Evolutionary Genetics of Cytoplasmic Incompatibility Genes cifA and cifB in Prophage WO of Wolbachia

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Abstract
The bacterial endosymbiont Wolbachia manipulates arthropod reproduction to facilitate its maternal spread through host populations. The most common manipulation is cytoplasmic incompatibility (CI): Wolbachia-infected males produce modified sperm that cause embryonic mortality, unless rescued by embryos harboring the same Wolbachia. The genes underlying CI, cifA and cifB, were recently identified in the eukaryotic association module of Wolbachia’s prophage WO. Here, we use transcriptomic and genomic approaches to address three important evolutionary facets of the cif genes. First, we assess whether or not cifA and cifB comprise a classic toxin–antitoxin operon in wMel and show that the two genes exhibit striking, transcriptional differences across host development. They can produce a bicistronic message despite a predicted hairpin termination element in their intergenic region. Second, cifA and cifB strongly coevolve across the diversity of phage WO. Third, we provide new domain and functional predictions across homologs within Wolbachia, and show that amino acid sequences vary substantially across the genus. Finally, we investigate conservation of cifA and cifB and find frequent degradation and loss of the genes in strains that no longer induce CI. Taken together, we demonstrate that cifA and cifB exhibit complex transcriptional regulation in wMel, provide functional annotations that broaden the potential mechanisms of CI induction, and report recurrent erosion of cifA and cifB in non-CI strains, thus expanding our understanding of the most widespread form of reproductive parasitism.

Key words: symbiosis, reproductive manipulation, gene loss, bacteriophage, prophage.

Introduction

The genus Wolbachia is the most widespread group of maternally transmitted endosymbiotic bacteria (Zug et al. 2012). They occur worldwide in numerous arthropods and nematodes and can selfishly manipulate reproduction (Werren et al. 2008), confer antiviral defense (Teixeira et al. 2008; Bian et al. 2010), and assist reproduction and development of their hosts (Hoerauf et al. 1999; Dedeine et al. 2001; Hosokawa et al. 2010). The most common parasitic manipulation is cytoplasmic incompatibility (CI), whereby Wolbachia-infected males produce modified sperm that can only be rescued by eggs infected with the same Wolbachia strain (Yen and Barr 1971). If the modified sperm fertilize eggs infected with no Wolbachia (unidirectional CI) or a genetically incompatible Wolbachia strain (bidirectional CI), then delayed histone decondensation, improper chromosome condensation, and cell division abnormalities result in embryonic arrest and death (Lassy and Karr 1996; Tram and Sullivan 2002; Serbus et al. 2008; Landmann et al. 2009). Other described reproductive manipulations include parthenogenesis (Stouthamer et al. 1990),...
male-killing (Hurst et al. 1999), and feminization (Rousset et al. 1992), all of which enhance the fitness of Wolbachia-infected females and assist the spread of the infected matriline through a population. These manipulations, once sustained, can also impact host evolution including speciation (Bordenstein et al. 2001; Jaenike et al. 2006; Brucker and Bordenstein 2013) and mating behaviors (Randerson et al. 2000; Moreau et al. 2001; Miller et al. 2010; Shropshire and Bordenstein 2016).

In addition to the aforementioned reproductive manipulations, Wolbachia strains affect host biology by provisioning nutrients (Hosokawa et al. 2010), altering host survivorship (Min and Benzer 1997) and fecundity (Stouthamer and Luck 1993; Dedeine et al. 2001), and importantly, protecting the host against pathogens (Teixeira et al. 2008; Kambris et al. 2009; Moreira et al. 2009; Bian et al. 2010; Hughes et al. 2011; Walker et al. 2011). The combination of reproductive manipulations that enable Wolbachia to spread in a population and the ability to reduce vector competence through pathogen protection have placed Wolbachia in the forefront of efforts to control disease carrying arthropod populations (Werren and Windsor 2000; Hilgenboecker et al. 2008; Zug 2011). Despite these applications, the mechanisms through which Wolbachia are transmitted across arthropod taxa are largely unknown, though they can be at least partially attributed to the widespread prevalence of Wolbachia across arthropod taxa (Werren and Windsor 2000; Hilgenboecker et al. 2008; Zug et al. 2012), and decades of research, only recently have the genes underlying CI been determined (Beckmann et al. 2017; LePage et al. 2017).

Two studies converged on the same central finding: Coexpression of a pair of syntenic genes recapitulates the CI phenotype (Beckmann et al. 2017; LePage et al. 2017). Uninfected Drosophila melanogaster males transgenically expressing the two genes from wMel Wolbachia caused CI-like embryonic lethality when crossed with uninfected females that was notably rescued by wMel-infected females (LePage et al. 2017). Additionally, the two wMel genes separately enhanced wMel-induced CI in a dose-dependent manner when expressed in infected males, and the CI was again rescued by wMel-infected females (LePage et al. 2017). In the other study, CI-like embryonic lethality was also recapitulated in D. melanogaster males through transgenic coexpression of homologous transgenes cidA and cidB, encoded by the wPip strain of Wolbachia that naturally infect Culex mosquitoes (Beckmann et al. 2017). These two genes are located in the recently discovered eukaryotic association module of temperate phage WO (Bordenstein and Wernegreen 2004; Chaefee et al. 2010) while Wolbachia are mainly vertically transmitted from mothers to offspring. The genes were proposed as candidate CI effectors due to the presence of one of the protein products in the spermatheca of infected female mosquitoes (Beckmann and Fallon 2013) and their absence in the wAu Wolbachia strain that lost CI function (Sutton et al. 2014).

