The CinB Nuclease from wNo Wolbachia Is Sufficient for Induction of Cytoplasmic Incompatibility in Drosophila

Guangxin Sun, Mengwen Zhang, Hongli Chen, Mark Hochstrasser

ABSTRACT Wolbachia is an obligate intracellular bacterium that can alter reproduction of its arthropod hosts, often through a mechanism called cytoplasmic incompatibility (CI). In CI, uninfected females fertilized by infected males yield few offspring, but if both are similarly infected, normal embryo viability results (called "rescue"). CI factors (Cifs) responsible for CI are pairs of proteins encoded by linked genes. The downstream gene in each pair encodes either a deubiquitylase (CidB) or a nuclease (CinB). The upstream gene products, CidA and CinA, bind their cognate enzymes with high specificity. Expression of CidB or CinB in yeast inhibits growth, but growth is rescued by expression of the cognate CifA protein. By contrast, transgenic Drosophila male germ line expression of both cifA and cifB was reported to be necessary to induce CI-like embryonic arrest; cifA expression alone in females is sufficient for rescue. This pattern, seen with genes from several Wolbachia strains, has been called the "2-by-1" model. Here, we show that male germ line expression of the cinB gene alone, from a distinct clade of cif genes from wNo Wolbachia, is sufficient to induce nearly complete loss of embryo viability. This male sterility is fully rescued by cognate cinAwNo expression in the female germ line. The proteins behave similarly in yeast. CinBwNo toxicity depends on its nuclease active site. These results demonstrate that highly divergent CinB nucleases can induce CI, that rescue by cognate CifA factors is a general feature of Wolbachia CI systems, and that CifA is not strictly required in males for CI induction.

IMPORTANCE Wolbachia bacteria live within the cells of many insects. Like mitochondria, they are only inherited from females. Wolbachia often increases the number of infected females to promote spread of infection using a type of male sterility called cytoplasmic incompatibility (CI): when uninfected females mate with infected males, most embryos die; if both are similarly infected, embryos develop normally, giving infected females an advantage in producing offspring. CI is being used against disease-carrying mosquitoes and agricultural pests. Wolbachia proteins called CifA and CifB, which bind one another, cause CI, but how they work has been unclear. Here, we show that a CifB protein singly produced in fruit fly males causes sterility in crosses to normal females, but this is rescued if the females produce the CifA partner. These findings clarify a broad range of observations on CI and will allow more rational approaches to using it for insect control.

KEYWORDS Wolbachia, cytoplasmic incompatibility, CinA, CinB, nuclease, toxin-antidote
the female germ line (3). Manipulations include parthenogenesis, male killing, and feminization of chromosomal males, but the most common is a mechanism called cytoplasmic incompatibility, or CI (4). In CI, when females are infected, fertilization by either infected or uninfected males yields normal numbers of viable embryos, but if an uninfected female mates with a *Wolbachia*-infected male, a high fraction of the resulting embryos die. The ability of *Wolbachia*-infected females to “rescue” viability provides a selective advantage to infected females and can drive *Wolbachia* infections into populations. Several other bacterial species are now known to cause CI, but *Wolbachia* is the most widespread and best studied (5).

*Wolbachia*-induced CI is being used in several ways to control mosquito-vectored disease, particularly by *Aedes aegypti* mosquitoes that transmit dengue virus and other arboviruses. In one approach, introduction of large numbers of *Wolbachia*-infected male mosquitoes into uninfected mosquito populations causes massive drops in mosquito number due to the postzygotic male sterility caused by CI (6). Another control method exploits the observation that *Wolbachia* infection severely limits the ability of insects to carry certain arboviruses. In this approach, male and female *A. aegypti* mosquitoes trans-infected with *Wolbachia*, usually the wMel strain, are released together in areas with endemic arbovirus-infected mosquitoes—principally those carrying dengue virus (7, 8). Due to the ability of the wMel strain to cause CI, the bacterial infection rapidly spreads, and dramatic reductions in local dengue fever cases have been reported (7, 9).

The molecular basis of CI had long been a mystery, but recently, the *Wolbachia* factors responsible were identified (10–12). The CI factors (Cifs) are encoded by two-gene operons, usually, but not always, found in WO prophage regions (13–15), with the upstream gene designated most generally as *cisA* and the downstream gene as *cisB* (Fig. 1A). Different CI-inducing *Wolbachia* strains carry distinct but related *cisA-cisB* operons, and some strains bear multiple divergent versions, which, in some cases, involve pseudogenes (13, 14). The downstream gene in each cognate gene pair usually encodes either a deubiquitylase (DUB) that cleaves ubiquitin from substrate proteins (CI-inducing DUB, or CidB) or a DNA-cleaving enzyme (CI-inducing nuclease, or CinB) (16, 17). The wPip strain has syntenic gene pairs of both types. When either CidB<sub>wPip</sub> or CinB<sub>wPip</sub> is expressed by itself in the yeast *Saccharomyces cerevisiae*, temperature-dependent growth inhibition is observed (10). Growth impairment depends on the catalytic activity of the respective enzymes.

CidA<sub>wPip</sub> and CinA<sub>wPip</sub> bind the cognate CidB<sub>wPip</sub> and CinB<sub>wPip</sub> proteins with high affinity, but the binding does not block the enzyme active sites of the latter proteins (10, 12). Nevertheless, coexpression in yeast of the cognate CifA protein from each operon suppresses the toxicity observed when CidB or CinB is expressed by itself (10). These observations suggest that suppression results from interference of the CifA factors with CifB enzyme-substrate interactions or from enzyme relocalization within the cell. Expression of the CifA proteins by themselves is not deleterious to yeast (18).

