- 1 Title: Evolution-guided mutagenesis of the cytoplasmic incompatibility proteins: Identifying
- 2 CifA's complex functional repertoire and new essential regions in CifB
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- 56 Short Title: Mutagenesis of CI proteins
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- Keywords: cytoplasmic incompatibility, *Wolbachia*, prophage WO, *Drosophila melanogaster*,
   Two-by-One model
- 25

#### 26 Abstract

27 Wolbachia are the world's most common, maternally-inherited, arthropod endosymbionts. Their 28 worldwide distribution is due in part to a selfish drive system termed cytoplasmic incompatibility (CI) that confers a relative fitness advantage to females that transmit Wolbachia to their offspring. 29 CI results in embryonic death when infected males mate with uninfected females but not infected 30 females. Under the Two-by-One genetic model of CI, males expressing the two phage WO 31 32 proteins CifA and CifB cause CI, and females expressing CifA rescue CI. While each protein is predicted to harbor three functional domains, there is no knowledge on how sites across these 33 34 Cif domains, rather than in any one particular domain, contribute to CI and rescue. Here, we use evolution-guided, substitution mutagenesis of conserved amino acids across the Cif proteins, 35 coupled with transgenic expression in uninfected Drosophila melanogaster, to determine the 36 37 functional impacts of conserved residues evolving mostly under purifying selection. We report that amino acids in CifA's N-terminal unannotated region and annotated catalase-related domain are 38 39 important for both complete CI and rescue, whereas C-terminal residues in CifA's putative domain of unknown function are solely important for CI. Moreover, conserved CifB amino acids in the 40 predicted nucleases, peptidase, and unannotated regions are essential for CI. Taken together, 41 42 these findings indicate that (i) all CifA amino acids determined to be involved in rescue are 43 correspondingly involved in CI, (ii) an additional set of CifA amino acids are uniquely important 44 in CI, and (iii) CifB amino acids across the protein, rather than in one particular domain, are all 45 essential for CI. We discuss how these findings advance an expanded view of Cif protein evolution and function, inform the mechanistic and biochemical bases of Cif-induced Cl/rescue, and 46 47 continue to substantiate the Two-by-One genetic model of CI.

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#### 50 Article summary

51 Wolbachia are maternally-transmitted, intracellular bacteria that occur in approximately half of arthropod 52 species worldwide. They can spread rapidly though host populations via the cytoplasmic incompatibility 53 (CI) drive system. CI causes embryonic death when infected males mate with infected females, but 54 offspring of infected females are rescued. Two proteins, CifA and CifB, underlie the genetic basis of CI and 55 rescue, but how amino acid sites across these proteins contribute to CI and/or rescue remain unknown. 56 Here, we employed evolution-guided, combinatorial mutagenesis on conserved amino acids to 57 understand their relative contributions to CI and rescue. The results of this study reveal a phenotypic 58 complexity underlying the expression of these proteins and provide relevance to the biochemical and 59 mechanistic bases of CI and rescue.

60

#### 62 Introduction

63 Wolbachia are maternally-inherited, intracellular  $\alpha$ -Proteobacteria that occur in 40-65% of 64 all arthropod species (Charlesworth et al., 2019; Hilgenboecker et al., 2008; Weinert et al., 2015; Zug and Hammerstein, 2012). Residing in the cells of reproductive tissues, Wolbachia commonly 65 cause a selfish drive system, cytoplasmic incompatibility (CI), that yields a relative advantage to 66 females that transmit Wolbachia and thus increases Wolbachia's rate of spread through the 67 68 matriline (Rasgon, 2008; Turelli, 1994). CI causes embryonic death when Wolbachia-infected 69 males mate with uninfected females and is rescued when the female is infected with a compatible 70 Wolbachia strain (Fig. 1A) (LePage and Bordenstein, 2013; Serbus et al., 2008; Taylor et al., 71 2018). CI can act as a form of reproductive isolation between populations of different infection states (Bordenstein et al., 2001; Brucker and Bordenstein, 2012; Jaenike et al., 2006; Shropshire 72 73 and Bordenstein, 2016). Additionally, this drive system has brought Wolbachia to the forefront of vector control efforts to combat Zika and dengue viruses because wMel Wolbachia from 74 75 Drosophila melanogaster flies confer resistance to RNA arboviruses when transinfected into Aedes mosquitoes (Aliota et al., 2016; Caragata et al., 2016; Kittayapong et al., 2018; Moreira et 76 al., 2009; O'Connor et al., 2012; O'Neill, 2018; X. Zheng et al., 2019). Notably, wMel-induced CI 77 78 is the focus of this study.

79 CI's microbial genetic basis can be described by the Two-by-One genetic model (Fig. 1B) 80 (Shropshire and Bordenstein, 2019) where the two phage WO genes *cifA* and *cifB* cause CI when 81 dually expressed in testes (Beckmann et al., 2017; LePage et al., 2017; Shropshire and Bordenstein, 2019), and *cifA* rescues CI when singly expressed in ovaries (Chen et al., 2019; 82 83 Shropshire et al., 2018; Shropshire and Bordenstein, 2019). The Cif proteins are divided into at least five phylogenetic clades, referred to as Types 1 - 5 (Bing et al., 2020; LePage et al., 2017; 84 85 Lindsey et al., 2018). To date, this genetic model has been well supported using Type 1 *cif* genes from *w*Mel (LePage et al., 2017; Shropshire et al., 2018; Shropshire and Bordenstein, 2019) and 86 is consistent with results with Type 1 and 4 genes from wPip of Culex pipiens mosquitos 87 (Beckmann et al., 2019b; Chen et al., 2019). However, the mechanism underlying CifA;B-induced 88 89 CI and CifA-induced rescue remains mostly unresolved and limited to in vitro assays and structural homology-based predictive annotations discussed below. 90

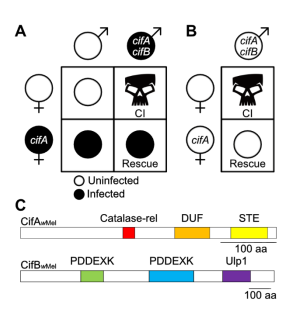
CifA<sub>wMel</sub> is weakly predicted to encode a catalase-related domain (catalase-rel), a domain
of unknown function 3243 (DUF), and a sterile-like transcription factor domain (STE) (Fig. 1C)
(Lindsey et al., 2018). Catalase-rel domains are predicted to catalyze the degradation of reactive
oxygen species (ROS) (Guy et al., 2005; Loew, 1900). DUF has a distant homology to globin-like
domains and Puf-family RNA-binding domains, which influence the stability of eukaryotic RNAs

96 (Kumar and Subramaniam, 2018; Nishanth and Simon, 2020). Finally, STE domains mediate 97 transcriptional induction in yeast (Wong Sak Hoi and Dumas, 2010). Structural homology 98 predictions identify the STE and Puf-family RNA-binding domains across four and five 99 phylogenetic Cif Types respectively, whereas the catalase-rel domain is only annotated in Type 100 1 (Bing et al., 2020; Lindsey et al., 2018).

Conversely, CifB<sub>wMel</sub> harbors two putative PD-(D/E)XK-like nuclease domains (PDDEXK), 101 102 and a ubiquitin-like-specific protease 1 domain (Ulp1) (Fig. 1C) (Beckmann et al., 2017; LePage 103 et al., 2017; Lindsey et al., 2018). The Ulp1 domain is restricted to Type 1 CifB and cleaves poly-104 ubiquitin chains in vitro (Beckmann et al., 2017). Since mutation of the CifB Ulp1's catalytic motif 105 ablates CifB's ability to contribute to CI when transgenically expressed in *D. melanogaster*, the Ulp1 domain has previously been described as the "enzymatic warhead" for CI induction 106 107 (Beckmann et al., 2017). The function of the remaining sequence, including the PDDEXK dimer of Type 1 CifB, has not been assessed. We previously demonstrated that this remaining sequence 108 109 comprises a CifB central core region found throughout Cif phylogenetic types (LePage et al., 2017). Moreover, the PDDEXK domains are also annotated in all phylogenetic Cif clades (Bing et 110 al., 2020; LePage et al., 2017; Lindsey et al., 2018), and in vitro assays indicate that the PDDEXK 111 112 domains of Type 4 CifB<sub>wPip</sub> nick DNA (Chen et al., 2019). As such, CifB's central core including 113 the PDDEXK domains is likely important, if not one of the central aspects of its ability to contribute 114 to CI. Finally, in vitro assays suggest that CifA and CifB can bind (Beckmann et al., 2017), but the 115 importance of this binding in vivo and for phenotypic output remains unknown.