The wMel homologs of these genes are designated CI factors cidA (locus WD0631) and cidB (locus WD0632), with cidA always encoded directly upstream of cidB (LePage et al. 2017). The gene set occurs in varying copy number across 11 total CI-inducing strains, and the copy number tentatively correlates with CI levels. Core sequence changes of the two genes exhibit a pattern of codivergence and in turn closely match bidirectional incompatibility patterns between Wolbachia strains. Homologs of CifA and CifB protein sequences belong to four distinct phylogenetic Types (designated Types I–IV) that do not correlate with various phylogenies of Wolbachia housekeeping genes or gpW (locus WD0640) in phage WO (LePage et al. 2017). The homologous sequences in wPip also cluster in Type I, though they are 66% and 76% different from wMel’s, respectively (Beckmann et al. 2017). Hereinafter we use cidA and cidB to refer to these genes, unless specifically referring to analyses of the wPip homologs, cidA and cidB. In vitro functional analyses revealed that cidB encodes deubiquitylase activity, and cidA encodes a protein that binds CidB (Beckmann et al. 2017). Mutating the predicted catalytic residue in the deubiquitylating domain of CidB results in a loss of the CI-like function in transgenic flies (Beckmann et al. 2017). Whether these genes or other alleles have additional enzymatic or regulatory roles and which other residues are important for function remain open questions.

There are important considerations for the location, organization, and characterization of these genes. Whether or not cidA and cidB form a strict, toxin–antitoxin operon is debatable, and likewise has important implications for how gene expression is regulated by Wolbachia during host infection. Support for the operon hypothesis is based on weak transcription across the junction between cidA and cidB, inferred to be due to the presence of bicistronic mRNA (Beckmann and Fallon 2013; Beckmann et al. 2017); an alternative explanation is transcriptional slippage. Quantitative transcription analyses and various computational predictions of operon structure do not support the operon hypothesis (LePage et al. 2017). Moreover and importantly, transgenic studies show that both cidA and cidB are required for induction of CI and thus cannot form a strict toxin (cidB)–antitoxin (cidA) system as both genes positively contribute to CI and can individually enhance Wolbachia-induced CI (LePage et al. 2017). However, like toxin–antitoxin systems, CidA binds CidB in vitro and expression of cidA rescues temperature-sensitive growth inhibition induced by cidB expression in Saccharomyces, via an as-yet-unknown mechanism (Beckmann et al. 2017).
As it stands now, the genes remain largely unannotated with the exception of a few small domains. If other predicted protein domains occur in CifA and CifB, they could allow for new hypotheses for the mechanism of Cif. Finally, the sequence diversity and/or loss of cif genes across the Wolbachia tree may give insights into the selective conditions that maintain the cif genes versus those that do not. Exploration of cif gene regulation, expression, and function thus can provide a framework for more targeted investigations of Wolbachia–host interactions, and potentially inform the deployment of Wolbachia-based arthropod control.

Materials and Methods

Expression

For analysis of RNAseq data, we used our published approach (Gutzwiller et al. 2015). Briefly, fastq sequences for 1-day-old male and female flies were mapped against the Wolbachia wMel reference genome (GenBank AE017196) using bwa mem v. 0.7.5a with default parameters in paired-end mode. Mapped reads were sorted and converted to BAM format using samtools v0.1.19 after which BAM files were used as input to Bedtools (bedcov) to generate pileups and count coverage at each position. For expression correlations between genes, the raw RNAseq counts were divided by (gene length–average at each position. For expression correlations between all genes was defined as (1 + r2)0.5, where 99 corresponds to read length (100) – 1. Within a growth stage these values were multiplied by 1e6/(sum of

Hairpin transcription termination elements (Gautheret and rna.igmors.u-psud.fr/toolbox/arnold/) was used to predict et al. 2013 ), with default parameters, in conjunction with

Wolbachia

99), where 99 corresponds to read length (100)

and 1492R (GGTTACCTTGTTACGACTT) with the same rRNA gene primers 27F (AGAGTTTGATCCTGGCTCAG) and

Exploration of

cif

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Operon Prediction In Silico

We used the dynamic profile of the transcriptome above to identify operons within the wMel genome using two different approaches. We used the program Rockhopper (McClure et al. 2013), with default parameters, in conjunction with the BAM files generated above to delineate likely operons across the entire genome. The Arnold web server (http://rna.igmors.u-psud.fr/toolbox/arnold/) was used to predict hairpin transcription termination elements (Gautheret and Lambert 2001; Macke et al. 2001).