Results from yeast growth studies strongly parallel effects seen by transgenic germ line expression of the same Cif proteins in *Drosophila melanogaster*. For example, female germ line expression of CidA<sup>*Mel*</sup> or CinA<sup>*Mel*</sup> was sufficient for rescue of transgenic CI caused by expression of the cognate *cisA-cisB* genes in the male germ line (12, 19). The enzymatic activity of the CifB enzymes is required for full CI in transgenic flies, as is true for yeast toxicity (10, 12, 20). Moreover, orthologous karyopherin-α proteins were shown to suppress CI and growth inhibition in flies and yeast, respectively (18).

By contrast, induction of CI in flies required not only expression of CifB, as would be predicted from the yeast studies, but also that of the partner CifA (12, 14). This requirement for both *cisA* and *cisB* for transgenic CI induction and *cisA* alone for rescue has been termed the “2-by-1” model (19). The dual gene requirement for induction has called into question the reliability of yeast growth assays as a surrogate for CI analysis in transgenic insects even though the CI rescue activity of CifA proteins was initially predicted from yeast studies (10).

In the current study, we analyze a divergent cinA-cinB system from the wNo
Wolbachia strain, which naturally infects *Drosophila simulans* (21). We had shown previously that wNo-infected *D. simulans* display embryonic cytological defects in incompatible crosses that are similar to those observed with incompatible transgenic *D. melanogaster* crosses involving *cidA-cidB* or *cinA-cinB* from wPip (12). However, we had not tested whether transgenic *cinA-cinB* behaves similarly. Here, we find that this is indeed the case. Unexpectedly, male expression of *cinB* alone induced strong CI in transgenic flies, and this was fully rescued by cognate *cinB* expression in the female germ line. Parallel results were seen with *cinA-cinB* expression in yeast, where it was further shown that the nuclease active site must be intact to observe toxicity. These findings indicate that catalytically active CifB is the fundamental CI-inducing factor and that CifA is the specific rescue factor. Our results also bear on current models for CI mechanisms. In particular, they argue against the generality of the 2-by-1 model and models that posit a primary role for CifA in CI induction.

**RESULTS**

Sequence comparisons of CinA and CinB homologs. As noted above, we had previously shown that the CinA and CinB factors from wPip Wolbachia are sufficient to recapitulate key features of CI in transgenic *D. melanogaster* (12). Primary sequence comparisons divide the *cidA-cidB* gene pairs from different Wolbachia strains into five distinct clades or types (22, 23). In four of these, types II to V, the downstream gene is predicted to encode an active nuclease; these operons would all be *cinA-cinB* operons in our nomenclature based on the (predicted) enzymatic activity encoded by *cinB*. The *cinB* nuclease is a type IV Cif protein. Because wPip(Pel) has both *cidA-cidB* (type I *cif*) and *cinA-cinB* gene pairs, one cannot infer which of the two, if not both, is responsible for causing CI. We therefore had turned to an analysis of *D. simulans* infected with wNo, a *Wolbachia* strain with only a single *cinA-cinB* operon, to verify that the embryonic cytological features of CI associated with a nuclease-encoding operon were similar to those seen with Wolbachia strains such as wMel, which only have a *cidA-cidB* (DUB-encoding) locus (12). However, in our previous analysis of wNo-induced CI, we had not tested whether the CinA and CinB factors from wNo could recapitulate CI transgenically.

The wNo CinA and CinB proteins have diverged considerably from the homologous proteins from wPip (Fig. 1A). Nevertheless, the nuclease catalytic residues are conserved in *cinB*, and it is also predicted to have two active PD-(D/E)XK (PDDEXK) nuclease domains (Fig. 1A and B). In the N-terminal nuclease domain (NTND) of CinB, the catalytic residues coordinate a divalent cation, whereas no cation binding was observed in the C-terminal ND (CTND) (24). Despite this, mutating catalytic residues in either domain abolishes DNase activity and blocks CI-like induction (12). The conservation of these residues in both domains among CifB types II to V (Fig. 1B) predicts that CinB will also function in CI in a way that depends on its nuclease activity.

Analysis of CinA and CinB expression in budding yeast. Growth analysis of *S. cerevisiae* expressing different Wolbachia Cif proteins has generally proven to be a reliable surrogate for transgenic CI analysis in insects (10, 12, 16). Expression of CidB proteins from wPip or wHa was found to cause temperature-sensitive growth defects in yeast that were suppressed by coexpression of their cognate CidA proteins (10, 18). The same was true for CinA and CinB expression (10). We therefore transformed yeast with plasmids carrying *cinA* or *cinB*, or both under galactose-inducible promoters and examined growth of the transformants (Fig. 1C). Galactose-induced production of CinB caused a reproducible defect in growth at higher temperatures, although the impairment was less severe than that previously seen with CinB expression. Importantly, the growth deficiency was relieved by *cinB* coexpression.

When a predicted catalytic lysine in either the NTND or CTND (K275 and K629, respectively) was mutated to alanine, yeast growth impairment was abrogated (Fig. 1C; Fig. S1 in the supplemental material). Unlike what had been observed with CinB, the active-site mutations caused a partial reduction in levels of the protein. This effect was relatively mild in the BY4741 strain background (Fig. 1D) but was more marked in a W303 strain (Fig. S1). The modest reduction of CinB mutant protein
The wNo-derived CinA and CinB proteins. (A) Comparison of the wNo and wPip CinA and CinB CI factors. The wNo proteins are from the type III clade of CI factors; the wPip proteins belong to the type IV branch. Five base pairs separate the stop codon of wNo \( \text{cinA} \) from the start codon of \( \text{cinB} \); in the wPip \( \text{cinA-cinB} \) operon, this gap is 51 bp. All CinB proteins include two intact nuclease domains (NDs), which both appear to be necessary for DNase activity and biological function. ID, identity. (B) Core sequences of the PDDEXK nuclease domains (NDs) from the four known clades of CI factors thought to encode active nucleases (types II to V). The conserved residues that constitute the active site based on the CinB\(_{\text{wPip}}\) (type IV) crystal structures (24) are indicated by arrowheads. A conserved QxxxY motif just downstream of the active site residues is marked by dots; this motif is characteristic of RecB-like

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amounts seen in the former strain would not predict complete loss of growth inhibition. Interestingly, coexpression of CinA<sub>wt</sub> reproducibly increased the levels of the cognate CinB<sub>wt</sub> proteins, whether WT or mutant. This suggests that CinB<sub>wt</sub> might be susceptible to degradation in yeast and that the (predicted) binding to CinA<sub>wt</sub> may protect it from such turnover.