The aforementioned protein annotations and biochemical data represent a foundation to 116 117 develop a more complete and nuanced understanding of how Cif proteins cause CI and rescue, 118 but it remains unclear what kind of impact each domain and unannotated region have on the 119 phenotypic output of these proteins. Here, we test the importance of conserved amino acids 120 across the Cif proteins to CI and rescue via site-directed substitution mutagenesis and transgenic expression in D. melanogaster. We report three key findings. First, conserved sites in CifA's N-121 122 terminal unannotated region and the catalase-rel domain are important in both CI and rescue. 123 Second, conserved sites in CifA's DUF are only involved in CI. Finally, all tested conserved sites in CifB are required for CI. Taken together, we identify sites in seven Cif mutants (both CifA and 124 125 CifB) essential for complete CI, and determine that CifA's N-terminus is involved in both CI and 126 rescue while the middle of the protein is only involved in CI. These results inform the mechanistic and biochemical basis of CI and rescue and lend further support for a Two-by-One genetic model 127 128 where both CifA and CifB are functionally crucial for expression of CI.





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Figure 1. Cytoplasmic incompatibility, the Two-by-One genetic model, and Cif protein architecture. (A) CI is caused when *Wolbachia*-infected males expressing *cifA* and *cifB* mate with uninfected females. If females are infected and express cifA then offspring are rescued. (B) The Two-by-One genetic model indicates that males expressing cifA and cifB, even in the absence of an infection, can cause CI that can be rescued by uninfected females expressing cifA. (C) Schematic showing the protein architecture of CifA and CifB from the *w*Mel *Wolbachia* of *D. melanogaster*. Annotations are based on a prior structural homology-based analyses (Lindsey et al., 2018).

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#### 139 Results

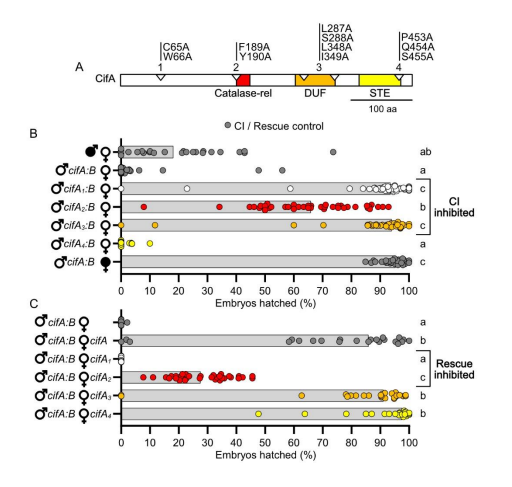
### 140 CifA mutants impact CI and rescue

141 Since CifA does not have any putative catalytic motifs, we used a previous sequence analysis of conserved amino acid residues in an alignment of phylogenetically diverse CifA 142 proteins (Lindsey et al., 2018) to select highly conserved sites across the protein for mutagenesis. 143 CifA1, CifA2, CifA3, and CifA4 have combinatorial, alanine substitutions in the N-terminal 144 145 unannotated region and putative catalase-rel, DUF, and STE domains, respectively (Fig. 2A). Alanine mutagenesis is used to analyze the importance of specific amino acids in protein 146 sequences without contributing significant structural variation to the protein (Cunningham and 147 Wells, 1989). We tested mutant *cifA* transgenes for their ability to (i) induce CI when dually 148 expressed with wild type *cifB* in tests of uninfected males and (ii) rescue when singly expressed 149 150 in ovaries of uninfected females. Using the GAL4-UAS system in D. melanogaster (Duffy, 2002), we predict that CI and/or rescue will not be recapitulated when CifA mutants are transgenically 151 152 expressed if the sites mutated are crucial to the respective function. Notably, we use the nos-GAL4:VP16 driver throughout this study since it was previously shown to enable strong transgenic 153 154 CI when dually expressing *cifA* and *cifB* transgenes (Shropshire and Bordenstein, 2019). Since

155 CI manifests as embryonic death, we measured the strength of CI induced under mutant 156 transgenic expression by measuring the percentage of *D. melanogaster* embryos that hatch into 157 larvae.

Consistent with prior studies (Shropshire and Bordenstein, 2019), dual cifA; B expression 158 159 in males yielded nearly complete CI-defining embryonic death when mated to uninfected females 160 (Mdn = 0% hatching) that was statistically comparable to *w*Mel-induced CI (p > 0.99), and it was 161 rescued by *w*Mel-infected females (Mdn = 95.4% hatching) (Fig. 2B). Transgenic dual expression 162 of either  $cifA_1$ ; B (Mdn = 93.7% hatching; p > 0.99) or  $cifA_3$ ; B (Mdn = 94.1% hatching; p > 0.99) in 163 males crossed to uninfected females revealed no statistically significant difference in hatching 164 relative to the compatible, rescue cross; thus, mutating conserved sites in CifA's unannotated region and putative DUF ablates CI. Conversely, transgenic expression of *cifA*<sub>4</sub>;*cifB* caused hatch 165 166 rates statistically comparable to *cifA*; *B*-induced CI (Mdn = 0% hatching; p > 0.99), suggesting that mutation of conserved sites in the putative STE did not impact *cifA*'s ability to contribute to CI. 167 168 Finally, transgenic expression of  $cifA_2$ ; B (Mdn=66.0%) yielded an intermediate phenotype whereby it was statistically different from both cifA;B-induced CI (p = 0.0006) and rescue of 169 transgenic CI (p = 0.0001), indicating that the putative catalase-rel mutant induces a partial CI 170 171 phenotype. Together, these results suggest the mutated sites in the unannotated region, catalase-172 rel, and DUF of CifA are important for CI-induction (Fig. 2B).

173 Next, we reciprocally tested if uninfected transgenic females singly expressing the same 174 cifA mutants rescue cifA;B-induced CI (Fig. 2C). As above, dual cifA;B expressing males induced near-complete embryonic death consistent with strong CI (Mdn = 0% hatching), and this lethality 175 176 could be rescued when the female expressed cifA (Mdn = 85.9% hatching). Transgenic 177 expression of  $cifA_1$  (Mdn = 0.00%; p < 0.0001) and  $cifA_2$  (Mdn=27.6%; p = 0.0390), which failed 178 to contribute to CI, also failed to rescue *cifA;B*-induced CI as compared to the standard transgenic rescue cross. Conversely, transgenic expression of  $cifA_3$  (Mdn=91.2%; P > 0.9999) and  $cifA_4$ 179 (Mdn=97.6%; P = 0.3039) rescued cifA; B-induced CI at levels comparable to the standard 180 181 transgenic rescue cross. These results suggest that the sites mutated in the unannotated and 182 catalase-rel regions of CifA are important for rescue (Fig. 2C).



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**Figure 2.** *cifA*<sub>1</sub> *and cifA*<sub>2</sub> *fail* to cause or rescue CI, *and cifA*<sub>3</sub>*can* rescue but fails to cause CI. (A) schematic showing the location of amino acid mutations in CifA relative to previously predicted domains (Lindsey et al., 2018). (B) Hatch rate experiment testing if *cifA* mutants can induce CI when dual expressed with *cifB* in uninfected males. (C) Hatch rate experiment testing if expressing *cifA* mutants can rescue transgenic CI when expressed in uninfected females. (B/C) Each dot represents the percent of embryos that hatched from a single male and female pair. Expressed genes are noted to the right of the corresponding sex. Gray bars represent median hatch rates for each cross and letters to the right indicate significant differences based on  $\alpha = 0.05$  calculated by Kruskal-Wallis and Dunn's test for multiple comparisons between all groups. Panel B was conducted three times and C was conducted twice. P-values are reported in Table S1.

#### 194

### 195 CifB mutants ablate Cl

Four CifB mutants were constructed based on a comparative sequence analysis of 196 197 conserved residues (Lindsey et al., 2018). All CifB mutations are similarly alanine substitutions, with the exception of one glycine mutation of a conserved alanine (Fig. 3A). Glycine was chosen 198 199 to replace alanine since it is comparably sized and would be less likely to impact protein structure than other amino acids. CifB1, CifB2, CifB3, and CifB4 have mutations in the N-terminal 200 unannotated region, first PDDEXK, second PDDEXK, and Ulp1 respectively (Fig. 3A). The Ulp1 201 mutation is the same used previously to test for the catalytic activity of the Ulp1 domain 202 203 (Beckmann et al., 2017). We predict that CI will not be recapitulated when CifB mutants are 204 transgenically expressed if the sites mutated are crucially important for CI-induction. As with CifA

205 mutants above, we tested mutant CifB for their ability to induce CI when dually expressed with 206 *cifA* in uninfected males.

207 As expected, dual *cifA*: B expression in uninfected males caused hatch rates statistically comparable to *w*Mel-induced CI (p > 0.99), and it could be rescued by *w*Mel-infected females 208 (Mdn = 93.9% hatching). However, transgenic expression of  $cifA;B_1$  (Mdn = 96.3%; p < 0.0001), 209  $cifA:B_2$  (Mdn = 95.6%; p < 0.0001),  $cifA:B_3$  (Mdn = 94.3%; p < 0.0001), and  $cifA:B_4$  (Mdn = 93.0%; 210 p < 0.0001) all failed to reduce hatch rates statistically comparable to *cifA*;*B*-induced CI (Mdn = 211 212 0.%). These results specify that all mutated conserved sites, rather than any one site or domain such as the previously reported catalytic site of Ulp1 (Beckmann et al., 2017), are important for 213 214 CifB in CI-induction.