Nucleic Acid Extractions and Quantitative Reverse Transcription Polymerase Chain Reaction

To identify Wolbachia gene expression in adult male and female D. melanogaster, RNA was extracted from individual, age-matched flies (1–3 days old, stock 145) using a modified Trizol extraction protocol. Briefly, 500 µl of Trizol was added to individual flies and samples homogenized using a pestle. After a 5-min incubation at room temperature, a 12,000 rcf centrifugation (at 4 °C for 10 min) was followed by a chloroform extraction. Aqueous phase containing RNA was extracted a second time with phenol: Chloroform before isopropanol precipitation of RNA. This RNA pellet was washed and resuspended in THE RNA Storage Solution (Ambion). RNA used in subsequent analyses was subjected to a short DNase treatment (10 min at 37 °C then 10 min at 75 °C to inactivate the enzyme). To detect the number of cifA and cifB transcripts as well as RNA levels across the junction between cifA and cifB, we utilized the RNA extracted from these flies and the SensiFAST SYBER Hi-ROX One-step RT mix (Bioline) and the Applied Biosystems StepOne Real-time polymerase chain reaction (PCR) system. Quantitative reverse transcription polymerase chain reaction (qRT-PCR) was performed with the following primer sets: cifA: ATAAAGGCGTTCAGCAGGA, cifB: TCAATGAGGCGCTTCTAGGT; cifA: TACGGGAAG TTATCATGACCA, cifB: TTGCCAGCCTATTCCATAA, cifA: TCTGGTTCATTCAAGAAAGAAAGAATC, cifB: begR: AACATCAAGATCTCCATCCA. As a reference for transcription activity of the core Wolbachia genome, we utilized the Wolbachia ftsZ gene (forward: TTTGTGTGCGCAATACCG; reverse: AGCAAGGCGTTCATTCC). We designed primers for ftsZ because as a core protein involved in cell division, the quantities of ftsZ would better correlate with bacterial numbers and activity. Reactions were performed in duplicate or triplicate in a 96-well plate and CT values generated by the machine were used to calculate the relative amounts of Wolbachia using the ΔΔCt (Livak) method.

To identify a bicistronic message encompassing cifA and cifB, we designed primers based on the 5′-region of cifA and the 3′-region of cifB (WD0631F: ATAAAGGCGTTCAGCAGGA; WD0632R: TTGCCAGCCTATTCCATAA). We extracted RNA from whole animals and performed a DNase treatment (as described above) before using the iScript first strand synthesis kit (Biorad) to generate cDNA. Negative controls included RT minus reactions. Resulting cDNA and negative controls were used in PCR reactions with the primers above and the following cycling conditions: 95 °C 5 min then 35 cycles of 95 °C for 1 min, 64 °C for 1 min, 72 °C for 2.5 min followed by a final extension of 72 °C for 10 min and using the HF Phusion enzyme mix (NEB). As a positive control, to confirm that we could amplify long mRNAs from these samples, we used the 16S rRNA gene primers 27F (AGATTTTGATCCTTGCTCAG) and 1492R (GGTACCTTGTTACGACTT) with the same reverse transcription conditions as above except that the annealing temperature was 55 °C.

Correlated Cif Trees and Distance Matrices

Quantifying congruence scores between the CifA and CifB trees was carried out with Matching Cluster (MC) and
Robinson–Foulds (RF) metrics using a custom python script previously described (Brooks et al. 2016) and the TreeCmp program (Bogdanowicz et al. 2012). MC weights topological congruency of trees, similar to the widely used RF metric. However, MC takes into account sections of subtree congruence and therefore is a more refined evaluation of small topological changes that affect incongruence. Significance in the MC and RF analyses was determined by the probability of 100,000 randomized bifurcating dendrogram topologies yielding equivalent or more congruent trees than the actual tree. Normalized scores were calculated as the MC and RF congruency score of the two topologies divided by the maximum congruency score obtained from random topologies. The number of trees that had an equivalent or better score than the actual tree was used to calculate the significance of observing that topology. Mantel tests were also performed on the CifA and CifB patristic distance matrices calculated in Geneious v8.1.9 (Kearse et al. 2012). A custom Jupyter notebook (Pérez and Granger 2007) running python v3.5.2 (http://python.org) was written in the QIME2 (Caporaso et al. 2010) anaconda environment. The Mantel test (Mantel 1967) utilized the scikit-bio v0.5.1 (scikit-bio.org) Mantel function run, using scikit-bio-distance matrix objects for each gene. The Mantel test was run with 100,000 permutations to calculate significance of the Pearson correlation coefficient between the two matrices using a two-sided correlation hypothesis.