In summary, these data indicate that the CinA<sub>wt</sub>-CinB<sub>wt</sub> cognate pair behaves similarly to those encoded by other Wolbachia cifA-cifB genes previously analyzed in yeast.

**Cognate-specific binding preferences of wNo and wPip CinA-CinB pairs.** The ability of CinA<sub>No</sub> to suppress the growth defect of yeast expressing CinB<sub>No</sub> (and to elevate CinB<sub>No</sub> levels) suggested that the two proteins interact, as was known to be true for CinA<sub>wPip-CinB<sub>wPip</sub></sub>, which form a tight complex (dissociation constant [K<sub>d</sub>], 25 nM) (12). To test this directly, recombinant His<sub>6</sub>-tagged CinA and glutathione S-transferase (GST)-tagged CinB proteins derived from wNo and wPip were coexpressed in *Escherichia coli* and tested for binding using GST pulldown assays (Fig. 2A). As expected, CinA<sub>wt</sub>-CinB<sub>wt</sub> showed a strong interaction (Fig. 2A, lane 1), which was likely a 1:1 complex based on our recent crystallographic study (24). Similarly, CinA<sub>No</sub>-CinB<sub>No</sub> displayed a readily detectable interaction (lane 4). In contrast, the noncognate pairs showed only weak cross-binding (lanes 2 and 3). Similar results were found when lysates from separate *E. coli* transformants expressing either recombinant CinA<sub>No</sub> or CinB<sub>No</sub> were mixed and evaluated by GST pulldown analysis (Fig. S2). Consistent with these binding data, suppression of CinB-induced yeast growth defects by CinA proteins showed cognate specificity: rescue was only seen when the CinA and CinB proteins came from the same *Wolbachia* strain (Fig. 2B).

**CinB<sub>No</sub> is sufficient for transgenic CI induction, and CinA<sub>No</sub> is sufficient for its rescue.** There are still only a few examples where the ability of specific cifA-cifB gene pairs to recapitulate CI by transgenic germ line expression in insects has been demonstrated. Moreover, it has been suggested that there may be additional *Wolbachia* genes that are responsible for CI (5, 13, 25). We therefore tested whether the highly divergent wNo-encoded CinA and CinB proteins (Fig. 1A) were indeed capable of causing CI.

In contrast to yeast growth impairment, expression of both cifA and cifB genes in the *D. melanogaster* male germ line has been reported to be necessary for triggering transgenic CI (19). Both proteins are expected to be present in mature sperm, although only CidA<sub>wt</sub> has been directly detected there (26). This has raised questions about whether CifA or CifB is the primary inducer of CI (20).

We first created transgenic flies that expressed the *cinA<sub>No</sub>-cinB<sub>No</sub>* coding sequences linked by a picornavirus T2A sequence; the latter element is expected to lead to expression of the two proteins in roughly equal amounts from a single mRNA (27). Expression of the CinA-T2A-CinB (wNo) construct in males that were crossed to wild-type (WT) females caused a nearly complete loss of viable embryos based on egg hatch rates (Fig. 3A). The *cinA<sub>No</sub>* gene by itself caused no reduction in viability, as expected, despite high relative mRNA levels (Fig. S3).

Surprisingly, expression of the *cinB<sub>No</sub>* gene alone in the male germ line induced a severe reduction in viable embryos that was indistinguishable from that induced by the *cinA<sub>No</sub>-cinB<sub>No</sub>* pair. This embryonic lethality was completely reversed by expression of *cinA<sub>No</sub>* in the female germ line (Fig. 3A). We also repeated our earlier analysis of transgenic CI in *D. melanogaster* caused by the *cinA<sub>Pip</sub>-cinB<sub>Pip</sub>* pair (12) (Fig. 3B).

**FIG 1 Legend (Continued)**

nucleases and has been suggested to function in DNA binding (33). Alignments were done with Clustal Omega, and the figure was made using MView (1.63). u, upstream or N-terminal ND; d, downstream or C-terminal ND. (C) Growth assays of yeast expressing wNo cinA and cinB alleles. BY4741 cells were transformed with plasmids expressing the indicated alleles from a galactose-inducible GAL1 promoter, and cultures were spotted in serial dilution onto selection plates with the indicated carbon source and grown for 2.5 days at either 30°C or 36°C. Red dots highlight strains expressing WT CinB<sub>wt</sub> with and without CinB<sub>wt</sub> coexpression. EV, empty vector; A, CinA<sub>wt</sub>; B, CinB<sub>wt</sub>. (D) Expression levels of different CinA and CinB proteins in the same BY4741 transformants from panel C were measured by immunoblot analysis. Both proteins were tagged with a FLAG epitope. The anti-PGK immunoblot served as a loading control.
Both cinAwNo and cinBwPip were needed for inducing strong embryonic lethality, in contrast to the sufficiency seen with cinBwPip. As expected, expression of cinAwNo in the female germ line did not rescue viability in crosses to males with the heterologous cinA-T2A-cinBwPip construct. In summary, these data strongly suggest that CI induction, at least in this transgenic model, does not require the cifA component of the cifA-cifB locus, in contrast to previous suggestions.

