might be responsible for 304 endocytosis from the plasma membrane in Chlamydomonas as it is in other organisms. To 305 determine where in the cell Arp2/3 complex-mediated actin networks might be acting, we looked 306 directly at the effects of loss of Arp2/3 complex function on the actin structures in the cell. Using 307 new protocols for the visualization of actin in Chlamydomonas developed by our lab (Craig et al. 308 2019), we stained wild-type cells and *arpc4* mutant cells with fluorescent phalloidin. In wild-type 309 cells, apical dots reminiscent of endocytic actin patches in yeast are typically seen near the 310 base of cilia (Figure 4A). We quantified the presence of these dots in the wild-type cells 311 compared to the *arpc4* mutant cells (Figure 4A-B). We found that while about 70% of wild-type 312 cells contain the dots, only about 5% of the *arpc4* mutant cells had dots (Figure 4B), suggesting 313 the Arp2/3 complex is required for the formation of this actin structure. This phenotype was 314 rescued by the expression of the ARPC4-V5 construct in the *arpc4* mutant cells (Figure 4C). 315 The reliance of this structure on the Arp2/3 complex. led us to further question whether these 316 dots could represent endocytic membrane remodeling.

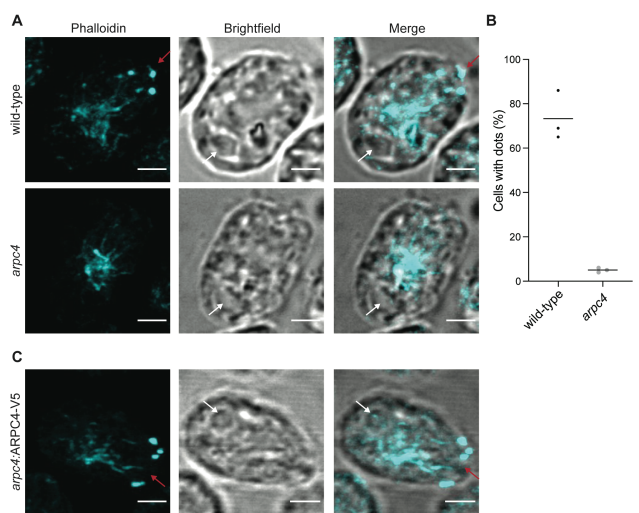


Figure 4. Loss of a functional Arp2/3 complex results in changes in actin distribution. A) Wild-type and *arpc4* mutant cells stained with phalloidin to visualize the actin network along with brightfield to show cell orientation. Images were taken as a z-stack using airsycan imaging and are shown as a maximum intensity projection. Red arrow is pointing to dots at the apex of the cell, and white arrow is pointing to the pyrenoid near the basal end of the cell. Scale bars represent 2µm. B) Percentage of cells with apical dots as shown in A. Percentages taken from 3 separate experiments where n=100. Line represents the mean. P<0.0001. C) Presence of apical dots in the *arpc4* mutant rescue expressing ARPC4-V5. Images were taken as a z-stack using airsycan imaging and are shown as a maximum intensity projection. Red arrow is pointing to dots at the apex of the cell, and white arrow is pointing to the pyrenoid near the basal end of the cell. Scale bars represent the mean. P<0.0001. C) Presence of apical dots in the *arpc4* mutant rescue expressing ARPC4-V5. Images were taken as a z-stack using airsycan imaging and are shown as a maximum intensity projection. Red arrow is pointing to dots at the apex of the cell, and white arrow is pointing to the pyrenoid near the basal end of the cell. Scale bars represent 2µm.

30 Endocytosis in Chlamydomonas is likely clathrin-dependent:

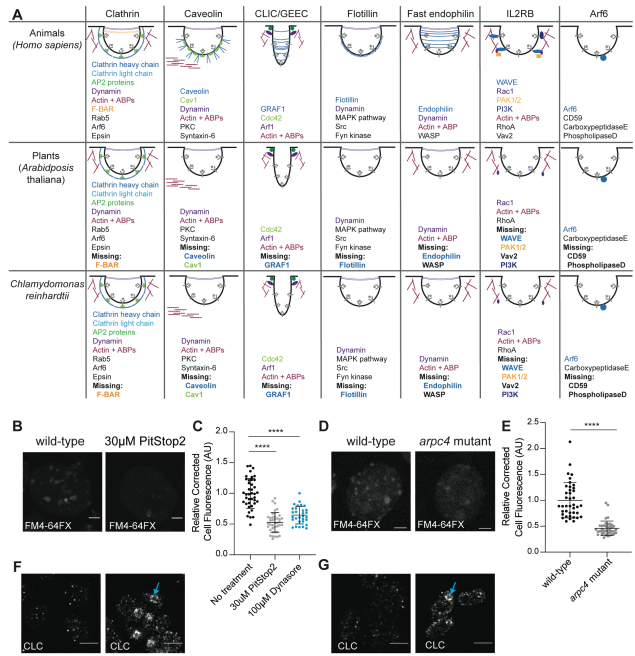
The Arp2/3 complex is conventionally thought to be involved in endocytosis in cell-walled veast to overcome turgor pressure (Aghamohammadzadeh and Ayscough 2009; Basu, Munteanu, and Chang 2014; Carlsson and Bayly 2014). Chlamydomonas cells also have a cell wall and since the apical actin dots resemble these endocytic pits (Goode, Eskin, and Wendland 2015; Adams and Pringle 1984; Ayscough et al. 1997), we hypothesized that Arp2/3 complex and actin-dependent endocytosis might be occurring in Chlamydomonas even though this process has not yet been directly demonstrated in this organism. To determine what kind of endocytosis was likely occurring in these cells, we compared the endocytosis-related proteins found in mammals and plants to those in *Chlamydomonas* (Figure 5A). We found that Chlamydomonas lacks much of the important machinery for almost all typical endocytosis processes, including caveolin for caveolin-mediated endocytosis, flotillin for flotillin-dependent

endocytosis, and endophilin for endophilin-dependent endocytosis (Figure 5A). However, most
 of the canonical clathrin-related endocytosis machinery could be found in *Chlamydomonas*, and
 thus, clathrin-mediated endocytosis is conserved to a higher extent than other endocytic
 mechanisms.