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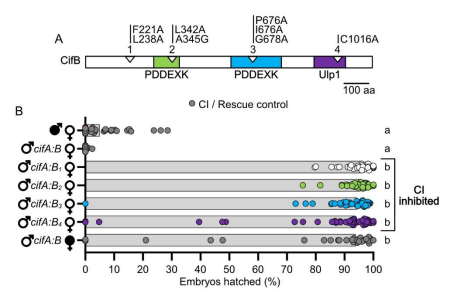


Figure 3. All *cifB* mutants fail to contribute to Cl. (A) schematic showing the location of mutations in CifB relative to previously predicted domains (LePage et al., 2017; Lindsey et al., 2018). (B) Hatch rate experiment testing if *cifB* mutants can induce Cl when dual expressed with *cifB* in uninfected males. Each dot represents the percent of embryos that hatched from a single male and female pair. Expressed genes are noted to the right of the corresponding sex. Gray bars represent median hatch rates for each cross and letters to the right indicate significant differences based on  $\alpha = 0.05$  calculated by Kruskal-Wallis and Dunn's test for multiple comparisons between all groups. Panel B was conducted three times. P-values are reported in Table S1.

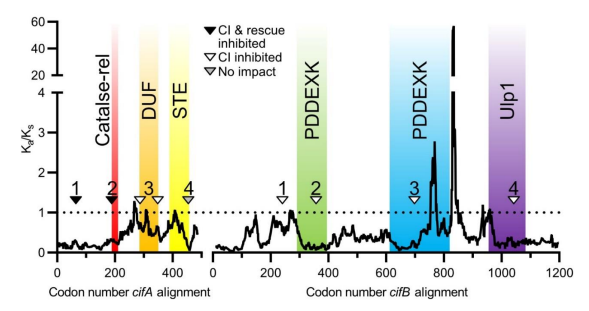
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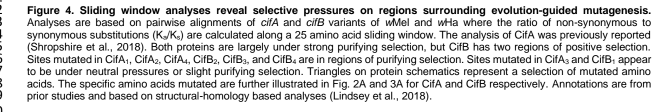
225 <u>CifB selection analysis.</u>

Prior selection analyses using a sliding window analysis indicated that CifA is under strong purifying selection, and CifA's N-terminal unannotated region and catalase-rel domain are under stronger purifying selection than C-terminal regions (Shropshire et al., 2018). Here, in order to assess how engineered mutations and phenotypes align with selective pressures, we replotted this analysis for CifA and, for the first time, applied a sliding window analysis of K<sub>a</sub> and K<sub>s</sub> (SWAKK) to assess the selective pressure on CifB (Fig. 4). SWAKK calculates the ratio of nonsynonymous to synonymous substitution rates ( $K_a/K_s$ ) in a pairwise alignment. Codons with ratios below one are considered under purifying selection, and those above one are under positive selection. In both cases, pairwise nucleotide alignments were *cif<sub>wMel</sub>* relative to *cif<sub>wHa</sub>*, which are modestly divergent Type 1 *cif* variants. For both CifA and CifB, we then plotted the amino acid substitutions to assess if they occur in regions of purifying or positive selection.

For CifA, all mutant sites were in areas of purifying selection (Fig. 4). However, the 237 strongest purifying selection was surrounding the codons we mutated in the N-terminal 238 239 unannotated region and catalase-rel domain. Intriguingly, the sites mutated near CifA's STE are 240 under strong purifying selection, but the rest of the domain evolves neutrally. These results are in-line with our observations that the mutations in the N-terminal unannotated region and catalase-241 242 rel domain are involved in both CI and rescue, whereas weaker selection may act on the DUF domain which based on the results here are only involved in CI. Additionally, as with CifA, the 243 majority of CifB is under strong purifying selection (Fig. 4). However, there are two clusters of 244 245 amino acids that appear to be under strong positive selection. These residues are on the C-246 terminal side of CifB's second PDDEXK domain and in the unannotated region between the 247 PDDEXK domain and Ulp1 domain. CifB<sub>1</sub>'s mutations fall within regions under neutral pressures, 248 whereas the other CifB mutants are in regions under strong purifying selection. 249



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### 261 <u>Cif structural predictions</u>

There are numerous ways to interpret the impact of a mutation on a protein's function. These can include non-exclusive changes to catalytic motifs, ligand binding sites, or changes in local or global structures that ablate, enhance, or otherwise modify the phenotypic output of the protein. The CifB<sub>4</sub> mutation in the Ulp1 domain involves a putative cysteine catalytic motif common to deubiquitinase domains (Beckmann et al., 2017). Otherwise, no other catalytic motifs or binding sites have been identified in CifA<sub>wMel</sub> or CifB<sub>wMel</sub>. As such, we aimed to investigate the impact of these mutations on the structure of CifA and CifB proteins.

269 The Iterative Threading ASSEmbly Refinement (I-TASSER) webserver was used to generate a list of structural homologs from the protein databank (PDB) for each wild-type and Cif 270 mutant protein and construct structural models based on these hits (Zhang, 2009). The shared 271 272 and unique PDB hits for wild-type and mutant proteins are summarized in Fig. S1A and detailed in Table S2. The top 10 PDB hits for each protein were used to create structural models (Fig. 273 274 S1B). Each model is generated with confidence measures in the form of C-scores and TM-scores. C-scores range from -5 to 2 where 2 is the highest confidence, and TM-scores range from 0-1 275 where 1 is the highest confidence (Zhang, 2009). The similarity between wild-type and mutant 276 277 structures was then assessed using the Alignment plugin in PyMOL 2.3.2 which provides values 278 for the root-mean-square deviation (RMSD) of atomic positions. Higher RMSDs indicates a 279 greater distance between the atoms of mutant proteins superimposed on the wild-type protein. 280 Structural models were generated for CifA (C-score = -2.74;TM =  $0.4\pm0.13$ ), CifA<sub>1</sub> (C-score = -2.74;TM =  $0.4\pm0.13$ ), CifA<sub>1</sub> (C-score = -2.74;TM = -2.1.42; TM = 0.54±0.15; RMSD = 19.7), CifA<sub>2</sub> (C-score = -2.84; TM = 0.39±0.13; RMSD = 1.9), CifA<sub>3</sub> 281 282  $(C-score = -1.39;TM = 0.54\pm0.15; RMSD = 20.2), and CifA<sub>4</sub> (C-score = -1.43;TM = 0.54\pm0.15;$ 283 RMSD = 19.8) (Fig. S1B). These low C-scores and TM-scores indicate that the I-TASSER 284 predictions for CifA are not robust, and that variation in structure between wild-type and mutant proteins could be the result of poor threading templates. However, these results suggest that CifA 285 is structurally most comparable to CifA<sub>2</sub> while the other structures are predicted to change to 286 287 comparable degrees. Crucially, since the most divergent model, CifA<sub>3</sub> remains rescue-capable and the second most divergent model CifA4 remains functional in both CI and rescue it is unlikely 288 289 that predictions of global structural variation assists an understanding of phenotypic ablation in 290 CifA.

I-TASSER was also used to identify PDB hits and create structures for wild-type and
 mutant CifB. The shared and unique PDB hits for wild-type and mutant proteins are summarized
 in Fig. S1C and detailed in Table S2. As above, I-TASSER protein structures were then created
 based on these threading templates and compared for RMSD. Structural models (Fig. S1D) were

generated for CifB (C-score = -1.02, TM-score =  $0.59\pm14$ ), CifB<sub>1</sub> (C-score = -0.68, TM-score =  $0.63\pm14$ ; RMSD = 1.7), CifB<sub>2</sub> (C-score = -1.03, TM-score =  $0.58\pm14$ ; RMSD = 10.6), CifB<sub>3</sub> (C-score = -1.07, TM-score =  $0.58\pm14$ ; RMSD = 2.2), CifB<sub>4</sub> (C-score = -0.65, TM-score =  $0.63\pm14$ ; RMSD = 2.2) (Fig. S1D). Together, these results suggest that all CifB mutants are structurally comparable to the wild-type protein. As with CifA, it remains unknown how small effects in protein structure may influence phenotypic ablation.