Although rescue of embryo viability by cinAwNo was robust and strongly implicated CI as the cause of the observed male sterility, it was important to document that the cytological features of transgenic CI induced by cinBwPip resembled natural CI rather than some other form of male sterility. Typically, CI leads to arrest during the first zygotic mitotic division due to asynchronous chromosome condensation and separation in the juxtaposed male and female pronuclei (4, 28). Not all embryos will necessarily arrest at this stage, but nuclear division stalls in a large fraction of embryos prior to the blastoderm stage (10, 11, 17).

We analyzed embryo cytology 1 to 2 h after egg deposition in transgenic CI crosses (Fig. 4A to E). Either expression of CinBwPip alone or in combination with CinAwNo in the
male germ line caused most resulting embryos to arrest early in development. Expression of CinA<sup>w<sub>No</sub></sup> in the female germ line strongly countered the arrest phenotype in both cases. When embryos from crosses between CinB<sup>w<sub>No</sub></sup> males and WT females were examined after a shorter (30-min) egg-laying period, we were able to observe first zygotic division defects; these ranged from asynchronous condensation of chromosomes between the juxtaposed male and female pronuclei to chromatin bridging in late anaphase or telophase (Fig. 4F). Therefore, the male sterility which occurs because of transgenic CinB<sup>w<sub>No</sub></sup> expression in males bears the hallmarks of natural CI previously observed in wNo-infected <i>D. simulans</i> and other examples of CI (12).

**DISCUSSION**

The current study includes several findings relevant to current models for the mechanism of <i>Wolbachia</i>-mediated CI (Fig. 5). The CinA and CinB CI factors from wNo behave in a manner very similar to the previously analyzed Cin (and Cid) proteins of wPip. Despite their strong sequence divergence, the CinB nucleases from both wNo and wPip can induce CI in fruit flies and toxicity in yeast. CinA<sup>w<sub>No</sub></sup> is sufficient for rescue of CI in transgenic <i>Drosophila</i> and suppression of toxicity in yeast. Paralleling their growth effects, the wNo CinA and CinB proteins bind one another in a cognate-specific fashion. Moreover, mutation of predicted catalytic residues of the nuclease-active centers in CinB<sup>w<sub>No</sub></sup> prevents its toxicity when expressed in yeast. Most importantly,
expression of \( \text{CinB}^{\text{wNo}} \) alone in the male germ line of transgenic flies induces highly penetrant CI. This embryonic lethality can be fully rescued by \( \text{CinA}^{\text{wNo}} \) expression in females.

One significant discrepancy in previous studies of the \( \text{cif} \) genes was the observation that yeast growth defects caused by heterologous Cif protein expression and transgenic male sterility caused by these same factors showed different \( \text{cif} \) gene dependencies. Specifically, \( \text{cifB} \)-encoded proteins were sufficient for growth impairment in yeast, whereas expression of both \( \text{cifA} \) and \( \text{cifB} \) genes in the \( \text{D. melanogaster} \) male germ line was needed to trigger transgenic CI (12, 19). On the other hand, the ability of cognate \( \text{cifA} \) genes by themselves to rescue transgenic CI matched what had been seen by yeast suppression analysis (10, 12).

The requirement for both CifA and CifB in transgenic CI induction but only CifA for rescue has been enumerated as the 2-by-1 model (Fig. 5A) (19). Although this genetic summary is usually interpreted in the context of a host modification (HM) mechanism for CI, it can be accommodated by a toxin-antidote (TA) scheme as well (Fig. 5B). However, recent versions of the 2-by-1 model (for example, reference 20) have posited a primary role for CifA in CI induction in which CifA, by an unknown mechanism, causes modifications in sperm precursors that, following fertilization, will be lethal to the embryo if not reversed, also by CifA, in the egg. CifB, in this view, has only an ancillary role in CI induction, possibly by maintaining proper levels or activity of CifA during spermatogenesis. While the plausibility of this model could be questioned on several
grounds, it does make a straightforward prediction, which is that CifA is strictly necessary for CI induction. Our data demonstrate that the wNo CifA (CinA) protein is dispensable for CI induction and instead show that wNo CifB (CinB) is sufficient; this embryonic lethality is completely blocked following mating with females expressing CinA<sup>wNo</sup> in the germ line (Fig. 3 and 4). These findings align well with results on yeast toxicity caused by CinB<sup>wNo</sup> alone and its suppression by CinA<sup>wNo</sup> (Fig. 1C; Fig. 2B). Very recently, transgenic expression of wPip CidA and CidB proteins in the malaria vector Anopheles gambiae has been reported (29). Expression of CidB<sup>wPip</sup> by itself in male mosquitoes induced very strong transgenic CI, and CidA<sup>wPip</sup> in females was sufficient to block the embryonic lethality caused by crossing to transgenic CidB<sup>wPip</sup> male mosquitoes. These and our results

**FIG 5** Models for cytoplasmic incompatibility (CI) caused by *Wolbachia*. (A) The 2-by-1 host modification (HM) model. Although the 2-by-1 model is strictly a genetic description, the most recently articulated version has been interpreted in the context of the HM mechanistic model shown here (20). CifA is regarded as the key CI-inducing factor that is responsible for modifications of sperm; CifB in this scheme has an undefined accessory role within the testes that is required for CI induction. In the fertilized egg, if CifA is secreted by infecting *Wolbachia* (blue ovals); it works in the opposite fashion to reverse the sperm modifications and rescues embryonic viability. *Wolbachia* is known not to be incorporated into mature sperm and is instead eliminated along with other organelles in the “waste bag” during spermiogenesis. There are multiple additional variants of the HM model that predate identification of the Cif proteins. (B) Toxin-antidote (TA) model. In this model, CifB is the fundamental CI-inducing factor and is postulated to be imported into the mature sperm (along with CifA); CifA may promote CifB levels or packaging into sperm or protect sperm precursors from CifB-induced toxicity. Upon fertilization, or possibly before, CifB enzymatic activity, either from the CidB deubiquitylase or CinB nuclease, alters the sperm-derived chromosomes in some way. CifB is proposed to be relatively long-lived, and CifA, the antidote that binds CifB, short-lived, so additional high-level expression of CifA in the egg is required to counter CifB activity.
together therefore argue that the fundamental inducers of CI are the CifB proteins, and this is true for both CinB nuclease and CidB deubiquitylase Wolbachia CI factors. Three different model organisms, D. melanogaster, A. gambiae, and S. cerevisiae, used for transgenesis studies of the cif genes, have now yielded evidence for a “1-by-1” genetic scheme in which CifB is the lone modification factor or toxin and CifA is the rescue factor or antidote.