Genomes Used in Comparative Analyses

In order to identify cif homologs across the Wolbachia genomes, we defined orthologs across existing, sequenced genomes using reciprocal best BlastP. We included Wolbachia genomes across five supergroups: Monophyletic clades of Wolbachia based on housekeeping genes, denoted by uppercase letters (O’Neill et al. 1992; Werren et al. 1995). Supergroups A and B are the major arthropod infecting lineages, whereas C and D infect nematodes (Bandi et al. 1998). Supergroup F Wolbachia infect a variety of hosts (Lo et al. 2002). Included in this analysis were nine type A strains (wRi, wSuzi, wHa, wMel, wMelPop, wAu, wRec, wUnii, and wVitA), seven type B strains (wPijJHB, wPipPel, wBo11-b, wNo, wTpre, wAlbB, and wDi), two type C strains (wOv and wOo), and one each type D (wBm) and type F (wCle). We included all genomic data available for each strain such that if multiple assemblies existed for each Wolbachia variant (such as in the case of wUnii) we included the union of all available contigs for that strain. Wolbachia orthologs were defined based on reciprocal best blast hits between amino acid sequences in Wolbachia genomes. An orthologous group of genes was defined by complete linkage such that all members of the group had to be the reciprocal best hit of all other members of the group. Information on strain phenotypes, hosts, and accession numbers can be found in table 1.

Cif Phylogenetics

CifA and CifB protein sequences were identified using BlastP searches of WOMelB WD0631 (NCBI accession number AAS14330.1) and WD0632 (AAS14331.1), respectively. Homologs were selected based on: 1) $E = \leq 10^{-30}$, 2) query coverage greater than 70%, and 3) presence in fully sequenced Wolbachia genomes. All sequences were intact with the exception of a partial WOSuziC CifA (WP_044471252.1) protein. The missing N-terminus was translated from the end of contig accession number CAOU020000024.1 and concatenated with partial protein WP_044471252.1 for analyses, resulting in 100% amino acid identity to WORic CifA (WP_012673228.1). In addition, two previously identified sequences (LePage et al. 2017), WORecB CifB and WORiB CifB, were not available in NCBI’s database and translated from nucleotide accession numbers JQAM01000018.1 and CP001391.1, respectively. The previously identified WOSol homologs (CifA: AGK87106 and CifB: AGK87078) (LePage et al. 2017) were also included in our analyses. All protein sequences were aligned with the MUSCLE (Edgar 2004) plugin in Geneious Pro version 8.1.7 (Kearse et al. 2012); the best models of evolution, according to corrected Akaike (Huvenich and Tsai 1993) information criteria, were estimated to be JTT-G using the ProtTest server (Abascal et al. 2005); and phylogenetic trees were built using the MrBayes (Ronquist et al. 2012) plugin in Geneious.

Protein Structure

All candidate CI gene protein sequences were individually assessed for the presence of domain structure using HHpred (https://toolkit.tuebingen.mpg.de/hhpred/; Söding et al. 2005)) with default parameters and the following databases: SCOPe70 (v.2.06), Pfam (v.31.0), SMART (v6.0), and COG/KOG (v1.0). Schematics were created in inkscape (https://inkscape.org/), to show regions with significant structural hits, as determined by probabilities greater than 50%, or greater than 20% and in the top five hits.

Protein Conservation

Protein conservation was determined with the Protein Residue Conservation Prediction tool (http://compbio.cs.princeton.edu/conservation/index.html; Capra and Singh 2007), using aligned amino acid sequences, Shannon entropy scores, a window size of zero, and sequence weighting set to “false.” Conservation was subsequently plotted in R version 3.3.2, and module regions were delineated according to the coordinates of the WOMelB modules within the alignment. CI gene conservation scores were calculated separately for Type I sequences, and for all Types together. For CifB Type I sequences, the WOVitA4 ortholog was left out, due to the extended C-terminus of that protein. Conservation scores were also calculated for “control proteins”: Wsp (Wolbachia surface protein),
known to be affected by frequent recombination events (Baldo et al. 2005), and FtsZ, which is relatively unaffected by recombination (Baldo, Dunning Hotopp, et al. 2006; Ros et al. 2009). Variation in amino acid conservation between modules and nonmodule regions was assessed in R version 3.3.2 with a one-way ANOVA including “region” (either the unique module number, or “nonmodule”) as a fixed effect, and followed by Tukey Honest Significant Difference for post hoc testing.

Cif Modules

The WOMeB structural regions delineated by HHpred were used to search for the presence of Cifs or remnants of Cifs across the Wolbachia phylogeny. Amino acid sequences of the WOMeB modules were queried against complete genome sequences (table 1) using TBlastN. Any hit that was at least 40% of the length and 40% identity, or at least 90% of the length and 30% identity of the WOMeB module was considered a positive match. Module presence was plotted across a Wolbachia phylogeny constructed using the five multilocus sequence typing (MLST) genes defined by Baldo, Dunning Hotopp, et al. (2006). Nucleotide sequences were aligned with MAFFT version 7.271 (Katoh and Standley 2013), and concatenated prior to phylogenetic reconstruction with RAxML version 8.2.8 (Stamatakis 2014), the GTRGAMMA substitution model, and 1,000 bootstrap replicates. We also searched for cif-like regions in Cardinium: An unrelated endosymbiont that can also cause CI in arthropods (Penz et al. 2012). Here, searches were performed with TBlastN and restricted to all available Cardinium sequence in NCBI GenBank (taxid: 273135).