346 We aimed to further probe the likelihood of clathrin-mediated endocytosis occurring in 347 Chlamydomonas. However, a mutant for the proteins involved in clathrin-mediated endocytosis 348 does not currently exist and methods of targeted mutagenesis in Chlamydomonas are not yet 349 reliable. Therefore, we turned to our best alternative PitStop2, which inhibits the interaction of 350 adaptor proteins with clathrin, halting clathrin endocytosis, despite the reported off-target effects 351 on global endocytosis in mammalian cells (Willox, Sahraoui, and Royle 2014) (Supplemental 352 **Figure 4**). Additionally, we used the dynamin inhibitor Dynasore, which is also thought to block endocytosis by inhibiting the GTPase activity of dynamin (Macia et al. 2006). For this 353 experiment, we used the fixable lipophilic dye FM 4-64FX (Cochilla, Angleson, and Betz 1999; 354 355 Gachet and Hyams 2005) (Thermo Scientific). This dye is impermeable to the plasma 356 membrane but is usually quickly endocytosed into cells showing bright foci where dye is 357 enriched in endocytosed compartments. Thus, we incubated the dye for only 1 minute to allow 358 enough time for internalization into endosomes but not enough for incorporation into various 359 cellular membrane structures. The ability of PitStop2-treated cells to internalize membrane was 360 measured by calculating the total cell fluorescence inside the cell after allowing dye to be 361 internalized (Figure 5B). We found that cells treated with 30µM PitStop2 or 100µM Dynasore have significantly decreased membrane internalization (Figure 5C), which further supports the 362 363 idea that endocytosis of some kind is occurring in these cells and that it is likely clathrin-364 mediated.

Next, we tested whether the endocytosis is Arp2/3 complex-dependent by using this membrane internalization assay on *arpc4* mutant cells compared to wild-type cells. We found that cells lacking a functional Arp2/3 complex have decreased total cell fluorescence (**Figure 5D-E**) suggesting the endocytosis in *Chlamydomonas* is Arp2/3 complex-dependent.

369 To better demonstrate the relationship between the Arp2/3 complex and endocytosis, we 370 used a clathrin light chain antibody to stain cells. In both PitStop2 treated cells and arpc4 mutant 371 cells, but not in untreated wild-type cells, we see a mislocalization of clathrin staining around the 372 pyrenoid (Supplemental Figure 8, Figure 5F-G). Although the reason for this accumulation of 373 clathrin around the pyrenoid is not clear, the interesting takeaway from this data is that 374 disruption of either endocytosis with PitStop2 or of Arp2/3 function results in defects in 375 membrane internalization and clathrin localization. These data support a role for the Arp2/3 376 complex in endocytosis in Chlamydomonas. 377



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378 379 Figure 5. Arp2/3 complex-dependent endocytosis is conserved in Chlamydomonas. A) Gene presence was determined using BLAST. Word colors correspond to diagram colors. B) Cells treated with 30µM PitStop2 were 381 incubated with FM4-64FX and imaged on a spinning disk confocal. Max intensity projections of z-stacks are shown. 382 Scale bars are 2µm, C) The background corrected fluorescence for each sample, including cells treated with 100µM 383 Dynasore. The mean is shown with error bars showing standard deviation. n=30 in 3 separate experiments. P<0.0001. 384 D) Wild-type and arpc4 mutant cells treated with FM4-64FX and imaged on a spinning disk confocal. Max intensity 385 projections of z-stacks are shown. Scale bars are 2µm. E) The background corrected fluorescence for each sample. 386 The mean is shown with error bars representing standard deviation. n=30 in 3 separate experiments. P<0.0001. F) 387 Wild-type and PitStop2 treated cells, and arpc4 mutant cells were stained with clathrin light chain antibody and imaged 388 using a spinning disk confocal. Cyan arrows point to accumulation around the pyrenoid. Scale bar represents 5µm. G) 389 Wild-type and arpc4 mutant cells were stained with clathrin light chain antibody and imaged using a spinning disk 390 confocal. Cyan arrows point to accumulation around the pyrenoid. Scale bar represents 5µm.

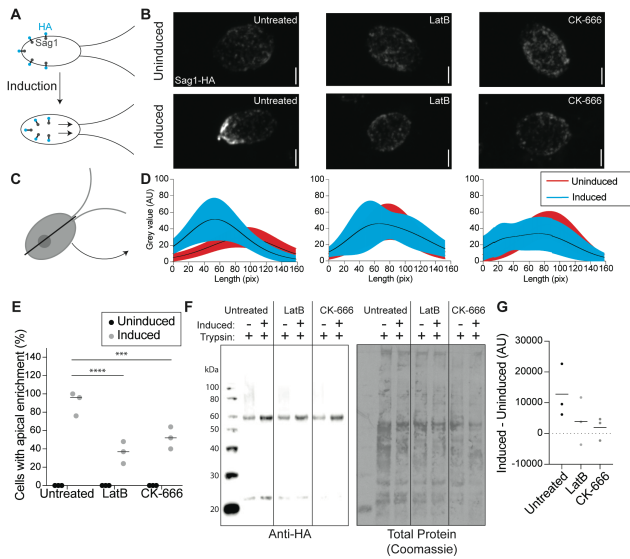
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The Arp2/3 complex is required for the internalization and relocalization of a membrane protein from the periphery of the cell to cilia:

395 Upon finding that there is likely Arp2/3 complex-dependent clathrin-mediated 396 endocytosis in Chlamydomonas, we next asked if this endocytosis could be responsible for the 397 relocalization and internalization of a known ciliary protein. SAG1 is a membrane protein that is 398 important for mating in Chlamydomonas cells (Belzile et al. 2013; Ranjan, Awasthi, and Snell 399 2019). When cells are induced for mating with dibutyryl-cAMP (db-cAMP), SAG1 must relocalize 400 from the cell periphery to cilia, where it facilitates ciliary adhesion between mating cells. This 401 relocalization of SAG1 is thought to occur through internalization of the protein followed by 402 internal trafficking on microtubules to the base of cilia (Belzile et al. 2013; Ranjan, Awasthi, and 403 Snell 2019).