Nevertheless, both the A and B factors from other Wolbachia cif operons were previously shown to be necessary for transgenic CI induction in Drosophila (11, 12), and here, we have reproduced our original finding showing that cinBmut is not sufficient for inducing strong transgenic CI but requires cinA compensation (Fig. 3B). The exact reason for the different CI gene requirements using genes from the same CI operon for expression in different insect models (Drosophila versus Anopheles) or from related CI operons of different Wolbachia strains (wPip versus wNo) is not known, but it likely derives from the level and localization of transgenic expression of these proteins in the host (see reference 17 for a fuller discussion). Our coexpression data in yeast suggest that CinA might stabilize the CifB protein, although this remains to be tested (Fig. 1D). The baker’s yeast model has been a useful predictor of the cellular effects of the CI factors, even though yeasts are not known to be natural hosts to Wolbachia.

By themselves, the current data do not distinguish between HM and TA mechanisms. The TA mechanism, but not HM, explicitly requires binding between modifier/toxin and rescue/antidote factors (Fig. 5). Given the results in the current work, this binding should occur between CifB brought in by sperm and CifA expressed in the female germ line. Support for this has recently been garnered using high-resolution crystal structures for several CifA-CifB binary complexes (24). From these structures, it was found that the interfaces between the A and B factors are dominated by charged and polar residues. Amino acids in these interfaces could be mutated to eliminate binding; this blocked the ability of the CifA rescue factors to suppress cognate CifB toxicity in yeast and to rescue transgenic CI in flies. If complementary mutations were made at the interfaces to partially restore CifA-CifB binding, yeast toxicity caused by the mutant CifB was suppressed by coexpression of the CifA interface mutant.

The current work aligns results from different model systems and makes clear which CI factors are fundamentally responsible for the induction and rescue of CI. Ultimately, determining the mechanism of CI will require identifying the relevant molecular targets of the CifB enzymes in the host and the timing of their action and determining how direct binding by the CifA factors blocks or reverses CifB action in vivo (17). Neither in vitro CinB nuclease activity nor CidB deubiquitylase activity is diminished by cognate CifA binding (10, 12). Therefore, CifA proteins may block or reverse enzyme-substrate interactions or alter enzyme localization (or both) in vivo. Whatever the CI mechanisms, they are likely to impact conserved physiological pathways given the wide host range of Wolbachia that cause CI and the even broader range of species in which heterologous expression of the CI factors mimics a toxin-antidote system. Understanding these mechanisms will allow more rational manipulation of CI for applications in crop protection and disease vector control.

**MATERIALS AND METHODS**

**Yeast and bacterial strains and plasmid construction.** MHY1774 (W303-1A) and MHY10139 (BY4741) yeast strains were used for yeast growth assays. E. coli TOP10F’ cells were used for DNA cloning, and Rosetta BL21(DE3 pLysS) cells were employed for protein expression. PCR products were generated with the primers listed in Table S2 in the supplemental material using HF Phusion DNA polymerase (New England Biolabs).

For galactose-inducible expression of proteins in yeast, pRS425GAL1 (LEU2) and pRS416GAL1 (URA3) expression plasmids were used (30). Constructs p425GAL1-Flag-Cina and p416GAL1-Flag-CinB were described earlier, as were the wPip equivalents (10, 18). Mutations were introduced into Flag-CinB in p416GAL1 by QuikChange (Agilent) mutagenesis using primers listed in Table S2.

For protein expression in E. coli and transgene expression in D. melanogaster, coding sequences of Cina, CinB, and Cina, T2A-CinB, optimized for expression in Drosophila, were synthesized by GenScript. To make transgenic flies, the pTiger vector, bearing a Gal4-binding UAS site, was used. Cina, CinB, and Cina, T2A-CinB coding sequences were separately cloned into pTiger using.
KpnI and SpeI restriction sites. For expression of proteins in E. coli, pET28a-pp, which encodes an N-terminal His tag, and pGEX-6P-1, which encodes an N-terminal GST tag, were used. The cinA<sup>reporter</sup> gene was subcloned from pRS425-CinA<sup>reporter</sup> into pET28a-pp using BamHI/Sall sites; cinB<sup>reporter</sup> was subcloned from pRS416-CinB<sup>reporter</sup> into pGEX-6P-1 using BamHI/NotI sites. pGEX-6P-1-CinB<sup>reporter</sup> was reported previously (12), while the coding sequence for CinA<sup>reporter</sup> was subcloned from pUAS-attb-CinA<sup>reporter</sup> (12) into pET28a-pp NdeI/Xhol sites. Also see Table S1.

Yeast growth analysis. MHY1774 or MHY10139 yeast were transformed with the plasmids indicated in the figures. To test growth, cultures were grown at 30°C in synthetic defined (SD) raffinose-containing medium lacking uracil and leucine for 2 days before being spotted in 6-fold serial dilution onto SD plates lacking uracil and leucine and containing either 2% galactose or glucose.