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### 302 Discussion

303 Cl's genetic basis involves a Two-by-One genetic model whereby *cifA;B* expressing males 304 cause CI and *cifA* expressing females rescue CI (Beckmann et al., 2017, 2019b; LePage et al., 305 2017; Shropshire et al., 2018; Shropshire and Bordenstein, 2019). However, the mechanistic 306 basis of *cifA*; *B*-induced CI and *cifA*-induced rescue remains largely unresolved. Here, we test the phenotypic impact of conserved sites (Lindsey et al., 2018) to CI and rescue across the CifwMel 307 308 proteins in vivo using site-directed mutagenesis and transgenic expression in D. melanogaster. 309 We discuss the relevance of these findings, particularly the complex functional repertoire of CifA, to the Toxin-Antidote (TA) and Host Modification (HM) models of CI (Beckmann et al., 2019a; 310 Shropshire et al., 2019) and the molecular basis of CifA;B-induced CI and CifA-induced rescue. 311

312 Crucially, phenotypic ablation caused by site-directed substitution mutagenesis has 313 multiple possible and non-exclusive interpretations. First, a mutated site may be part of a catalytic 314 motif. If so, mutagenesis may result in a significant inhibition of enzymatic activity related to CI and/or rescue induction. Second, the mutated site may be crucial for binding to host ligands, 315 316 binding between CifA and CifB, or binding to nucleotide products. Additionally, amino acid 317 substitutions may impact protein structure which in turn could also ablate function. Global folding 318 abnormalities may block the enzymatic function of trans-acting domains or prevent binding elsewhere in the protein. Local or non-local changes to the structure can inhibit processes close 319 to the mutated site in tertiary structure (Mi et al., 2019; Yang et al., 2020). These hypotheses are 320 321 applicable to all mutant variants. Below, we place our mutagenesis work in the context of prior in 322 vivo assays and bioinformatics to develop hypotheses regarding the mechanistic basis of CI and 323 rescue.

First and foremost, our results indicate that CifA has several amino acids with overlapping functions in CI and rescue, as well as amino acids of specific importance to CI. Thus, it is plausible that, CifA's functional role in CI is comparable to that in rescue. Among the two sets of mechanistic models for CI, HM-based models predict that expression of CifA and CifB cause CI through host modifications during spermatogenesis, and rescue occurs by reversing those modifications in the

329 embryo (Shropshire et al., 2019). In the context of these results, CifA's role in modifying host 330 factors in the testes may be comparable to its activity in rescue. Indeed, the HM-based mistiming 331 model posits that CI causes a delay in nuclear apposition during the first mitosis due to slowed development of the male pronucleus, and rescue occurs when the development of the female 332 pronucleus is comparably delayed (Ferree and Sullivan, 2006; Tram and Sullivan, 2002). Thus, 333 under the mistiming model. CifA's overlapping function in CI and rescue could indicate that it is 334 335 the primary driver of CI-induction and rescue, whereas CifB and CI-specific sites in CifA may 336 provide adjunct functions necessary to access those targets when expressed in the testes. 337 Conversely, the TA model predicts that CifB is paternally-transferred and the primary factor that 338 causes an embryonic toxicity, and CifA is necessary to prevent self-induced toxicity in the testes 339 and to rescue CifB-toxicity in the embryo (Beckmann et al., 2019a). Unlike the HM model, the TA 340 model predicts that CifA must function exactly as an antidote when expressed in testes and 341 embryos. Thus, our data are also in-line with these expectations, but as with the HM model, the 342 TA model must be expanded to explain why sites in CifA's DUF domain are only essential for CI-343 induction if CifA's primary function is as an antidote in both tissues. Crucially, there is no evidence 344 in the literature that CifB is transferred to the embryo, and we have proposed a HM model wherein 345 CifA is both a primary inducer of CI in testes and rescue in embryos, while CifB is an accessory 346 protein; this scenario enables a simple, one-step mutation scenario to the evolution of bidirectional 347 CI between different strains of Wolbachia (Shropshire et al., 2018). More work will be necessary 348 to understand CifA's cell biology and biochemistry to support it.

In the context of CifA, mutations in the N-terminal region completely ablate both CI and 349 350 rescue. Intriguingly, prior selection analyses revealed that this region is under stronger purifying 351 selection than the C-terminal end of the protein, suggesting that conservation in CifA's N-terminus 352 may have a strong impact on its phenotypic output (Shropshire et al., 2018). Given that these Nterminal regions in both CifA and CifB are unannotated, we cannot yet provide a specific 353 explanation for why these regions ablate CI and rescue. However, it is notable that N-terminal 354 355 regions in wPip homologs of both CifA and CifB are predicted to encode ankyrin-interacting domains (Bonneau et al., 2018). Thus, these mutations may be responsible for ablating the 356 binding capacity of Cif proteins to host ligands through either sequence or structural modifications 357 358 of the protein, which requires further study.

When conserved sites within CifA's putative catalase-rel domain were mutated and dually expressed with CifB in males, 34.05% of embryos died indicating that CI capability was inhibited but not ablated. Similarly, when expressed in females crossed to CI-inducing CifA;B males, only 34.05% of embryos survived, indicating that rescue was also significantly weaker. Thus, these

363 amino acids are important for causing complete CI and rescue phenotypes. Catalases are 364 enzymes that are involved in the decomposition of hydrogen peroxide and protect cells from 365 reactive oxygen species (ROS) damage (Loew, 1900). Some catalase-like domains are involved in host immune pathways that use ROS to combat disease (Govind, 2008; Zug and Hammerstein, 366 2015), and high levels of ROS can cause male infertility in organisms as diverse as Drosophila 367 and humans (Homa et al., 2015; Yu and Huang, 2015). Notably, while CifA is annotated with a 368 369 catalase-rel domain (Lindsey et al., 2018), the closest sequence homolog is from Helicobacter 370 pylori, which shares only ~22% sequence identity (Guy et al., 2005) and has no obvious active sites (Lindsey et al., 2018). Thus, it is unknown if CifA's catalase-rel is capable of degrading ROS, 371 372 but it may otherwise attract or interact with ROS while not degrading them. For example, oxidative 373 posttranslational modifications (PTM) can shift phenotypic output (Cai and Yan, 2013). CifA's 374 catalase-rel may attract ROS, enabling PTM of itself, CifB, or other host targets. Since oxidative PTMs can be reversible (Cai and Yan, 2013), rescue may in part occur through the removal of 375 376 these PTMs in the embryo. Alternatively, CifA may help to localize ROS to host targets to induce 377 oxidative damage or otherwise modify host targets.

378 Moreover, Wolbachia presence is correlated with increases in ROS in D. melanogaster, D. simulans, A. albopictus, A. polynesiensis, and T. urticae (Brennan et al., 2012, 2008; Zug and 379 380 Hammerstein, 2015), yielding additional support for CifA's function as a catalase-related protein 381 (Lindsey et al., 2018). Wolbachia-induced increases in ROS levels correlate with DNA damage in 382 D. simulans spermatocytes (Brennan et al., 2012) and an increase in lipid hydroperoxides in D. melanogaster, which are markers for ROS-induced oxidative damage (Driver et al., 2004). 383 384 Intriguingly, the immune-related gene kenny (key) is upregulated in Wolbachia-infected D. 385 melanogaster, and experimental upregulation of key in uninfected male flies yielded increased 386 ROS levels, DNA damage, and decreased hatching that can be rescued when mated to infected females (Biwot et al., 2019). Together, these data support a role for ROS in Cl's mechanism, but 387 more work is necessary to link CifA's catalase-rel domain to ROS variation and determine if ROS 388 389 are directly responsible for Cl/rescue-induction or are otherwise a symptom of other modifications 390 in gametogenesis.

The DUF in CifA shares distant homology to Puf-family RNA-binding proteins and is the only putative domain shared in all five CifA clades (Bing et al., 2020). Notably, mutating conserved residues in CifA's DUF domain revealed their specific importance to CI-induction but not for rescue. As such, this was the only domain in CifA wherein mutated sites were differentially important between the two phenotypes. RNA-binding proteins are important in transcriptional regulation and can influence the stability, localization, and translation of bound RNA. Puf-family

397 RNA-binding proteins typically influence the stability of mRNAs involved in cell maintenance, 398 embryonic development, and other processes (Forbes and Lehmann, 1998; Macdonald, 1992; 399 Parisi and Lin, 1999). For example, the Drosophila Puf-family RNA Pumilio (pum) is crucially involved in the establishment of patterning and abdominal segmentation in early embryonic 400 development by suppressing the translation of maternal hunchback RNA in the Drosophila 401 embryo (Forbes and Lehmann, 1998; Weidmann and Goldstrohm, 2012). Moreover, pum in 402 403 spermatogenesis negatively regulates the expression of p53 which is involved in DNA repair and 404 apoptosis, increases apoptosis, and reduces sperm production and fertility (Chen et al., 2012). 405 Intriguingly, mitochondrial protein p32 is a candidate suppressor of CI based on in vitro pull-down 406 assays using Drosophila lysates, and p32 regulates p53 activation (Ghate et al., 2019), 407 suggesting that *pum* in spermatogenesis may influence similar pathways in CI. Additionally, 408 Wolbachia infection has been shown to have a considerable impact on the fly transcriptome including sRNA profiles (Baião et al., 2019; Pinto et al., 2013; Y. Zheng et al., 2019). On their 409 410 own, these correlations do not sufficiently link transcription with CI. However, as described above, 411 key is significantly upregulated and causes rescuable hatch rate defects when experimentally 412 overexpressed (Biwot et al., 2019). Additionally, Wolbachia upregulate the sRNA nov-miR-12 413 which negatively regulates *pipsqueak* (*psg*), a DNA-binding protein that impacts chromatin 414 structure (Horowitz and Berg, 1996; Siegmund and Lehmann, 2002), and knockdown of psq 415 causes CI-like embryonic abnormalities and hatch rates in D. melanogaster (Y. Zheng et al., 416 2019). Thus, CifA's DUF may influence the expression of RNAs involved in the CI pathway, and by mutating the conserved site it may ablate the domain's ability to regulate these RNAs. It is also 417 418 important to note that since the DUF mutant only prevented CifA from contributing to CI, it 419 supports prior hypotheses that CifA has distinct mechanistic input to CI and rescue (Shropshire 420 and Bordenstein, 2019). The phenotypic plasticity of CifA may be caused by distinct protein 421 conformations in testes and ovaries or DUF-associated targets may only be present in testes. 422 More work will be necessary to confirm that CifA can bind RNAs and what impact this binding has 423 on downstream processes.