We examined whether actin and the Arp2/3 complex were required for the transport of 404 405 HA-tagged SAG1 to the apex of the cell and cilia for agglutination during mating (Figure 6A). 406 Using immunofluorescence, we observed cells treated with either 10µM LatB to depolymerize 407 IDA5 or 250µM CK-666 to perturb the Arp2/3 complex (Figure 6, Supplemental Figure 4). 408 Before induction, SAG1-HA localized to the periphery of the cell (Figure 6B, top). 30 minutes 409 after induction with db-cAMP, SAG1-HA relocalized to the apex of the cell and to cilia in 410 untreated cells (Figure 6B, left). In both LatB and CK-666 treated cells, this apical enrichment 411 was greatly decreased (Figure 6B, middle and right). To quantify this, line scans were drawn 412 through the cell from the apex to the basal region (Figure 6C-D). The percentage of cells with 413 apical enrichment was calculated, and it was found that untreated cells had a higher percent of 414 apical enrichment when compared with LatB or CK-666 treated cells (Figure 6E). Thus, cells 415 with perturbed Arp2/3 complex-mediated filamentous actin show decreased efficiency of SAG1-416 HA relocalization.

We next asked if this decrease in relocalization in cells with actin and Arp2/3 complex 417 418 inhibition could be due to a decrease in the internalization of SAG1-HA through a process that 419 seems to require endocytosis. To investigate this, we used a method first described by Belzile et 420 al. 2013, where cells were induced and treated with a low percentage (0.01%) of trypsin, which 421 will hydrolyze exterior proteins but cannot enter the cell. In untreated cells, we see an increase 422 in SAG1-HA protein levels following induction because SAG1-HA is internalized and becomes 423 protected from trypsin (Figure 6F). In cells treated with either 10µM LatB or 250µM CK-666 we 424 see a decrease in this trypsin protection as shown in the western blot (Figure 6F). This was 425 further quantified by subtracting the amount of protein before induction from the amount of 426 protein present after induction, which gives a value representing the amount of SAG1-HA 427 protected from trypsin due to internalization in these cells (Figure 6G). The decrease in SAG1-428 HA following induction in cells with decreased filamentous actin and Arp2/3 complex function 429 indicates a role for Arp2/3 complex-mediated actin networks in internalization of this specific 430 ciliary membrane protein.



432 433 Figure 6. The Arp2/3 complex is required for the relocalization and internalization of the ciliary protein SAG1 434 for mating. A) When mating is induced SAG1-HA is internalized and relocalized to the apex of the cells and cilia for 435 agglutination. B) Maximum intensity projections of z-stacks taken using spinning disk confocal microscopy of SAG1-436 HA. Left is untreated, middle is treated with 10µM LatB, and right is treated with 250µM CK-666. Top row of images 437 are uninduced and bottom row of images are induced with db-cAMP. Scale bar represents 2um. C) Diagram 438 representing line scans taken through the cells in z-stack sum images. D) Line scans were taken from the apex of the 439 cell to the basal region of the cell in untreated cells (left), LatB treated cells (middle), and CK-666 (right). Lines scans 440 were normalized and fit with a gaussian curve. The curves were averaged. Black lines represent mean and then shaded 441 regions represent standard deviation. Grey represents uninduced samples, green represent induced samples. 0 on the 442 v-axis represents the apical region of the cell. n=30 from a single representative experiment, E) Percentage of cells 443 with apical enrichment based on E for uninduced (black) and induced (grey) cells for each treatment group. The mean 444 is shown with error bars representing standard deviation. n=30 for 3 separate experiments for each treatment. F) 445 Western blot showing amount of SAG1-HA in uninduced and induced cells in each treatment group all treated with 446 0.01% trypsin. G) Intensity of the bands in H were normalized to the total protein as determined by amido black staining 447 and quantified in ImageJ was used to subtract uninduced from induced to give a representation of the amount of SAG1-448 HA internalized with induction. Line represents mean of 3 separate experiments.

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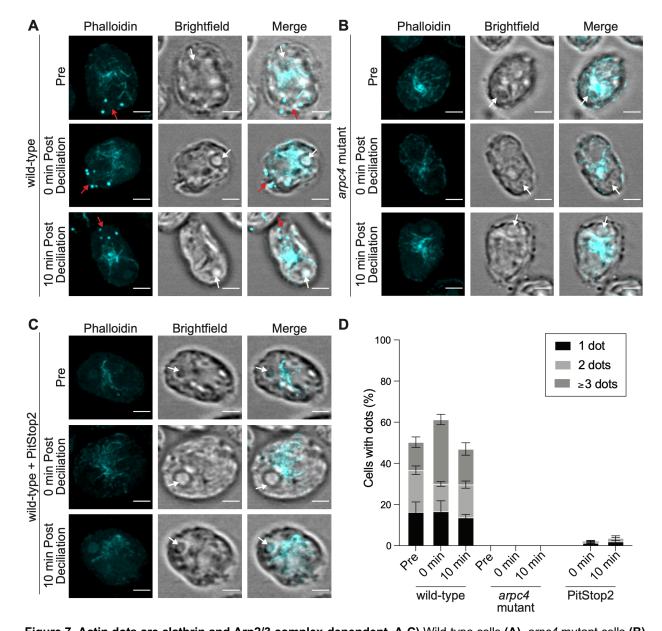
451 Actin dots increase in an Arp2/3 complex and clathrin-dependent manner following

452 deciliation:

453 Having established that the Arp2/3 complex is required for ciliary assembly, membrane 454 dye internalization, and the endocytosis of a known ciliary protein, we wondered if these 455 functions could be connected given that arpc4 mutant cells have defects in maintaining cilia 456 from non-Golgi sources (Figure 3). Therefore, we returned to the Arp2/3 complex-dependent 457 actin dots seen in wild-type cells that are reminiscent of endocytic pits in yeast. Because ciliary 458 membrane and proteins can come from the plasma membrane (Dentler, 2013), we suspected 459 there would be an increase in these actin dots immediately following deciliation. We used phalloidin to visualize the actin cytoskeleton of wild-type cells before and immediately following 460 461 deciliation, as well as 10 minutes later (Figure 7A). We saw an increase in both the percentage 462 of cells with dots and the number of dots per cell immediately following deciliation that then 463 returned to normal by 10 minutes (Figure 7A and D). This is consistent with the results shown in Figure 1E-F and confirms that the defect seen in ciliary assembly is due to an event 464 465 occurring very early in ciliary assembly, even within the first few minutes after deciliation.