For immunoblot testing of Wolbachia protein expression in yeast, coexpression cultures in SD-raffinose lacking uracil and leucine were diluted to an optical density at 600 nm (OD<sub>600</sub>) of 0.1 in SD-galactose medium lacking uracil and leucine and cultured for 12 to 16 h at 30°C until reaching an OD<sub>600</sub> of 0.8 to 1.0. Cells were harvested and treated for 5 min at room temperature with 0.1 M NaOH (31). Cell pellets were lysed in SDS sample buffer, and clarified lysates were analyzed by immunoblotting (18). Mouse anti-FLAG M2 (Sigma; 1:10,000) or mouse anti-PGK (yeast phosphoglycerate kinase; Abcam; 1:10,000) was used for immunoblotting along with sheep anti-mouse-horseradish peroxidase (HRP) NXX931V secondary antibody (GE Healthcare; 1:5,000 or 1:10,000, respectively). All yeast growth and Western blot data shown are representative of at least two biological replicates.

Protein-binding analysis. To test interactions between various CinA and CinB proteins, GST fusion protein pulldown assays were performed. Two different expression methods were used. In the first, pET28a-pp-CinA (confering kanamycin resistance) and pGEX-6P-1-CinB (confering ampicillin resistance) plasmids were co-transformed into Rosetta E. coli BL21(DE3 pLyS5). Bacteria were grown at 37°C in 75 mL LB (Luria Broth) medium containing kanamycin and ampicillin to an OD<sub>600</sub> of 0.6 to 0.8, induced with 600 μM IPTG (isopropyl-β-thiogalactoside), and incubated for ~20 h at 18°C. Cultures were centrifuged at 4°C and 7,000 × g for 10 min.

Cell pellets were resuspended in GST tag lysis buffer (50 mM Tris-HCl, pH 7.5, 150 mM NaCl, 0.01% Tween 20, 10 mM imidazole, pH 8.0, 2 mM phenylmethylsulfonyl fluoride (PMSF), 20 μg/mL DNase I, and 1 mg/mL chicken egg white lysozyme), kept on a rotator at 4°C for 30 min, and then lysed by sonication. Crude lysates were centrifuged at 21,000 × g for 30 min at 4°C. Clarified lysates were incubated with 50 μL Pierce glutathione-agarose beads (Thermo, USA) prewashed with wash buffer (50 mM Tris-HCl, pH 8.0, and 150 mM NaCl) and rotated at 4°C for 1 h. Beads were washed six times with 1 mL wash buffer, and proteins were eluted by adding two bed volumes of elution buffer (50 mM Tris-HCl, pH 8, 150 mM NaCl, and 20 mM reduced glutathione) and incubated on a rotator at room temperature for 15 min. Inputs and eluted fractions were resolved by SDS-PAGE. The proteins were visualized by GelCode Blue staining (Thermo Fisher, USA) and imaged on a GBox (Syngene).

In the second method, the expression plasmids were transformed individually into competent Rosetta cells. Cultures were grown and pelleted as above. Cells expressing His<sub>6</sub>-CinA were resuspended in His tag lysis buffer (50 mM Tris-HCl, pH 7.5, 150 mM NaCl, 0.01% Tween 20, 10 mM imidazole, pH 8.0, 2 mM PMSF, 20 μg/mL DNase I, and 1 mg/mL lysozyme), while those expressing GST-CinB were resuspended in GST tag lysis buffer. Cells were incubated on a rotator for 30 min at 4°C and then lysed by sonication as above. Clarified GST-CinB-containing lysates were first incubated with glutathione-agarose resin and rotated at 4°C for 1 h. After removing the flowthrough, the beads were washed three times with 1 mL GST tag wash buffer, followed by addition of clarified His<sub>6</sub>-CinA lysates and rotation at 4°C for 1 h. The beads were washed five times with 1 mL wash buffer, with all subsequent steps as in the previous method.

Hatch rate analysis of transgenic D. melanogaster crosses. To generate the wNo Cif transgenic flies used for hatch rate analyses, DNA constructs were sent to BestGene, Inc. for microinjection into D. melanogaster embryos. Flies transgenic for cinA<sup>reporter</sup>, cinB<sup>reporter</sup>, and cinA-T2A-cinB<sup>reporter</sup> were generated previously (12). White Canton-S (wCS; WT; catalog no. 189) and MTD driver (catalog no. 31777; GAL4) fly strains were gifts or were obtained from the Bloomington Stock Center. Fly strain 9744 (containing an attP insertion site on the third chromosome) was used for all genetic constructs for site-directed attP/B integration by the PhiC31 integrase, except the 9723 strain was chosen for site-specific integration of cinA<sup>reporter</sup> on the second chromosome. Flies were reared on standard corn meal-based solid media. All flies for the hatch rate assays were homozygous for the integrated genes except for the cinA-T2A-cinB<sup>reporter</sup> line, which showed a high degree of sterility when virgin red-eyed individuals were mated. This made it unfeasible to maintain a homozygous line; the sterility was likely due to leaky transgene expression, which is known to occur with the UAS system (10). For cinA-T2A-cinB<sup>reporter</sup>, red-eyed progeny of white-eyed males and red-eyed virgin females were used for maintenance of the heterozygous line as well as for hatch rate experiments.

Except for the cinA-T2A-cinB<sup>reporter</sup> flies, parental flies used in the hatch rate experiments were generated by crossing MTD-GAL4 virgin females to the corresponding transgenic males. The crosses were maintained at ~22°C (temperature was temporarily lowered to 18°C for overnight virgin collection) on a standard diet. Virgin flies of the desired genotype and sex were collected, aged at 25°C for 2 to 4 additional days, and used to set up 1-by-1 crosses for hatch rate determination. For each cross, a virgin female and male were mated as described (12). All crosses were incubated at 25°C overnight (~17 h) (initial incubation), and the original apple juice plates were set aside and replaced with freshly yeasted plates, which were kept at 25°C for 24 h (additional incubation). Adult flies were then removed and frozen at ~80°C for future expression analysis. Both sets of plates were incubated at 25°C for another 24 h before the number of hatched and unhatched embryos was counted. Embryo numbers from the two
sets of plates were pooled, and any mating pair with fewer than 15 total embryos laid was discarded. The counting was not blinded. One-way analysis of variance (ANOVA) with multiple-comparison statistical analysis was performed using GraphPad Prism (v.9) software.