The final CifA domain shares homology to STE transcription factor proteins which are found predominantly in fungi and encode a sequence-specific DNA-binding motif that influences yeast reproduction through pheromone-responsive elements (Wong Sak Hoi and Dumas, 2010). Mutation of conserved sites within the STE had no impact on either CI or rescue. This was surprising since the STE domain appeared structurally conserved across four of the five CifA phylogenetic Types (Bing et al., 2020; Lindsey et al., 2018). However, while the sites mutated in the domain were conserved, the remainder of the domain appears to evolving mostly neutrally

431 (Shropshire et al., 2018), suggesting the domain may be of lesser importance than N-terminal 432 regions. Alternatively, since transgenes are expressed via host transcription and translation 433 machinery, transgenic expression bypasses the need to export the proteins outside of Wolbachia and through the host-derived membranes that surround Wolbachia within the cell (Cho et al., 434 2011; Fattouh et al., 2019). As such, it is possible that the STE domain has an essential functional 435 role in translation initiation and/or in protein export within Wolbachia, which may or may not be 436 437 related to CI and/or rescue induction. Additional research will be necessary to determine if any 438 component of the STE domain is necessary for CI and whether this domain is essential when 439 expressed inside Wolbachia.

440 Type I CifB have two domains with homologs in the PDDEXK nuclease family, but they do 441 not encode a canonical PD-(D/E)XK catalytic motif (Knizewski et al., 2007; Lindsey et al., 2018). 442 This nuclease family is heavily involved in DNA restriction, repair, recombination, and binding (Knizewski et al., 2007). Mutations in conserved sites in either domain of Type I CifB<sub>wMel</sub> ablated 443 444 CI phenotypes. Interestingly, homologs of CifB proteins from all five phylogenetic Types harbor 445 putative nuclease domains (Lindsey et al., 2018), and the Type IV CifB have functional nucleases 446 with canonical catalytic motifs and can induce CI upon dual expression with CifA (Chen et al., 447 2019). However, it remains biochemically unclear how these DNA nicks contribute to wild-type CI 448 induction, and how this can be rescued by CifA expressing females. Moreover, while we show 449 here that mutating conserved residues in either PDDEXK domain ablates CI induction, this does 450 not confirm its role as a nuclease. For instance, these domains may be essential for the 451 localization of Cif proteins to host DNA or other host targets. More work will be necessary to 452 determine if CifB's PDDEXK domains are indeed active nucleases and why mutating these 453 conserved sites ablates CI. Though, we note that ablation of CI by site-specific mutagenesis 454 across the Cif proteins highlights the utility of a mechanistically-agnostic, gene nomenclature, like the CI factor (cif) gene nomenclature, as it is increasingly clear that CifB's role as a deubiquitinase 455 or nuclease alone does not define its role in CI (Beckmann et al., 2019a; Shropshire et al., 2019). 456

457 CifB's final domain is a Ulp1 domain that contains the only known catalytic motif within the 458 Cif proteins and is responsible for the deubiquitinase activity observed in vitro (Beckmann et al., 459 2017). Previous reports show that mutating the conserved cysteine active site ablates CI function 460 in CifB<sub>wPip</sub> (Beckmann et al., 2017). Here, we confirm that mutating the same cysteine active site 461 ablated CI in the CifB<sub>wMel</sub> protein. However, it is premature to claim that the Ulp1 domain is a "catalytic warhead" for CI (Beckmann et al., 2017) because several sites, when mutated, ablate 462 463 the CI phenotype. Instead, it is evident that this site in the UIp1 domain plays an important role in 464 Cl-induction but, when analyzed in context of results from mutating other regions, it is not the only

465 crucially important component of the protein. More work will be necessary to dissect the relative 466 importance of the unannotated region, the nuclease domains, and the Ulp1 in Cl's mechanism.

467 In conclusion, we report conserved amino acids in CifA and CifB that are essential for CI and rescue phenotypes. For CifA, conserved sites in the unannotated region and catalase-rel 468 469 domain were important for CifA-induced CI and rescue, while the mutated sites in the DUF was 470 specifically important to CI. For CifB, mutating conserved sites in an unannotated region, both 471 PDDEXK nuclease domains and the Ulp1 domain were important in CifB-induced CI. These works 472 provide additional support for the necessity of expressing both CifA and CifB proteins to cause CI and the importance of CifA's complex functional repertoire and new essential regions in CifB to 473 474 the genotype-phenotype relationship and mechanism underpinning CI.

475

### 476 Materials and methods

#### 477 <u>Creating transgenic flies.</u>

cifA<sub>1</sub> and cifB<sub>1</sub> mutant transgene variants were synthesized de novo at GenScript and 478 479 cloned into a pUC57 plasmid. Site-directed mutagenesis was then performed by GenScript to 480 produce the remaining three mutant variants of each gene (Fig 2A, 3A). UAS transgenic *cifA* and cifB mutant flies were then generated following previously described protocols (LePage et al., 481 2017). Briefly, each gene was subcloned into the pTIGER plasmid, which is a pUASp-based 482 483 vector designed for germline expression. *cifA* and *cifB* transgenes were integrated into the *attp40* and attp2 attachment sites in the D. melanogaster genome using PhiC31 integrase via embryonic 484 485 injections at BestGene (LePage et al., 2017). All wild-type and transgenic nucleotide and amino 486 acid sequences are reported in Table S3.

487

### 488 Fly rearing and strains.

D. melanogaster stocks  $y^1w^*$  (BDSC 1495), nos-GAL4:VP16 (BDSC 4937), and UAS 489 490 transgenic lines homozygous for *cifA*, *cifA* mutants, *cifB*, *cifB* mutants, *cifA*;*B*, and lines dual homozygous for *cifA* or *cifB* mutants with wild-type counterparts were maintained on a 12-hour 491 492 light/dark cycle at 25°C on 50mL of standard media. Dual transgenic lines were generated through standard genetic crossings and were all homozygous viable. Uninfected lines were produced by 493 494 tetracycline treatment as previously described (LePage et al., 2017). Infection status for all lines was regularly confirmed by PCR using Wolb F and Wolb R3 primers (Casiraghi et al., 2005). 495 Genotyping was confirmed by PCR and Sanger sequencing using the primers in Table S4. 496

497

498 <u>CI measurement assays.</u>

499 CI was measured using hatch rate assays. To control for the paternal grandmother age 500 effect on CI (Layton et al., 2019), virgin nos-GAL4:VP16 females were collected for the first 3 501 days of emergence and aged 9-11 days before crossing to nonvirgin UAS transgenic males. Collections for maternal and paternal lineages were separated by a 7-day period. Individual male 502 503 and female mating occurred in 8-oz Drosophila stock bottles with a grape-juice agar plate smeared with yeast and secured to the opening of each bottle with tape. Only the first emerging 504 505 and youngest males were used to control for the younger brother effect and age effects on CI 506 (Reynolds and Hoffmann, 2002; Yamada et al., 2007). Grape-juice agar plates were produced as 507 previously described (LePage et al., 2017). The flies and bottles were incubated at 25°C for 24 508 hours, at which time the grape plates were replaced with fresh plates and stored for an additional 509 24 hours. After this, the initial number of embryos on each plate were counted. The plates were 510 incubated at 25°C and after 30 hours, the number of unhatched embryos was counted. The 511 percentage of embryos that hatched was calculated by dividing the number of hatched embryos 512 by the total number of embryos and multiplying by 100. Plates with fewer than 25 embryos were 513 excluded from analysis as previously described (LePage et al., 2017; Shropshire et al., 2018).

514

#### 515 <u>Selection analysis.</u>

Selection analysis of CifB was conducted using a sliding window analysis of K<sub>a</sub> and K<sub>s</sub> (SWAKK). The SWAKK 2.1 webserver first generated a 1196 codon alignment (with gaps) between *cifB<sub>wMel</sub>* and *cifB<sub>wHa</sub>* using ClustalW2 and then calculated the Ka/Ks ratio across the gene alignment using a sliding window of 25 codons and a jump size of 1 codon for SWAKK. The *cifA* SWAKK analysis was previously published and is based on the same parameters described above for cifB (Shropshire et al., 2018).