We wondered if this increase in dots would result in dots in the *arpc4* mutant cells which have almost not dots normally. We found that in the *arpc4* mutants dots were never observed, before or after deciliation (**Figure 7B and D**), suggesting these dots are dependent on the Arp2/3 complex. Next, we wanted to see if the dots really were due to clathrin-mediated endocytosis, so we treated cells with PitStop2 and looked for this same increase in dots. This treatment almost fully blocked the appearance of dots following deciliation and completely eliminated the presence of cells with 3 or more dots (**Figure 7C-D**), suggesting a clathrin-

473 dependent mechanism, as well as an Arp2/3-dependent mechanism, is related to these dots.



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Figure 7. Actin dots are clathrin and Arp2/3 complex-dependent. A-C) Wild-type cells (A), arpc4 mutant cells (B), and wild-type cells treated with 30 µM PitStop2 (C) stained with phalloidin to visualize the actin network before deciliation, immediately following deciliation, and 10 minutes following deciliation. Brightfield images are to visualize cell orientation. Images were taken as a z-stack using airyscan imaging and are shown as a maximum intensity projection. Scale bar represents 2µm. Red arrows point to dots at the apex of the cell, and white arrows point to the 481 pyrenoid at the opposite end of the cell. D) The percentage of cells with 1 dot, 2 dot, or 3 dots in each condition. 482 Quantification based on sum slices of z-stacks taken using a spinning disk confocal. n=100 in 3 separate experiments. 483 For wild-type, the total number of cells with dots is significantly different for the 0 min time point (**) and the number of 484 dotted cells with 3 or more dots is significantly different for the 0 time point (****).

485 486

487 Ciliary membrane proteins follow different paths from the plasma membrane to the cilia:

488 Finally, to specifically determine if ciliary membrane and therefore membrane proteins 489 were coming from a pool in the plasma membrane we did an experiment first described in W. 490 Dentler 2013. Surface proteins were biotinylated, then cells were deciliated. After the cilia

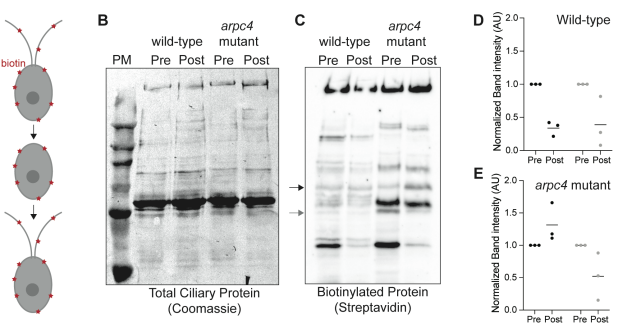
491 regrew, they were isolated and probed for biotinylated protein (Figure 8A). Any biotinylated

492 protein present in the newly grown and isolated cilia must have come from a pool in the plasma 493 membrane. While some proteins returned in both wild-type and *arpc4* mutant cells, some 494 appeared to a lesser degree in arpc4 mutant cells compared to wild-type cells (Figure 8B-E, 495 black arrow and black bars) and some returned to a higher degree in arpc4 mutant cells (Figure 8B-E, grev arrow and grev bars). Other biotinylated proteins found in wild-type cilia 496 497 were not found in the arpc4 mutant cilia before or after deciliation, so there is a mechanism for 498 delivery of proteins to the cilia from the plasma membrane that Arp2/3 is absolutely essential for 499 (Figure 8B-C). This suggests there are multiple paths to the ciliary membrane, some of which 500 are Arp2/3 complex-independent and some that are Arp2/3 complex-dependent. This may 501 represent lateral diffusion and endocytosis respectively.

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506 507 Figure 8. Ciliary membrane proteins have multiple paths from the plasma membrane. A) Cells were biotinylated, 508 deciliated, and then allowed to regrow before cilia were isolated and probed for biotinylated protein. B) Total protein in 509 wild-type and arpc4 mutant ciliary isolate investigated by western blot and Coomassie. C) Wild-type and arpc4 mutant 510 cells ciliary isolate was investigated by western blot and probed using streptavidin. Black arrow shows ciliary protein 511 present to a higher degree in wild-type cells than the arpc4 mutant cells. Grey arrows show ciliary protein that is present 512 to a higher degree in arpc4 mutant cells than in wild-type cells. D) Bands represented by black and grey arrows are 513 quantified for the wild-type cells. Data acquired from 3 separate experiments. E) Bands represented by black and grey 514 arrows are quantified for the arpc4 mutant cells. Data represented as the mean from 3 separate experiments. Error 515 bars represent standard deviation.

516 517

518 DISCUSSION

519 In this study, we investigate the Arp2/3 complex of Chlamydomonas reinhardtii that 520 functions to maintain and assemble cilia. This complex potentially lacks the ARPC5 subunit, 521 although it is possible that a highly divergent ARPC5 exists. In yeast, deletion of any of the 522 genes encoding Arp2/3 complex members causes severe defects and even lethality, but these 523 defects differ in severity depending on the complex members deleted, suggesting that complex 524 members have varying degrees of importance in Arp2/3 complex function (Winter, Choe, and Li 525 1999). The role of ARPC5 in actin nucleation is being investigated, but some groups have found 526 it unnecessary for overall function of the complex (Gournier et al. 2001; von Loeffelholz et al. 2020). Furthermore, our data show that knocking out function of the ARPC5-less 527

528 *Chlamydomonas* Arp2/3 complex genetically or chemically results in phenotypes in ciliary 529 assembly and maintenance, suggesting that the wild-type complex is active. Because the 530 Arp2/3 complex has known functions in membrane dynamics, this led us to pursue models of 531 Arp2/3 complex-dependent membrane trafficking to cilia.