Hidden Markov Model Searches

To identify cif homologs in draft Wolbachia genome assemblies we used the program suite HMMER (Eddy 2011). We defined cif Types based on our phylogenetic trees (fig. 4) and used aligned amino acids from these Types as input to HMMBUILD, using default parameters. We then searched six Wolbachia WGS assemblies (NCBI project numbers PRJNA310358, PRJNA279175, PRJNA322628) using HMMSEARCH with –F3 1e-20 –cut_nc and –domE 1e-10. Regardless of thresholds used, or cif type of HMM, resulting hits did not differ.

Results

cifA and cifB Are Cotranscribed but Differentially Regulated in wMel

In bacteria, genes are commonly grouped into a single transcriptional unit under the control of one promoter, referred to as an operon. Because cifA and cifB are syntenic across prophase WO of Wolbachia and both involved in CI, we aimed to assess whether cifA and cifB are cotranscribed. We performed RT-PCR using primers that amplify the entire region from the

Table 1

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<th>Supergroup</th>
<th>Strain</th>
<th>Host</th>
<th>Reproductive Phenotypes</th>
<th>Accession Number</th>
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Note.—Reproductive phenotypes include: CI, parthenogenesis-inducing (PI), male-killing (MK), obligate mutualism (OM), no phenotype discovered after assessment (None), and phenotype was not assayed (Undetermined).
start of cifA to the end of cifB (~2.5 kb in total). cDNA amplification of the region from wMel-infected male and female flies was successful (fig. 1A), and the transcript was confirmed to be cifA–cifB using Sanger sequencing from the forward and reverse ends of the cDNA amplicon (supplementary file S2, Supplementary Material online). We could not amplify a larger transcript from the loci flanking cifA and cifB, suggesting that the cifA–cifB transcript is a discrete unit.

Operons are often comprised of loci encoding related processes that can therefore be coregulated conveniently through the control of transcription from one promoter. To assess whether cifA and cifB are coregulated, we reasoned that strictly coregulated loci will have correlated gene expression across host development and similar total expression levels in whole animals. We therefore utilized an existing RNAseq data set for Wolbachia in Drosophila melanogaster, covering 24 life cycle stages and 3 time samplings each for adult males

FIG. 1—Expression of cifA and cifB in adult flies. (A) Amplification of cifA–cifB bicistronic message from cDNA generated from adult flies. Positive amplification occurred in both male and female adult flies. RT minus controls included. (B) RNAseq expression from 1-day-old female and male Drosophila melanogaster flies. Raw reads were mapped to the wMel assembly (using bwa) and coverage visualized using the Integrated Genomics Viewer (v2.3.77). The start of the cifB open reading frame is denoted by a vertical, dotted line.
and females (Gutzwiller et al. 2015). We mapped reads to the existing wMel assembly (see Materials and Methods) and calculated Pearson correlation coefficients of normalized expression values across host development for between all gene pairs. cifA is expressed at much higher absolute levels than cifB (fig. 1B; 8-fold higher based on RPKM values across both genes), and cifA and cifB expression is weakly, negatively correlated (Pearson r = 0.40; P-value: 0.014), suggesting that the expression level of one could have a negative influence on the level of the other. To confirm differential expression of cif genes in wMel, we performed a quantitative RT-PCR analysis of gene expression from 3-day-old male and female flies (fig. 2). We observed transcripts covering the junction between cifA and cifB. However, transcripts covering this junction were more similar to the expression levels in cifA, whereas expression of cifB was 9-fold less, supporting results from the RNAseq analysis.

As a possible explanation for the large absolute differences in transcription, we examined the intergenic sequence between cifA and cifB and identified a Rho-independent transcription terminator at nucleotides 618649–618668. This terminator region is predicted to form a GC-rich hairpin (50% GC compared with the Wolbachia wMel genome wide 35%) in newly synthesized mRNA message proximal to the RNA polymerase. There are two explanations for how the terminator might explain the transcript abundance differences between cifA and cifB, and both have an impact on the operon hypothesis. First, cifA and cifB have their own promoters, but occasionally the genes are cotranscribed as a bicistronic message due to an imperfect hairpin terminator at the end of cifA. In this model, cifA and cifB do not form an operon. Alternatively, the cifA and cifB operon has a single promoter upstream of cifA, and the imperfect terminator provides a mechanism to control transcriptional differences between cifA and cifB in the operon. Functionally resolving whether cifA and cifB have the same or different promoters will be the ultimate arbiter of the two models.

In order to identify loci with similar expression patterns during host development, we clustered all wMel genes based on their similarity in expression across Drosophila development (supplementary fig. S1, Supplementary Material online). cifA did not group with cifB in wMel (fig. 3), suggesting that these two genes are not similarly expressed. Indeed, the pattern of cifA expression differs strikingly from that of cifB. For example, cifA is relatively highly expressed during late embryogenesis and in adults, whereas cifB is relatively highly expressed during the first two-thirds of embryogenesis, and during larval stages (fig. 3A). Curiously, the expression profile of cifA in flies during development is most closely correlated with the wsp locus WD1063 (fig. 3B).