522

### 523 <u>Predicting mutational impact on protein structure.</u>

The effect of mutations on protein structure was evaluated with the I-TASSER protein 524 525 prediction tool (Zhang, 2009). I-TASSER generated protein tertiary structure predictions for Cif 526 proteins and their mutants using the on-line server with default settings. Structures are build based on the top ten hits generated by querying the PDB. Hits were provided Z-scores that 527 528 characterize the similarity to the query sequence. Higher Z-scores represent more confident 529 matches. Z-scores are reported in the source data provided with this manuscript and PDB hits 530 are detailed in Table S2. C-scores and TM-scores were generated for each tertiary structure. C-531 scores range from -5 to 2 where 2 is the highest confidence. TM-scores range from 0-1 where 1 532 is the highest confidence.

533

### 534 Statistical analysis.

All statistical analyses for hatch rates were conducted in GraphPad Prism 8. Hatch rate statistical comparisons were made using Kruskal-Wallis followed by a Dunn's multiple comparison test. All p-values from statistical comparisons are provided in Table S1. Figure aesthetics were edited using Affinity Designer.

539

## 540 Data availability

- All data generated in this study are available in the supplement of this manuscript.
- 542

# 543 Acknowledgments

544 We thank members of the Bordenstein lab, especially Brittany Leigh, for helpful comments and critiques during the course of this study. This work was supported by National Institutes of Health 545 546 awards R01 AI132581 and R01 AI143725, National Science Foundation award IOS 1456778, 547 and the Vanderbilt Microbiome Initiative to S.R.B., and a National Science Foundation Graduate Research Fellowship DGE-144519 to J.D.S. Any opinion, findings, and conclusions or 548 549 recommendations expressed in this material are those of the authors(s) and do not necessarily 550 reflect the views of the National Institutes of Health, the National Science Foundation, or Vanderbilt University. 551

552

## 553 Competing interests

554 J.D.S. and S.R.B. are listed as inventors on a provisional patent relevant to this work. S.R.B. is a 555 coinventor on two other pending patents related to controlling arthropods.

556

## 557 Author roles

J.D.S: Conceptualization, Formal Analysis, Funding Acquisition, Investigation, Methodology,
Supervision, Validation, Visualization, Writing – Original Draft Preparation, Writing – Review &
Editing. M.K.: Formal Analysis, Investigation, Writing – Review & Editing. S.R.B.:
Conceptualization, Funding Acquisition, Methodology, Supervision, Writing – Original Draft
Preparation, Writing – Review & Editing

### 565 References

- Aliota, M.T., Peinado, S.A., Velez, I.D., Osorio, J.E., 2016. The *w*Mel strain of *Wolbachia* reduces
   transmission of Zika virus by *Aedes aegypti*. Sci. Rep. 6, 28792.
   https://doi.org/10.1038/srep28792
- Baião, G.C., Schneider, D.I., Miller, W.J., Klasson, L., 2019. The effect of *Wolbachia* on gene expression in
   *Drosophila paulistorum* and its implications for symbiont-induced host speciation. BMC
   Genomics 20, 465. https://doi.org/10.1186/s12864-019-5816-9
- Beckmann, J., Bonneau, M., Chen, H., Hochstrasser, M., Poinsot, D., Merçot, H., Weill, M., Sicard, M.,
   Charlat, S., 2019a. The Toxin–Antidote Model of Cytoplasmic Incompatibility: Genetics and
   Evolutionary Implications. Trends Genet. https://doi.org/10.1016/j.tig.2018.12.004
- 575 Beckmann, J., Ronau, J.A., Hochstrasser, M., 2017. A *Wolbachia* deubiquitylating enzyme induces 576 cytoplasmic incompatibility. Nat. Microbiol. 2, 17007.
- 577 https://doi.org/10.1038/nmicrobiol.2017.7
- Beckmann, J., Sharma, G.D., Mendez, L., Chen, H., Hochstrasser, M., 2019b. The *Wolbachia* cytoplasmic
   incompatibility enzyme CidB targets nuclear import and protamine-histone exchange factors.
   eLife 8, e50026. https://doi.org/10.7554/eLife.50026
- Bing, X.-L., Zhao, D.-S., Sun, J.-T., Zhang, K.-J., Hong, X.-Y., 2020. Genomic analysis of *Wolbachia* from
   *Laodelphax striatellus* (Delphacidae, Hemiptera) reveals insights into its "Jekyll and Hyde" mode
   of infection pattern. Genome Biol. Evol. https://doi.org/10.1093/gbe/evaa006
- Biwot, J.C., Zhang, H.-B., Liu, C., Qiao, J.-X., Yu, X.-Q., Wang, Y.-F., 2019. *Wolbachia*-induced expression
  of kenny gene in testes affects male fertility in *Drosophila melanogaster*. Insect Sci.
  https://doi.org/10.1111/1744-7917.12730
- Bonneau, M., Atyame, C., Beji, M., Justy, F., Cohen-Gonsaud, M., Sicard, M., Weill, M., 2018. *Culex pipiens* crossing type diversity is governed by an amplified and polymorphic operon of
   *Wolbachia*. Nat. Commun. 9. https://doi.org/10.1038/s41467-017-02749-w
- 590Bordenstein, S.R., O'Hara, F.P., Werren, J.H., 2001. Wolbachia-induced incompatibility precedes other591hybrid incompatibilities in Nasonia. Nature 409, 707–710. https://doi.org/10.1038/35055543
- Brennan, L.J., Haukedal, J.A., Earle, J.C., Keddie, B., Harris, H.L., 2012. Disruption of redox homeostasis
   leads to oxidative DNA damage in spermatocytes of *Wolbachia*-infected *Drosophila simulans*.
   Insect Mol. Biol. 21, 510–520. https://doi.org/10.1111/j.1365-2583.2012.01155.x
- Brennan, L.J., Keddie, B.A., Braig, H.R., Harris, H.L., 2008. The endosymbiont *Wolbachia pipientis* induces
   the expression of host antioxidant proteins in an *Aedes albopictus* cell line. PloS One 3, e2083.
   https://doi.org/10.1371/journal.pone.0002083
- 598 Brucker, R.M., Bordenstein, S.R., 2012. Speciation by symbiosis. Trends Ecol. Evol. 27, 443–451. 599 https://doi.org/10.1016/j.tree.2012.03.011
- Cai, Z., Yan, L.-J., 2013. Protein Oxidative Modifications: Beneficial Roles in Disease and Health. J.
   Biochem. Pharmacol. Res. 1, 15–26.
- Caragata, E.P., Dutra, H.L.C., Moreira, L.A., 2016. Inhibition of Zika virus by *Wolbachia* in *Aedes aegypti*.
   Microb. Cell 3, 293–295. https://doi.org/10.15698/mic2016.07.513
- Casiraghi, M., Bordenstein, S.R., Baldo, L., Lo, N., Beninati, T., Wernegreen, J.J., Werren, J.H., Bandi, C.,
   2005. Phylogeny of *Wolbachia pipientis* based on gltA, groEL and ftsZ gene sequences: clustering
   of arthropod and nematode symbionts in the F supergroup, and evidence for further diversity in
   the *Wolbachia* tree. Microbiol.-Sgm 151, 4015–4022. https://doi.org/10.1099/mic.0.28313-0
- 608 Charlesworth, J., Weinert, L.A., Araujo, E.V., Welch, J.J., 2019. *Wolbachia, Cardinium* and climate: an 609 analysis of global data. Biol. Lett. 15, 20190273. https://doi.org/10.1098/rsbl.2019.0273