Previously, three models for the trafficking of membrane proteins to cilia have been 532 533 proposed regarding where ciliary vesicles fuse relative to a diffusion barrier composed of 534 septins (Hu Qicong et al. 2010), which delineates ciliary membrane and cell body plasma 535 membrane (Nachury, Seeley, and Jin 2010). The first is that Golgi vesicles containing ciliary 536 proteins fuse with the ciliary membrane inside the cilium itself. Proteins, both membrane and soluble, have been found to be transferred from the Golgi to the cilia on or in cytoplasmic 537 538 vesicles (Wood and Rosenbaum 2014). Second, Golgi vesicles containing ciliary proteins fuse 539 outside but near the cilium still within the diffusion barrier (Papermaster, Schneider, and 540 Besharse 1985; Nachury et al. 2007; Zuo, Guo, and Lipschutz 2009). In Chlamydomonas, this 541 was first described for mastigoneme proteins, which were found to be transferred from the Golgi and then exocytosed for use on the exterior of the cell (Bouck 1971). In the third model, Golgi 542 543 vesicles containing proteins fuse with the plasma membrane outside the diffusion barrier where 544 they somehow move in the plane of the plasma membrane across this barrier, perhaps through 545 lateral diffusion that requires remodeling or passing through the diffusion barrier. Evidence for 546 this path was shown using Hedgehog signaling protein Smoothened, which was found to 547 relocalize in a dynamin-independent manner from the plasma membrane to the cilia 548 immediately after stimulation in pulse labeling studies (Milenkovic, Scott, and Rohatgi 2009).

549 Our data support a fourth model, likely occurring in concert with other models, in which 550 membrane proteins are recruited to the cilium from a reservoir in the cell body plasma 551 membrane. We find that immediately following deciliation the Arp2/3 complex is required for 552 ciliary assembly, clathrin-mediated endocytosis, and redistribution of ciliary proteins from the 553 plasma membrane (Figure 9A). We hypothesize that ciliary membrane proteins and membrane 554 targeted to the plasma membrane of the cell outside the diffusion barrier can be endocytosed 555 and trafficked to cilia, either within or outside of the diffusion barrier in an actin and Arp2/3 556 complex-dependent manner.

557 Although our data does not eliminate the possibility of Arp2/3 complex function in supply 558 of ciliary membrane and protein stored in other endosomal compartments, ciliary localization of 559 proteins initially labeled on the cell surface with biotin (Figure 8) suggests that some ciliary 560 membrane proteins incorporated during assembly are coming directly from the plasma membrane itself. An endocytic mechanism of trafficking in intracellular ciliogenesis has been 561 investigated previously in mammalian RPE1 cells. The ciliary pocket found at the base of 562 563 primary and motile cilia formed intracellularly has been found to be an endocytically active region (Molla-Herman et al. 2010) but clathrin-mediated endocytosis was not required for 564 565 ciliogenesis in those cells. The Bardet Biedl Syndrome complex (BBsome), which is involved in 566 regulation of ciliary membrane protein composition, has been shown to interact with clathrin 567 directly at the ciliary pocket to facilitate membrane sorting in trypanosomes (Langousis et al. 568 2016). Further, some BBsome complex members resemble coat proteins such as clathrin (Jin et al. 2010) suggesting a direct role for the this cilium regulatory complex in membrane budding 569 570 functions. It has also been found that disruption of recycling endosomes reduces the localization 571 of polycystin-2 to cilia, suggesting a role for recyclying endosomes in the localization of proteins 572 to the cilia (Monis, Faundez, and Pazour 2017). Even in *Chlamydomonas*, clathrin heavy chain 573 has been found to localize at the base of cilia (Kaplan et al. 2012). While the mechanism was 574 unknown, it has been shown that plasma membrane surface-exposed proteins are relocalized to 575 cilia during ciliary regeneration (W. Dentler 2013), a result we were able to recapitulate and 576 demonstrate depends, in part, upon the Arp2/3 complex.

577 Altogether, this leads us to hypothesize that the role of the Arp2/3 complex in ciliary 578 assembly is through endocytic recruitment from a ciliary protein reservoir in the plasma

579 membrane before newly synthesized protein and Golgi-derived membrane are capable of 580 supplying additional materials (**Figure 9B**). While this model provides a possible route that 581 some ciliary proteins and membranes take to the cilia, we believe this is one of several paths 582 that can be taken to the cilia. Trafficking to cilia is likely cargo- and time-dependent, and which 583 path proteins take may tell us the order and speed in which they populate the cilium for 584 subsequent function.

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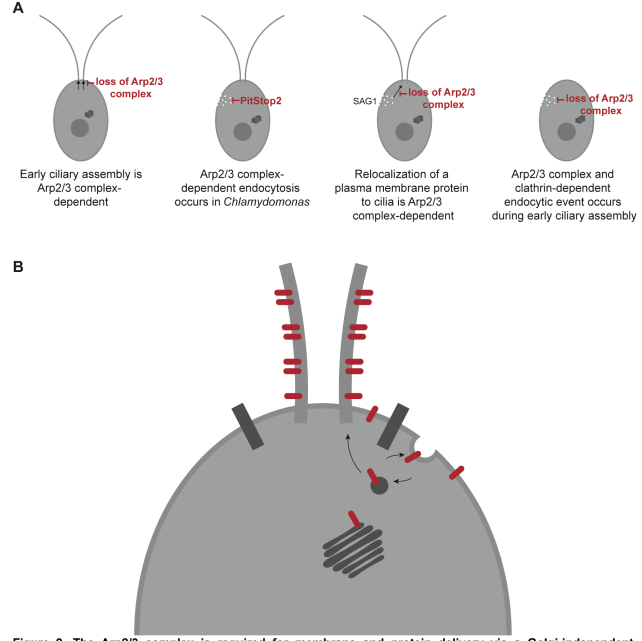


Figure 9. The Arp2/3 complex is required for membrane and protein delivery via a Golgi-independent, endocytosis-like process. A) Arp2/3-mediated actin networks are required for ciliary assembly in *Chlamydomonas* particularly during the initial stages. These actin networks are also required for clathrin-mediated endocytosis, and for the endocytosis-like relocalization of a ciliary protein from the plasma membrane to the cilia. Finally, a large endocytic event occurs immediately following deciliation that is Arp2/3 complex-mediated and dependent on clathrin-mediated endocytosis. B) Proposed model of membrane protein and membrane transport from the plasma membrane to the cilia through endocytosis.