Because of the dramatic absolute difference in cifA and cifB transcript levels, computational methods for operon prediction do not support their cotranscription. For example, after mapping reads to the wMel assembly, we used the resulting BAM files as input to Rockhopper (McClure et al. 2013). The program was able to correctly identify known operons in wMel (such as the T4SS WD0004–WD0008 and the ribosomal protein operon), but it did not identify cifA and cifB as an operon.

In summary, although a bicistronic message of cifA and cifB was detected by qPCR, their absolute and relative expression levels are drastically different. A termination signal in their intergenic sequence may limit expression of the bicistronic message and could explain the much higher absolute level of cifA. Given the negative correlation across growth stages, some entity that activates cifA transcription, or a cifA product itself, could repress cifB transcription. Clearly, cifA and cifB are not a traditional operon and functionally resolving their promoter(s) will reveal much about the regulation of the reproductive manipulations induced by Wolbachia.

New Protein Domain Predictions Are Variable across the Cif Phylogeny

We recovered the four previously identified phylogenetic Types (LePage et al. 2017). Here, our analyses include additional strains that cause reproductive parasitism beyond CI (parthenogenesis and male-killing, table 1), and the more divergent Type IV paralogs for cifA, so far identified in B-Supergroup Wolbachia. We recover a set of Type III alleles from wUn, a strain that induces parthenogenesis in the parasitoid wasp, Muscidifurax uniraptor (Stouthamer et al. 1993). The wBo1–b strain, a male-killer that has retained CI
capabilities (Hornett et al. 2008), has alleles belonging to both Type I and Type IV.

Homologs and predicted protein domains of CifA and CifB for all four phylogenetic Types (LePage et al. 2017) from Wolbachia strains that cause CI, parthenogenesis, male-killing, or no reproductive phenotype were characterized by HHpred homology and domain structure prediction software (Söding et al. 2005). Search parameters are described in the
methods. Several new prominent protein domains, herein referred to as “modules,” were identified for each CifA and CifB protein sequence (table 2).

For CifA, three modules were annotated (fig. 4A, table 2). First, the most N-terminal module (ModA-1) is only recovered in Type I variants, with distant homology (~22% amino acid identity) to Catalase-rel that is predicted to catalyze the breakdown of hydrogen peroxide (Chelikani et al. 2004). The probability of the module being homologous to Catalase-rel is low (prob = 21–24), but the consistent recovery of structure in this region across Type I alleles is notable. The second CifA module in the central region (ModA-2) has homology to a domain of unknown function (Types I, II, and IV, prob = 27–64), globin-like domains (Types II and IV, prob = 21–30), and Puf family RNA-binding domains (Types III and IV, prob = 25–49). The last CifA module in the C-terminal region (ModA-3) has hits to an STE-like transcription factor in all Types (prob = 27–42). In general, the CifA proteins showed distant homology to known domains, but we consistently recovered the same regions of structure within CifA protein Types.

For CifB, three modules were also defined (fig. 4B, table 2). The first (ModB-1) and second (ModB-2) most N-terminal regions of all Types both have matches to the PDDEXX nuclease family (prob = 57–98) and various other restriction endonucleases such as NucS, HSDR_N, and MmcB (prob = 50–91). The third module, found only in the Type I C-terminus (ModB-3), has very strong homology to a number of ubiquitin-modification and protease-like domains (prob = 71–96). This was expected, as ModB-3 contains the predicted catalytic residue associated with CI function in CidB, known to have deubiquitylase activity (Beckmann et al. 2017). WOVitA4 (Type I) has an extended C-terminus not present in any other alleles, and within that extended C-terminus is an additional structural domain, with homology to a Herpesvirus tegument protein (prob = 53), and a phosphohydrolyase-associated domain (prob = 57). CifB Type IV alleles (WOAlbB, WOPip2, and wBol1-b) were not included in the phylogenetic reconstruction, as they are highly divergent and not reciprocal blasts of WOMelB cifB. Despite their divergence, these Type IV CifB alleles have similar structures to Type II and III alleles: Two PDDEXX-like modules, and no Ulp-1-like module 3 (supplementary fig. S3, Supplementary Material online). Full structural schematics with exact coordinates and homology regions for each allele are available in the Supplementary Material online (supplementary figs. S2 and S3, Supplementary Material online), as are all significant domain hits with associated probabilities and extended descriptions (supplementary tables S1 and S2, Supplementary Material online).