**Cytological analysis of embryos.** To prepare embryos for cytological analysis, ~100 males and ~100 females (2- to 4-day-old virgins) were used. The methods were described previously (12) with the exception that embryo collection was repeated 2 to 3 times to collect sufficient 1- to 2-h embryos.

In preparation for microscopic analysis, the melanin was removed, and embryos were treated as described (12). Propidium iodide (PI; Sigma-Aldrich) mounting medium was used for DNA staining, and stained embryos were mounted on glass slides and sealed under a coverslip with nail polish. Imaging was performed with a Zeiss Axioskop microscope with AxioCam MRm camera using 10× and 40× objective lenses.

To assess the cytology of very early stage embryos from incompatible crosses with cinB<sup>fl</sup>, cinA<sup>fl</sup>, T2A-cinB<sup>fl</sup>, and T2A-cinA<sup>fl</sup> transgenic males, a slightly different method was employed. Roughly 100 virgin female wCS WT flies and ~100 transgenic cin<sup>fl</sup> or cinA<sup>fl</sup> males were crossed as described previously (12). Embryos were collected from fresh apple juice plates after 30 min, directly dechorionated in 10 mL 50% fresh bleach for 1 to 3 min, washed once in 10 mL fresh embryo wash buffer (0.6% NaCl, 0.04% Triton X-100) for 5 s, and fixed immediately with methanol or formaldehyde solution. The methanol fixation method (as in the images used for Fig. 4) (12) and the formaldehyde fixation method (32) were described previously. Embryos were washed and stored in methanol. Embryo collection was repeated 3 to 4 times to collect sufficient 30-min embryos. All ensuing steps were as above, except the embryos were stained with 40 μL fresh Hoechst 33342 (Thermo Fisher Scientific) at 1:1,000 in phosphate-buffered saline/bovine serum albumin/Triton X-100/sodium azide (PBTA) buffer (10). Imaging was done as above using 10× and 100× objective lenses.

**Transgenic fly mRNA expression analysis by reverse transcriptase quantitative PCR.** To analyze expression levels of w<sup>cinA</sup>, cinB<sup>fl</sup>, and cinA-T2A-cinB<sup>fl</sup> transgenes in flies used for hatch rate assays (Fig. S3 in the supplemental material), 11 female or male flies were pooled and kept frozen at −80°C until processed. Untransformed flies were used as a negative control. Real-time PCR was performed as described previously (24).

**Statistical analysis.** All statistical analyses were done with GraphPad Prism (v.9) software. For comparisons between more than two data sets, a nonparametric Kruskal-Wallis one-way ANOVA test followed by Dunn’s multiple comparisons was used in hatch rate analyses. Pairwise χ<sup>2</sup> (Fisher’s exact) test was used for the cytological analyses to infer statistically significant differences between normal and defective cytological phenotypes. A parametric t test was used to compare transgene mRNA levels of transgenic flies.

**SUPPLEMENTAL MATERIAL**

Supplemental material is available online only.

**FIG S1**, EPS file, 0.6 MB.

**FIG S2**, EPS file, 0.3 MB.

**FIG S3**, EPS file, 0.2 MB.

**TABLE S1**, EPS file, 0.7 MB.

**TABLE S2**, EPS file, 1.3 MB.

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We declare no conflicts of interest.

**REFERENCES**

1. Duron O, Hurst GDD. 2013. Arthropods and inherited bacteria: from counting the symbionts to understanding how symbionts count. BMC Biol 11:45. https://doi.org/10.1186/1741-7007-11-45.

2. Zug R, Hammerstein P. 2012. Still a way of variance (Wolbachia) analysis of recent data suggests that 40% of terrestrial arthropod species are infected. PLoS One 7:e38544. https://doi.org/10.1371/journal.pone.0038544.

3. Werren JH, Baldo L, Clark ME. 2008. Wolbachia: master manipulators of invertebrate biology. Nat Rev Microbiol 6:741–751. https://doi.org/10.1038/nrmicro1969.

4. Serbus LR, Casper-Lindley C, Landmann F, Sullivan W. 2008. The genetics and cell biology of Wolbachia-host interactions. Annu Rev Genet 42:683–707. https://doi.org/10.1146/annurev.genet.41.110306.130354.

5. Gottoh T, Noda H, Ito S. 2007. Cardinium symbionts cause cytoplasmic incompatibility in spider mites. Hereditas (Edinb) 98:13–20. https://doi.org/10.1083/jhdy.6808881.

6. Bourtis K, Dobson SL, Xi Z, Rasgon JL, Calvitti M, Moreira LA, Bossin HC, Moretti R, Bonan LA, Hughes GL, Mavingui P, Gilles JR. 2014. Harnessing mosquito-Wolbachia symbiosis for vector and disease control. Acta Trop 132:510–63. https://doi.org/10.1016/j.actatropica.2013.11.004.

7. O’Neill SL. 2018. The use of Wolbachia by the World Mosquito Program to interrupt transmission of Aedes aegypti transmitted viruses. Adv Exp Med Biol 1062:355–360. https://doi.org/10.1007/978-1-4939-6727-1_24.

8. Hoffmann AA, Montgomery BL, Popovic J, Iurbe-Omaeaei I, Johnson PH, Muzzi F, Greenfield M, Durkan M, Leong YS, Dong Y, Cook H, Asford J, Callahan AG, Kenny N, Omodei C, McGraw E, Ryan PA, Ritchie SA, Turelli M, O’Neill SL. 2011. Successful establishment of Wolbachia in Aedes populations to suppress dengue transmission. Nature 476:454–457. https://doi.org/10.1038/nature10356.