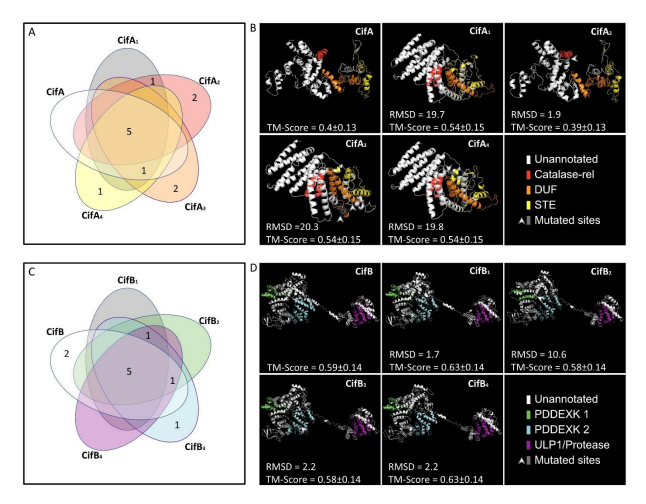
610 Chen, D., Zheng, W., Lin, A., Uyhazi, K., Zhao, H., Lin, H., 2012. Pumilio 1 Suppresses Multiple Activators 611 of p53 to Safeguard Spermatogenesis. Curr. Biol. 22, 420–425. 612 https://doi.org/10.1016/j.cub.2012.01.039 613 Chen, H., Ronau, J.A., Beckmann, J., Hochstrasser, M., 2019. A Wolbachia nuclease and its binding 614 partner provide a distinct mechanism for cytoplasmic incompatibility. Proc. Natl. Acad. Sci. 116, 615 22314–22321. https://doi.org/10.1073/pnas.1914571116 616 Cho, K.-O., Kim, G.-W., Lee, O.-K., 2011. Wolbachia bacteria reside in host Golgi-related vesicles whose 617 position is regulated by polarity proteins. PloS One 6, e22703. 618 https://doi.org/10.1371/journal.pone.0022703 619 Cunningham, B.C., Wells, J.A., 1989. High-resolution epitope mapping of hGH-receptor interactions by 620 alanine-scanning mutagenesis. Science 244, 1081–1085. 621 https://doi.org/10.1126/science.2471267 Driver, C., Georgiou, A., Georgiou, G., 2004. The contribution by mitochondrially induced oxidative 622 623 damage to aging in *Drosophila melanogaster*. Biogerontology 5, 185–192. 624 https://doi.org/10.1023/B:BGEN.0000031156.75376.e3 625 Duffy, J.B., 2002. GAL4 system in Drosophila: a fly geneticist's Swiss army knife. Genes. N. Y. N 2000 34, 626 1–15. https://doi.org/10.1002/gene.10150 627 Fattouh, N., Cazevieille, C., Landmann, F., 2019. Wolbachia endosymbionts subvert the endoplasmic 628 reticulum to acquire host membranes without triggering ER stress. PLoS Negl. Trop. Dis. 13, 629 e0007218. https://doi.org/10.1371/journal.pntd.0007218 630 Ferree, P.M., Sullivan, W., 2006. A genetic test of the role of the maternal pronucleus in Wolbachiainduced cytoplasmic incompatibility in Drosophila melanogaster. Genetics 173, 839–847. 631 632 https://doi.org/10.1534/genetics.105.053272 Forbes, A., Lehmann, R., 1998. Nanos and Pumilio have critical roles in the development and function of 633 634 Drosophila germline stem cells. Dev. Camb. Engl. 125, 679–690. 635 Ghate, N.B., Kim, J., Shin, Y., Situ, A., Ulmer, T.S., An, W., 2019. p32 is a negative regulator of p53 636 tetramerization and transactivation. Mol. Oncol. 13, 1976–1992. https://doi.org/10.1002/1878-637 0261.12543 638 Govind, S., 2008. Innate immunity in *Drosophila*: Pathogens and pathways. Insect Sci. Online 15, 29–43. 639 https://doi.org/10.1111/j.1744-7917.2008.00185.x 640 Guy, B., Krell, T., Sanchez, V., Kennel, A., Manin, C., Sodoyer, R., 2005. Do Th1 or Th2 sequence motifs 641 exist in proteins?: Identification of amphipatic immunomodulatory domains in Helicobacter 642 pylori catalase. Immunol. Lett. 96, 261–275. https://doi.org/10.1016/j.imlet.2004.09.011 Hilgenboecker, K., Hammerstein, P., Schlattmann, P., Telschow, A., Werren, J.H., 2008. How many 643 644 species are infected with Wolbachia? - a statistical analysis of current data. Fems Microbiol. Lett. 645 281, 215–220. https://doi.org/10.1111/j.1574-6968.2008.01110.x 646 Homa, S.T., Vessey, W., Perez-Miranda, A., Riyait, T., Agarwal, A., 2015. Reactive Oxygen Species (ROS) in 647 human semen: determination of a reference range. J. Assist. Reprod. Genet. 32, 757–764. 648 https://doi.org/10.1007/s10815-015-0454-x 649 Horowitz, H., Berg, C.A., 1996. The Drosophila pipsqueak gene encodes a nuclear BTB-domain-containing 650 protein required early in oogenesis. Dev. Camb. Engl. 122, 1859–1871. 651 Jaenike, J., Dyer, K.A., Cornish, C., Minhas, M.S., 2006. Asymmetrical reinforcement and Wolbachia infection in *Drosophila*. PLoS Biol. 4, e325. https://doi.org/10.1371/journal.pbio.0040325 652 653 Kittayapong, P., Kaeothaisong, N.-O., Ninphanomchai, S., Limohpasmanee, W., 2018. Combined sterile 654 insect technique and incompatible insect technique: sex separation and quality of sterile Aedes 655 *aegypti* male mosquitoes released in a pilot population suppression trial in Thailand. Parasit. 656 Vectors 11, 657. https://doi.org/10.1186/s13071-018-3214-9

657 Knizewski, L., Kinch, L.N., Grishin, N.V., Rychlewski, L., Ginalski, K., 2007. Realm of PD-(D/E)XK nuclease 658 superfamily revisited: detection of novel families with modified transitive meta profile searches. 659 BMC Struct. Biol. 7, 40. https://doi.org/10.1186/1472-6807-7-40 660 Kumar, G.A., Subramaniam, K., 2018. PUF-8 facilitates homologous chromosome pairing by promoting 661 proteasome activity during meiotic entry in C. elegans. Development dev.163949. 662 https://doi.org/10.1242/dev.163949 Layton, E.M., On, J., Perlmutter, J.I., Bordenstein, S.R., Shropshire, J.D., 2019. Paternal grandmother age 663 664 affects the strength of Wolbachia-induced cytoplasmic incompatibility in Drosophila 665 melanogaster. mBio 10. https://doi.org/10.1128/mBio.01879-19 666 LePage, D., Bordenstein, S.R., 2013. Wolbachia: Can we save lives with a great pandemic? Trends 667 Parasitol. 29, 385–393. https://doi.org/10.1016/j.pt.2013.06.003 668 LePage, D.P., Metcalf, J.A., Bordenstein, Sarah R., On, J., Perlmutter, J.I., Shropshire, J.D., Layton, E.M., Funkhouser-Jones, L.J., Beckmann, J., Bordenstein, Seth R., 2017. Prophage WO genes 669 670 recapitulate and enhance Wolbachia-induced cytoplasmic incompatibility. Nature 543, 243–247. 671 https://doi.org/10.1038/nature21391 672 Lindsey, A., Rice, D.W., Bordenstein, Sarah R., Brooks, A.W., Bordenstein, Seth R., Newton, I.L.G., 2018. Evolutionary genetics of cytoplasmic incompatibility genes *cifA* and *cifB* in prophage WO of 673 674 Wolbachia. Genome Biol. Evol. 10, 434–451. https://doi.org/10/gcvmkm 675 Loew, O., 1900. A new enzyme of general occurrence in organisms. Science 11, 701–702. 676 https://doi.org/10.1126/science.11.279.701 677 Macdonald, P.M., 1992. The Drosophila pumilio gene: an unusually long transcription unit and an 678 unusual protein. Dev. Camb. Engl. 114, 221–232. Mi, D., Ou, X., Li, P., Peng, G., Liu, Y., Guo, R., Mu, Z., Li, F., Holmes, K., Qian, Z., 2019. Glycine 29 Is 679 680 Critical for Conformational Changes of the Spike Glycoprotein of Mouse Hepatitis Virus A59 681 Triggered by either Receptor Binding or High pH. J. Virol. 93. https://doi.org/10.1128/JVI.01046-682 19 683 Moreira, L.A., Iturbe-Ormaetxe, I., Jeffery, J.A., Lu, G., Pyke, A.T., Hedges, L.M., Rocha, B.C., Hall-684 Mendelin, S., Day, A., Riegler, M., Hugo, L.E., Johnson, K.N., Kay, B.H., McGraw, E.A., van den 685 Hurk, A.F., Ryan, P.A., O'Neill, S.L., 2009. A Wolbachia symbiont in Aedes aegypti limits infection 686 with dengue, Chikungunya, and Plasmodium. Cell 139, 1268–1278. 687 https://doi.org/10.1016/j.cell.2009.11.042 688 Nishanth, M.J., Simon, B., 2020. Functions, mechanisms and regulation of Pumilio/Puf family RNA 689 binding proteins: a comprehensive review. Mol. Biol. Rep. 47, 785–807. https://doi.org/10.1007/s11033-019-05142-6 690 691 O'Connor, L., Plichart, C., Sang, A.C., Brelsfoard, C.L., Bossin, H.C., Dobson, S.L., 2012. Open release of 692 male mosquitoes infected with a Wolbachia biopesticide: field performance and infection 693 containment. PLoS Negl. Trop. Dis. 6, e1797. https://doi.org/10.1371/journal.pntd.0001797 694 O'Neill, S.L., 2018. The use of Wolbachia by the World Mosquito Program to interrupt transmission of 695 Aedes aegypti transmitted viruses. Adv. Exp. Med. Biol. 1062, 355–360. https://doi.org/10.1007/978-981-10-8727-1 24 696 Parisi, M., Lin, H., 1999. The Drosophila pumilio gene encodes two functional protein isoforms that play 697 698 multiple roles in germline development, gonadogenesis, oogenesis and embryogenesis. Genetics 699 153, 235-250. 700 Pinto, S.B., Stainton, K., Harris, S., Kambris, Z., Sutton, E.R., Bonsall, M.B., Parkhill, J., Sinkins, S.P., 2013. 701 Transcriptional regulation of *Culex pipiens* mosquitoes by *Wolbachia* influences cytoplasmic 702 incompatibility. PLoS Pathog. 9, e1003647. https://doi.org/10.1371/journal.ppat.1003647