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601 METHODS

- 602 Strains:
- The wild-type *Chlamydomonas* strain (CC-5325) and the *arpc4* mutant (LMJ.RY0402.232713)
- from the *Chlamydomonas* resource center. The *arpc4*:ARPC4-V5 strain was made by cloning
- 605 the gene into pChlamy4 (*Chlamydomonas* resource center). Colonies were screened for the 606 absence (in the case of the mutant) or presence (in the case of the rescue) by PCR using the
- 607 primers AAAAGAATTCATGGCGCTCTCACTCAGGCCATA and
- 608 AAAATCTAGACAGAAGGCAAGGGAGCGCAGGAA. The *nap1* mutant was a gift from Fred
- 609 Cross, Masayuki Onishi, and John Pringle. The SAG1-HA strain was a gift from William Snell.
- 610 Cells were grown and maintained on 1.5% Tris-Acetate Phosphate Agar (TAP) plates
- 611 (*Chlamydomonas* resource center) under constant blue (450-475 nm) and red light (625-660
- nm). For experiments, cells were grown in liquid TAP media (*Chlamydomonas* resource center)
- overnight under constant red and blue light with agitation from a rotator. To induce gametes for
- mating for the SAG1-HA experiments, cells were grown in liquid M-N media (*Chlamydomonas*
- resource center) overnight with constant red and blue light and agitation.
- 616
- 617 Ciliary studies:
- For steady state experiments, cells were treated with specified drugs [either 100μM CK-666,
- 619 250μM CK-666 (Sigma, Burlington, MA), 250μM CK-689 (Sigma, Burlington, MA), 10μM LatB
- 620 (Sigma, Burlington, MA), 10μM CHX (Sigma, Burlington, MA), or 36μM BFA (Sigma, Burlington,
- 621 MA)] and incubated with agitation for the allotted times. Following any incubation (as well as a
- 622 pre sample), cells were diluted in an equal volume of 2% glutaraldehyde and incubated at 4°
- 623 Celsius until they sediment (within 24hrs). Following sedimentation, cells were imaged using a
- Zeiss DIC scope with a 40X objective. Cilia were then measured using the segmented linefunction in ImageJ. One cilia per cell was measured and 30 cilia total were measured.
- 626
- For regeneration experiments, a pre sample was taken by adding cells to an equal volume of
- 628 2% glutaraldehyde. Then cells were deciliated with 115μ L of 0.5N acetic acid for 45 seconds.
- After this short incubation, the pH was returned to normal by adding 120μ L of 0.5N KOH. A 0-
- 630 minute sample was again taken by adding cells to an equal volume of 2% glutaraldehyde. Then
- cells were incubated with agitation and allowed to regrow cilia for the allotted time period with
- samples taken at the indicated time points by adding cells to an equal volume of 2%
 glutaraldehyde. Cells in glutaraldehyde were allowed to incubate at 4° Celsius until
- 634 sedimentation (within 24hrs). Then, cells were imaged using the same Zeiss DIC scope with a
- 40X objective. Cilia were then measured using the segmented line function in ImageJ. One cilia
- 636 per cell was measured and 30 cilia total were measured.
- 637
- 638 Click-iT OPP Protein Synthesis Assay:
- 639 Cells were grown overnight in TAP. The following day cells were deciliated as described above
- and allowed to regrow either with or without cycloheximide (10µM) to block protein synthesis. 1
- hour following deciliation, cells were mounted onto poly-lysine coverslips. Cells on coverslips
- 642 were incubated with Click-iT OPP reagent containing the O-propargyl-puromycin (OPP) which is
- 643 incorporated into nascent polypeptides for 30 minutes. OPP was removed and cells were

washed once in PBS. Cells were then fixed with 4% PFA in 1X HEPES for 15 minutes, then

645 permeabilized with 0.5% Triton-X 100 in PBS for 15 minutes. Cells were washed twice with

- 646 PBS. Detection was performed by incubating coverslips with 1X Click-iT OPP Reaction Cocktail
- 647 that includes 1X Click-iT OPP Reaction Buffer, 1X Copper Protectant, 1X Alexafluor picolyl
- azide, and 1X Click-iT Reaction Buffer Additive for 30 minutes protected from light. This was
- removed and Reaction Rinse Buffer was added for 5 minutes. This was removed and coverslips
- 650 were washed twice with PBS, allowed to dry fully, and mounted with Fluormount-G.
- 651

652 Cells were then imaged on a Nikon Spinning Disk Confocal. Z-stacks were obtained then

- combined into sum slices for quantification of maximum intensity projections for viewing. In the
 summed images, the integrated density and area of individual cells was obtained, as well as the
 background fluorescence. These were then used to calculate CTCF, which was then normalized
 to the "Pre" sample for each cell.
- 657

658 Phalloidin staining and quantification:

Procedure adapted from (Craig et al. 2019). Cells were mounted onto poly-lysine coverslips and fixed with fresh 4% paraformaldehyde in 1X HEPES. Coverslips with cells were then permeabilized with acetone and allowed to dry. Cells were rehydrated with PBS, stained with atto-phalloidin-488 (Sigma, Burlington, MA), and finally washed with PBS and allowed to dry before mounting with Fluoromount-G (Craig et al. 2019). Cells were imaged using the Nikon Spinning Disk Confocal. Z-stacks were obtained, and in ImageJ, maximum intensity projections were created for viewing.