9. Uhartani A, Indiriani C, Ahmad RA, Tantowijoyo W, Arguni E, Ansari MR, Supriyati E, Wardana DS, Metika Y, Ernesia I, Nurhayati I, Prabowo E, Andari B, Green BR, Hodgson L, Cutscher Z, Rancés E, Ryan PA, O’Neill SL, Dufault SM, Tanamas SK, Jewell NP, Anders KL, Simmons CP, AWARE Study Group. 2021. Efficacy of Wolbachia-infected mosquito deployments for the control of dengue. N Engl J Med 384:2177–2186. https://doi.org/10.1056/NEJMoa2030243.

January/February 2022 Volume 13 Issue 1 e03177-21 mbio.asm.org
21. Poinsot D, Merçot H. 1997. A Wolbachia deubiquitylating enzyme induces cytoplasmic incompatibility. Nat Microbiol 2: 17007. https://doi.org/10.1038/nmicrobiol.2017.7.

20. Shropshire JD, Kalra M, Bordenstein SR. 2020. Evolution-guided mutagenesis of the cytoplasmic incompatibility proteins: identifying CifA’s complex functional repertoire and new essential regions in CifB. PLoS Pathog 16:e1008794. https://doi.org/10.1371/journal.ppat.1008794.

19. Shropshire JD, Bordenstein SR. 2019. Two-by-one model of cytoplasmic incompatibility. Nature 543:243–247. https://doi.org/10.1038/nature21391.

18. Beckmann JF, Sharma GD, Mendez L, Chen H, Hochstrasser M. 2019. The Wolbachia cytoplasmic incompatibility enzyme CidB targets nuclear import and protamine-histone exchange factors. Elife 8:e50026. https://doi.org/10.7554/eLife.50026.

17. Chen H, Zhang M, Hochstrasser M. 2020. The biochemistry of cytoplasmic incompatibility caused by endosymbiotic bacteria. Genes 11:857. https://doi.org/10.1038/s41564-020-00998-6.

16. Xiao Y, Chen H, Wang H, Zhang M, Chen X, Berk JM, Zhang L, Wei Y, Li W, Cui W, Wang F, Wang Q, Cui C, Li T, Chen C, Ye S, Zhang L, Ji X, Huang J, Wang W, Wang Z, Hochstrasser M, Yang H. 2021. Structural and mechanistic insights into the complexes formed by Wolbachia cytoplasmic incompatibility factors. Proc Natl Acad Sci U S A 118:e2107699118. https://doi.org/10.1073/pnas.2107699118.

15. Bing XL, Zhao DS, Sun JT, Zhang KJ, Hong XY. 2020. Genomic analysis of Wolbachia in the mosquito vector. Nat Microbiol 5:17007. https://doi.org/10.1038/s41564-020-00998-6.

14. Lindsey ARI, Rice DW, Bordenstein SR, Brooks AW, Bordenstein SR, Weill M, Sicard M, Charlat S. 2019. The toxin-antidote model of cytoplasmic incompatibility: synthetic recapitulation by transgenic expression of Wolbachia. Genome Biol Evol 13:evab111. https://doi.org/10.1093/gbe/evab111.

13. Gillespie JJ, Driscoll TP, Verhoeve VI, Rahman MS, Macaluso KR, Azad AF. 2021. Wolbachia CinB Induces CI does the female host bear a physiological cost? Evolution 51:180–186. https://doi.org/10.1534/genetics.111.136291.

12. Chen H, Ronau JA, Beckmann JF, Hochstrasser M. 2019. A Wolbachia nu- clease and its binding partner provide a distinct mechanism for cytoplasmic incompatibility. Proc Natl Acad Sci U S A 116:22314–22321. https://doi.org/10.1073/pnas.1914571116.

11. LePage DF, Metcalf JA, Bordenstein SR, On J, Perlmutter JI, Shropshire JD, Layton EM, Funkhouser-Jones LJ, Beckmann JF, Bordenstein SR. 2017. Prophage WO genes recapitulate and enhance Wolbachia-induced cytoplasmic incompatibility. Nature 543:243–247. https://doi.org/10.1038/nature21391.

10. Beckmann JF, Ronau JA, Hochstrasser M. 2017. A Wolbachia deubiquitylating enzyme induces cytoplasmic incompatibility. Nat Microbiol 2: 17007. https://doi.org/10.1038/nmicrobiol.2017.7.

9. Adams KL, Abernathy DG, Willett BC, Selland EK, Itoe MA, Catteruccia F. 2021. Wolbachia cifB induces cytoplasmic incompatibility in the malaria mosquito vector. Nat Microbiol 6:1575–1582. https://doi.org/10.1038/s41564-021-00998-6.

8. Mumberg D, Müller R, Funk M. 1994. Regulatable promoters of Saccharomyces cerevisiae: comparison of transcriptional activity and their use for heterologous expression. Nucleic Acids Res 22:5767–5768. https://doi.org/10.1093/nar/22.25.5767.

7. Kushnirov VV. 2000. Rapid and reliable protein extraction from yeast. Yeast 16:857–860. https://doi.org/10.1002/1097-0061(20000630)16:9<857::AID-YEA561>3.0.CO;2-B.

6. Rothwell WF, Sullivan W. 2000. Fluorescent analysis of Drosophila embryos, p 149. In Sullivan W, Ashburner M, Hawley RS (ed), Drosophila, 2nd ed. Drosophila protocols. Cold Spring Harbor Laboratory Press, Cold Spring Harbor, NY.

5. Sisakova E, Stanley LK, Weiserova M, Szczelkun MD. 2008. A RecB-family nuclease motif in the Type I restriction endonuclease EcoR124I. Nucleic Acids Res 36:3939–3949. https://doi.org/10.1093/nar/gkn333.