703	Rasgon, J.L., 2008. Using Predictive Models to Optimize <i>Wolbachia</i> -Based Strategies for Vector-Borne
704	Disease Control, in: Aksoy, S. (Ed.), Transgenesis and the Management of Vector-Borne Disease.
705	Springer New York, New York, NY, pp. 114–125. https://doi.org/10.1007/978-0-387-78225-6_10
706	Reynolds, K.T., Hoffmann, A.A., 2002. Male age, host effects and the weak expression or non-expression
707	of cytoplasmic incompatibility in Drosophila strains infected by maternally transmitted
708	Wolbachia. Genet. Res. 80, 79–87.
709	Serbus, L.R., Casper-Lindley, C., Landmann, F., Sullivan, W., 2008. The genetics and cell biology of
710	Wolbachia-host interactions. Annu. Rev. Genet. 42, 683–707.
711	https://doi.org/10.1146/annurev.genet.41.110306.130354
712	Shropshire, J.D., Bordenstein, S.R., 2019. Two-by-one model of cytoplasmic incompatibility: synthetic
713	recapitulation by transgenic expression of <i>cifA</i> and <i>cifB</i> in <i>Drosophila</i> . PLOS Genet. 15,
714	e1008221. https://doi.org/10.1371/journal.pgen.1008221
715	Shropshire, J.D., Bordenstein, S.R., 2016. Speciation by symbiosis: the microbiome and behavior. mBio 7,
716	e01785-15. https://doi.org/10.1128/mBio.01785-15
717	Shropshire, J.D., Leigh, B., Bordenstein, Sarah R., Duplouy, A., Riegler, M., Brownlie, J.C., Bordenstein,
718	Seth R., 2019. Models and nomenclature for cytoplasmic incompatibility: caution over
719	premature conclusions – a response to Beckmann et al. Trends Genet. 0.
720	https://doi.org/10.1016/j.tig.2019.03.004
721	Shropshire, J.D., On, J., Layton, E.M., Zhou, H., Bordenstein, S.R., 2018. One prophage WO gene rescues
722	cytoplasmic incompatibility in <i>Drosophila melanogaster</i> . Proc. Natl. Acad. Sci. U. S. A. 115, 4987–
723	4991. https://doi.org/10.1073/pnas.1800650115
724	Siegmund, T., Lehmann, M., 2002. The <i>Drosophila</i> Pipsqueak protein defines a new family of helix-turn-
725	helix DNA-binding proteins. Dev. Genes Evol. 212, 152–157. https://doi.org/10.1007/s00427-
726	002-0219-2
727	Taylor, M.J., Bordenstein, S.R., Slatko, B., 2018. Microbe Profile: <i>Wolbachia</i> : a sex selector, a viral
728	protector and a target to treat filarial nematodes. Microbiology 164, 1345–1347.
729	https://doi.org/10.1099/mic.0.000724
730	Tram, U., Sullivan, W., 2002. Role of delayed nuclear envelope breakdown and mitosis in <i>Wolbachia</i> -
731	induced cytoplasmic incompatibility. Science 296, 1124–1126.
732	https://doi.org/10.1126/science.1070536
733	Turelli, M., 1994. Evolution of incompatibility-inducing microbes and their hosts. Evol. Int. J. Org. Evol.
734	48, 1500–1513. https://doi.org/10.1111/j.1558-5646.1994.tb02192.x
735	Weidmann, C.A., Goldstrohm, A.C., 2012. <i>Drosophila</i> Pumilio Protein Contains Multiple Autonomous
736	Repression Domains That Regulate mRNAs Independently of Nanos and Brain Tumor. Mol. Cell.
737	Biol. 32, 527–540. https://doi.org/10.1128/MCB.06052-11
738	Weinert, L.A., Araujo-Jnr, E.V., Ahmed, M.Z., Welch, J.J., 2015. The incidence of bacterial endosymbionts
739	in terrestrial arthropods. Proc R Soc B 282, 20150249. https://doi.org/10.1098/rspb.2015.0249
740	Wong Sak Hoi, J., Dumas, B., 2010. Ste12 and Ste12-Like Proteins, Fungal Transcription Factors
741	Regulating Development and Pathogenicity. Eukaryot. Cell 9, 480–485.
742	https://doi.org/10.1128/EC.00333-09
743	Yamada, R., Floate, K.D., Riegler, M., O'Neill, S.L., 2007. Male development time influences the strength
744	of Wolbachia-induced cytoplasmic incompatibility expression in Drosophila melanogaster.
745	Genetics 177, 801–808. https://doi.org/10.1534/genetics.106.068486
746	Yang, H., Ahmad, Z.A., Song, Y., 2020. Molecular insight for the role of key residues of calreticulin in its
740	binding activities: A computational study. Comput. Biol. Chem. 85, 107228.
748	https://doi.org/10.1016/j.compbiolchem.2020.107228
749	Yu, B., Huang, Z., 2015. Variations in Antioxidant Genes and Male Infertility. BioMed Res. Int. 2015.
750	https://doi.org/10.1155/2015/513196
, 50	

- Zhang, Y., 2009. I-TASSER: fully automated protein structure prediction in CASP8. Proteins 77 Suppl 9,
   100–113. https://doi.org/10.1002/prot.22588
- Zheng, X., Zhang, D., Li, Y., Yang, C., Wu, Y., Liang, X., Liang, Y., Pan, X., Hu, L., Sun, Q., Wang, X., Wei, Y.,
  Zhu, J., Qian, W., Yan, Z., Parker, A.G., Gilles, J.R.L., Bourtzis, K., Bouyer, J., Tang, M., Zheng, B.,
  Yu, J., Liu, J., Zhuang, J., Hu, Zhigang, Zhang, M., Gong, J.-T., Hong, X.-Y., Zhang, Z., Lin, L., Liu, Q.,
  Hu, Zhiyong, Wu, Z., Baton, L.A., Hoffmann, A.A., Xi, Z., 2019. Incompatible and sterile insect
  techniques combined eliminate mosquitoes. Nature. https://doi.org/10.1038/s41586-019-1407-
- 758

- Zheng, Y., Shen, W., Bi, J., Chen, M.-Y., Wang, R.-F., Ai, H., Wang, Y.-F., 2019. Small RNA analysis provides
   new insights into cytoplasmic incompatibility in *Drosophila melanogaster* induced by *Wolbachia*.
   J. Insect Physiol. 118, 103938. https://doi.org/10.1016/j.jinsphys.2019.103938
- Zug, R., Hammerstein, P., 2015. Wolbachia and the insect immune system: what reactive oxygen species
   can tell us about the mechanisms of Wolbachia–host interactions. Front. Microbiol. 6.
   https://doi.org/10.3389/fmicb.2015.01201
- Zug, R., Hammerstein, P., 2012. Still a host of hosts for *Wolbachia*: analysis of recent data suggests that
   40% of terrestrial arthropod species are infected. PLOS ONE 7, e38544.
- 767 https://doi.org/10.1371/journal.pone.0038544
- 768 769



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Figure S1. Summary of I-TASSER structural predictions for CifA, CifB, and their mutants. (A, C) Venn-diagrams showing the number of PDB hits shared between wild-type and mutant (A) CifA and (C) CifB proteins. (B, D) I-TASSER uses these PDB hits to generate structural predictions for (C) CifA and (D) CifB. TM-scores range from 0-1 where 1 is the highest confidence. RMSD scores are from pairwise alignments of mutant proteins with the wild-type in PyMol. Higher RMSD scores represent more distance between the superimposed proteins. Mutated sites in the tertiary structure are indicated with a white arrow. Domain annotations were based on previous sequence analyses (Lindsey et al., 2018). Details regarding the PDB hits are reported in Table S2.

Table S1. P values associated with all statistical comparisons made in main and extended data hatch rate and cytology figures.
 M=male, F=female ,+=Wolbachia infected, -=Wolbachia uninfected.

Table S2. Protein structural prediction software I-TASSER identifies homologous protein domains found in all of our Cif homologs,
 and those that differ.

- 782 Table S3. Nucleotide and protein sequences used in this study.
- 783 **Table S4.** Primers used for genotyping and sanger sequencing if cif transgenes.
- 784 Source Data File 1. All raw data from main figures, supplemental figures, and replicate data experiments in this study.