- 666
- 667 Electron microscopy:

668 Cells (1mL of each strain) were deciliated via pH shock by adding 115μL of 0.5N acetic acid for 669 45 seconds followed by 120μL of 0.5N KOH to bring cells back to neutral pH. Cells were

allowed to regrow cilia for 30 minutes. A pre sample and a 30-minute post-deciliation sample

- 671 were fixed in an equal volume of 2% glutaraldehyde for 20 minutes at room temperature.
- 672 Samples were then pelleted using gentle centrifugation for 10 minutes. The supernatant was
- 673 removed, and cells were resuspended in 1% glutaraldehyde, 20mM sodium cacodylate. Cells
- 674 were incubated for 1 hour at room temperature and then overnight at 4° Celsius. This protocol
- 675 was first reported in (W. L. Dentler and Adams 1992).
- 676
- 677 SAG1-HA Immunofluorescence:
- 678 Procedure adapted from (Belzile et al. 2013). SAG1-HA cells were grown overnight in M-N 679 media to induce gametes. These cells were then treated with either 10μ M LatB for 1 hour or
- 680 250μM CK-666 for 2 hours. Following treatment, mating was induced by adding db-cAMP
- 681 (ChemCruz, Santa Cruz, CA) to a final concentration of 13.5mM and incubating for 30 minutes.
- 682 Cells were adhered to coverslips and fixed with methanol. Cells were then dried and rehydrated
- with PBS and incubated with 100% block (5% BSA, 1% fish gelatin) for 30 minutes. The 100%
 block was replaced with new 100% block containing 10% normal goat serum for another 30-
- 684 block was replaced with new 100% block containing 10% normal goat serum for another 30-685 minute incubation. The primary antibody (rat anti-HA, Sigma, Burlington, MA) was diluted
- 1:1000 in 20% block in PBS. Coverslips were incubated at 4° Celsius in a humidified chamber
- 687 overnight. The primary antibody was removed and washed away with 3 10-minute PBS washes.
- The secondary (anti-rat IgG-Alexafluor 488, Invitrogen, Carlsbad, CA) was added and coverslips
- 689 were incubated at room temperature for 1 hour. This was followed by 3 more 10-minute PBS
- 690 washes and finally mounting with Fluoromount-G. Cells were imaged using a Nikon widefield
- 691 microscope. Z-stacks were obtained, and maximum intensity projections were created for 692 visualization and sum slices were created for quantification using ImageJ.
- 693

Images were quantified by using line scans from the apex of the cells to the basal region of the cells farthest away from the apex. Line scans were then normalized, and background subtracted before being combined into single graphs. Using the line scans, the intensity of signal at the basal region of the cells was subtracted from the signal at the apical region. Finally, cells with a difference over 30 were considered to be apically enriched and this was quantified as percentage of cells with apical staining.

700

701 SAG1-HA western blot:

702 Procedure adapted from (Belzile et al. 2013). SAG1-HA cells were grown overnight in M-N 703 media to induce gametes. These cells were then treated with either 10µM LatB for 1 hour or 704 250µM CK-666 for 2 hours. Following treatment, mating induction was done by adding db-cAMP 705 to a final concentration of 13.5mM and incubating for 10 minutes. Cells were then treated with 706 0.01% trypsin for 5 minutes, pelleted (at 500xg for 2 minutes), resuspended in lysis buffer (5% alvcerol, 1% NP-40, 1mM DTT, 1X protease inhibitors), and then lysed with bead beating. Cell 707 708 debris was spun down at 14000xg for 15 minutes. An equal amount of protein was loaded to a 709 10% SDS-PAGE gel. The resulting gel was transferred to membrane which was then blocked 710 with 5% milk in PBST. The primary antibody (rabbit anti-HA, Cell Signaling, Danvers, MA) 711 diluted to 1:1000 in 1% BSA, 1% milk was added and incubated overnight at 4° Celsius. Primary 712 antibody washed off with 3 10-minute PBST washes. Secondary antibody (anti rabbit IgG, 713 Invitrogen, Carlsbad, CA) was diluted to 1:5000 in 1% milk. 1% BSA was added, and the blot 714 was incubated for 1 hour. Membrane was probed with West Pico Chemiluminescent Pico 715 Substrate (Invitrogen, Carlsbad, CA). The same membrane was stripped of antibody and total 716 protein was determined with amido black staining. Band intensity was measured in ImageJ and 717 normalized to total protein

- 718
- 719 Membrane stain:
- 720 FM 4-64FX membrane stain (Thermo, Waltham, MA) was diluted to a stock concentration of
- 200µg/mL. Cells were adhered to poly-lysine coverslips. After a 5-minute incubation, cells were
- tilted off and 5µg/mL of ice-cold stain in Hank's Buffered Salt Solution (HBSS) without
- magnesium or calcium was added for 1 minute. The stain was tilted off and cells were fixed with
- ice cold 4% paraformaldehyde in HBSS without magnesium or calcium for 15 minutes.
- 725 Coverslips were then rinsed 3 times for 10 minutes each in ice cold HBSS without magnesium
- or calcium. Finally, cells were mounted with Fluoromount-G and imaged using the Nikon
- 527 Spinning Disk Confocal. Z-stacks were taken and combined into sum projections using ImageJ.
- The background corrected total cell fluorescence was then calculated by taking the integrated
- density and subtracting the sum of the area and the mean background intensity.
- 730
- 731 Clathrin light chain immunofluorescence:
- 732 Cells were grown overnight in TAP media. Cells were deciliated using low pH shock. Cells are
- then adhered to coverslips and fixed with 4% PFA in 1X HEPES. Cells were then dried and
- rehydrated with PBS and incubated with 100% block (5% BSA, 1% fish gelatin) for 1 hour. The
- primary antibody (goat anti-clathrin light chain, Abcam, Cambridge, UK or rabbit anti-acetylated
- tubulin, Cell Signaling, Danvers, MA) was diluted 1:1000 in 20% block in PBS. Coverslips were
- incubated at 4° Celsius in a humidified chamber overnight. The primary antibody was removed
- and washed away with 3 10-minute PBS washes. The secondary (donkey anti-goat IgG-
- Alexafluor 488, Invitrogen, Carlsbad, CA or goat anti-rabbit IgG-Alexafluor 568, Invitrogen,
- 740 Carlsbad, CA) was added and coverslips were incubated at room temperature for 1 hour. For
- cells stained with DAPI, DAPI (Biotium, Fremont, CA) was added for the last 10 minutes of
- secondary antibody incubation. This was followed by 3 more 10-minute PBS washes and finally
- mounting with Fluoromount-G. Cells were imaged using a Nikon widefield microscope. Z-stacks

were obtained, and maximum intensity projections were created for visualization and sum sliceswere created for quantification using ImageJ.