CifA and CifB Codiverge

Initial phylogenetic trees based on core amino acid sequences of Type I–III variants of CifA and CifB exhibited similar trees (LePage et al. 2017). Here, we statistically ground the inference of codivergence using the largest set of Wolbachia homologs to date. We quantified congruence between the CifA and CifB phylogenetic trees for Types I–III (supplementary file S1, Supplementary Material online) using MC and RF tree metrics (Robinson and Foulds 1981; Bogdanowicz et al. 2012; Bogdanowicz and Giaro 2013), with normalized distances ranging from 0.0 (complete congruence) to 1.0 (complete incongruence). Results show strong levels of congruence between CifA and CifB (P < 0.00001 for both, normalized MC = 0.06 and normalized RF = 0.125). To further statistically validate the inference of codivergence, we measured the correlation between patristic distance matrices for CifA and CifB using the Mantel test (Mantel 1967). Results demonstrate a high degree of correlation between patristic distance matrices, and through permutation show that independent evolution of CifA and CifB is highly unlikely (Pearson correlation coefficient = 0.905, P = 0.00001).

| Table 2 Predicted Structural Modules of Cif Proteins |
|-----------------|--------|-----------------|-----------------|
| Protein | Module | Size Range (AA) | Homology |
| CifA | ModA-1 | 21–22 | Catalase-rel, decomposes hydrogen peroxide into water and oxygen |
| ModA-2 | 65–264 | DUF3243 domain of unknown function |
| ModA-3 | 47–74 | PDDEXX, PD-(D/E)XX nuclease superfamily |
| ModB-1 | 103–133 | Endonuclease NucS |
| ModB-2 | 122–205 | Restriction endonuclease-like family |
| ModB-3 | 95–147 | HSDR_N, type I restriction enzyme R protein N terminus |

*Colors next to modules are used throughout the Figures. *
*Only present in Type I.*
**Fig. 4**—Phylogenetic relationships and representative predicted protein structure of Cif protein Types. (A) CifA and (B) CifB. Alleles are in bold next to their corresponding accession number, and pink shapes around branches designate monophyletic “Types.” Representative structures are shown for each type, with the length of the protein indicated at the C-terminus. If genes differed by only a few amino acids a single representative is shown. Allele names use the previously described naming convention with a WO prefix referring to particular phage WO haplotype, and the w prefix indicating a phage WO-like island (LePage et al. 2017). The N-terminus of CifA WOSuziC (* in figure) was translated from the end of another contig and concatenated to get the full-length protein (see Materials and Methods). WOMelB and WOMelPop are identical at the amino acid level, as are WOPipJHB and WOPip2.
Cif Proteins Evolve Rapidly

Amino acid sequence conservation across the full length of the Cif proteins was determined and compared with Wolbachia amino acid sequences of genes that either have signatures of recombination and directional section (Wsp) or have not undergone extensive recombination and directional selection (FtsZ, cell division protein). Wsp protein sequences exhibit considerable divergence (mean conservation $= 0.85$), with very few sites in a row being completely conserved (fig. 5A). In contrast, FtsZ is relatively conserved (mean conservation $= 0.94$), and most of the divergence is clustered at the C-terminus (fig. 5B). Mean conservation for the Cif protein sequences was lower than Wsp—0.83 for Type I CifA (fig. 5C) and 0.82 for Type I CifB alleles (fig. 5E, table 3). When all Cif alleles were considered, mean conservation was even further reduced—0.58 for CifA (fig. 5D) and 0.43 for CifB (fig. 5F). The lower average conservation of CifB genes is in part due to the many insertions and deletions in the alignment, and the missing C-terminal deubiquitylase region, ModB-3, of the Type II and III alleles. Thus, several CifB proteins apparently lack this activity, and whether these variants cause CI remains to be determined. Although the CifB proteins are highly divergent, the catalytic residue (red dot in fig. 5E and F) in the deubiquitylating module of CifB is unique to and completely conserved for the Type I alleles.

The Cif proteins have extensive amounts of diversity, with completely conserved amino acids distributed across the length of the protein, and not confined to any particular

Table 3

<table>
<thead>
<tr>
<th>Protein</th>
<th>Region*</th>
<th>Type I</th>
<th>All</th>
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<tr>
<td>CifA</td>
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<tr>
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<td>0.70</td>
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<tr>
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<tr>
<td></td>
<td>CifB</td>
<td>0.82</td>
<td>0.43</td>
</tr>
</tbody>
</table>

*Module number is defined in table 2.
*Only annotated in Type I.
regions (fig. 5C–F, supplementary tables S3–S6, Supplementary Material online). There were significant differences in the level of conservation between modules and nonmodule regions for the Type I alignments of both CifA ($F_{3,490} = 4.276, P = 0.0054$) and CifB ($F_{3,1195} = 9.703, P = 1.5e-06$) (table 3). The only modules that had significantly higher conservation than the nonmodule regions of the alignment were ModB-1 ($P = 0.0173$) and ModB-2 ($P = 0.0011$). The wMel strain contains the (P)D-(D/E)XK motif in ModB-1 (blue dots in fig. 5E and F) (Kosinski et al. 2005), but it is less than 80% conserved across strains despite the higher average conservation of this module. In contrast, wMel does not contain the catalytic motif in ModB-2, also a PDDEXX nucleotide-like domain. The ModB-2 (P)D-(D/E)XK motif is present in Type IV alleles such as WOPip2 that when mutated no longer induces growth defects in yeast (Beckmann et al. 2017). ModA-3 is significantly less conserved than the nonmodule regions of Type I CifA ($P = 0.0300$).