- 746
- 747 Biotin ciliary isolation:

Procedure adapted from (W. Dentler 2013). 100mL of cells were grown in TAP for each 748 condition until they reached an OD₇₃₀ of 1.6 or above. Cells were then spun down and 749 750 resuspended in M1 media and allowed to grow overnight. The next day cells were spun down at 751 1800rpm for 3 minutes and resuspended in HM Media (10mM HEPES, 5mM MgSO4, pH 7.2). 752 Solid biotin (Thermo, Waltham, MA) was added to 20µg/mL for each strain and incubated for 5 minutes with agitation. Cells were diluted with 10 volumes of fresh M1 media before being spun 753 754 down at 1800rpm for 3 minutes. After all cells were pelleted, they were washed with fresh M1 media three times. A pre sample was set aside (100mL) and the remainder of the cells were 755 756 resuspended in 4.5 pH M1 media for 45 seconds before being spun down again at 1800rpm for 757 3 minutes. Cells were then resuspended in pH 7.0 media and allowed to regrow their cilia for 4 758 hours. A sample was taken pre-biotinylation to use as a control for non-specific streptavidin 759 binding.

760

761 Meanwhile, the cilia were isolated from the pre sample. The samples were centrifuged for 3 762 minutes at 1800rpm. Supernatant was drained and each pellet was resuspended in 2 mL of 763 10mM HEPES (pH 7.4). This was repeated 2 times. Then each pellet was resuspended in 1 mL 764 of fresh ice-cold 4% HMDS (10mM HEPES pH 7.4, 5mM MgSO4, 1mM DTT, 4% w/v sucrose). 765 Cells were deciliated by incubating with 25mM dibucaine for 2 minutes. Then ice cold HMDS 766 with 0.5mM EGTA was added (1mL per 1.5mL of cells). This was then centrifuged for 3 minutes 767 at 1800rpm. Supernatant was collected for each sample. Then HMDS with 25% sucrose was 768 layered beneath the supernatant (2 mL of 25% HMDS for 1mL of supernatant) to create an 769 interface. This was centrifuged at 4° Celsius for 10 min at 2400rpm with no brake to avoid 770 disrupting interface where cilia should now be located. Cilia were removed, pelleted at 21130xg 771 for 30 minutes, then resuspended in lysis buffer (5% glycerol, 1% NP-40, 1mM DTT, 1X 772 protease inhibitors). This was repeated with the post samples 4 hours following deciliation. An 773 equal amount of protein was loaded to a 10% SDS-PAGE gel. The resulting gel was transferred 774 to PVDF membrane. The membrane was washed 2x with PBSAT (PBST + 0.1% BSA), then 775 incubated with HRP-conjugated streptavidin (Thermo, Waltham, MA) for 1 hour. The membrane 776 was then washed 3 times with PBSAT (10 minutes each) and 3 times with PBST (15 minutes 777 each). Membrane was probed with West Pico Chemiluminescent Pico Substrate (Invitrogen. 778 Carlsbad, CA). The same membrane was stripped of antibody and incubated with Coomassie 779 Brilliant Blue to observe total protein.

- 780
- 781 Homology modeling and sequence studies:

Arp2/3 homology model was created using the Modeller plugin in UCSF Chimera. The template used was 1U2Z (Nolen, Littlefield, and Pollard 2004; Sali and Blundell 1993; Pettersen et al.

- 2004). Percent identity and similarity is calculated in relation to the human Arp2/3 complex
- 785 members using a MUSCLE alignment in Geneious. The homology model was visualized and
- conservation was mapped on the protein surface using Chimera (Pettersen et al. 2004).
- 787
- 788 Statistical analysis:

789 Statistical analyses were done if GraphPad Prism Version 9. Superplots were created using the

- method in (Lord et al. 2020). For any experiments comparing 2 groups (**Figure 3D, 5C, and 5E**)
- an unpaired student's t-test was used to determine P value. For experiments comparing multiple
- samples at a single time point (Figure 1A, 1B, and 2B), an ANOVA was used. Finally, for any
- graphs covering several time points (Figure 1C, 1D, 2C, 3B, and 6E), multiple comparisons

were performed (Tukey's and Sidak's). For any percentages shown (Figure 7D), Chi-squared
analysis was performed. For all experiments **** P<0.0001, *** P<0.001, ** P<0.01, * P<0.1.

797

798 **ACKNOWLEDGEMENTS**:

Our most sincere gratitude to William Dentler for providing his expertise especially in looking at the electron microscopy images and for his helpful advice, William Snell for generously providing the SAG1-HA strain, Masayuki Onishi for generously providing the *nap1* mutant strain, Henry Higgs for his feedback on version 1 of the manuscript, Ann Lavanway for assistance with microscopy, and the Avasthi lab for all their help throughout the project. We would also like to thank David Sept and Courtney M Schroeder for the help with the original version of this paper and for providing helpful comments throughout the process.

We also thank our funding sources including the Madison and Lila Self Graduate
Fellowship at the University of Kansas Medical Center and the MIRA (R35GM128702). Finally,
we thank the BioMT core at Dartmouth College (NIH/NIGMS COBRE award P20-GM113132),
the Genomics and Molecular Biology Shared Resources Core (NCI Cancer Center Support
Grant 5P30CA023108-37), and the KIDDRC NIH U54 HD 090216 at the University of Kansas
Medical Center, Kansas City, KS 66160.

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