**Cif Module Presence Generally Predicts Reproductive Phenotype**

We used the wMel-predicted Cif modules as a seed to search for the presence of homologous modules across Wolbachia genome sequences using TBLastN (fig. 6), with the intent of discovering cif-like regions or remnants in strains with other phenotypes. In strains with more divergent Cif Types, we report modules that were expected based on the HHpred results, but not recovered with TBLastN due to sequence divergence from WOMeLB. Additionally, we recover homologous modules outside of the annotated cif open reading frames, such as the chromosomal region with a ModB-3 (Ulp1-like) region in wNo. The ModB-3 in wNo module is generic, found within a hypothetical protein (WP_041581315.1). Whether or not these cif-like regions outside of prophage WO contribute to CI remains to be determined. All Supergroup A and B strains, with the exception of wAu and wTpre (non-CI inducing strains), contained at least one recovered module.

Importantly, all strains that are known to be capable of inducing or rescuing CI have two or more recovered modules, though they do not necessarily have ModB-3, which contains the catalytic residue implicated in CI function (Beckmann et al. 2017). The non-CI strains have fewer recovered modules: One module (ModB-2) in wUni, and no modules in wCle, wAU, wTpre and the nematode-infecting strains (Supergroups C and D). wUni is a unique case, where we identified cif alleles in the genome, but recovered only one module. Most wUni modules are either missing (fig. 4A) or divergent enough from WOMeLB that they were not considered a positive match. wAlbB and wNo, both CI-inducing strains with Type III and IV alleles, have fewer recovered modules, but this is congruent with the more divergent nature of those Cif Types. It is notable that despite the phylogenetic distance from WOMeLB, more modules are recovered from the CI-inducing strains than the non-CI inducing strains within the same Type.

The high number of modules in wSuzi and wRi is due to the presence of a duplicated set of Type I variants. We recovered many modules in wSuzi, which is a strain not known to induce CI, but is sister to wRi, which can induce CI (Hamm et al. 2014; Cattel et al. 2016). This discrepancy between cif presence and absence of a reproductive phenotype might be explained by the disrupted Type II cif in wSuzi. The split WOSuziC sequence was concatenated to allow for a more robust phylogenetic reconstruction (fig. 4), but it is in fact disrupted by a transposase (Conner et al. 2017). However, having a functional set of Type I cif alleles appears to be sufficient for CI-induction in other strains (Beckmann et al. 2017; LePage et al. 2017), so it is not clear how inactivation of the Type II alleles here may affect the final CI phenotype in wSuzi. Strain wDi, infecting the Asian citrus psyllid Diaphorina citri, has no identified reproductive phenotype, but only contains two modules: ModB-1 and ModB-3. For wHa, we recovered duplicates of all the modules. These represent a highly disrupted copy of the gene set harboring frameshifts that were annotated as pseudogenes.

The lack of evidence for homologous cif genes in the C, D, and F Supergroup Wolbachia agrees with previous findings (LePage et al. 2017) that CI-function is restricted to the A + B Supergroup clade (likely due to WO phage activity), and the absence of WO phages for the nematode-infecting strains (Gavotte et al. 2007). The loss of CI within the A and B Supergroups is likely a derived trait due to the rapid evolution of prophage WO (Ishmael et al. 2009; Kent, Salichos, et al. 2011) and relaxed selection after transition to a new reproductive phenotype. The low number of modules identified in such strains is consistent with gene degradation and loss. Additionally, we recover no cif-like regions in Cardinium, a member of the Bacteroidetes, and an independent transition to a CI phenotype (Penz et al. 2012).

To further explore the conservation of the cif genes across the sequenced Wolbachia, and to uncover diversity that may be present in other genomes, we searched the WGS databases for recently sequenced genomic scaffolds from Wolbachia infecting the Nomada bees (wNleu, wNla, wNpa, and wNfe) (Gething and Bleidorn 2016), Drosophila incomputa (wInC_Cu) (Wallau et al. 2016), and Laodelphax striatellus (wStri) (GenBank accession number NZ_LRUH00000000.1) using HMMER. Only for wStri do we have direct evidence of CI induction (Noda et al. 2001) yet the wStri WGS projects contain only one cif locus (CifB) with an unusual structure not found in any of the other Types. On the basis of HHPred analyses, the wStri homolog contains a deubiquitylase region in the middle of the protein, with two downstream regions that have homology to glucosyl transferases and lipases, respectively (supplementary fig. S4, Supplementary Material online). The wnc_Cu WGS project contained one each of CifA and CifB alleles. The CifA allele from wnc_Cu is a typical Type
I protein containing three modules: An N-terminal Catalase-rel domain and an internal DUF3242 domain, followed by the STE-like transcriptional factor domain. Because these are incomplete genome projects, it is possible that other \textit{cif} homologs have been missed due to the current sequencing coverage. Alternatively, it is possible that other, as yet undiscovered, mechanisms of reproductive manipulation exist in these strains. In contrast, the \textit{Nomada}-associated \textit{Wolbachia} contain a large repertoire of \textit{cif} homologs, including Types I, II, IV, and several homologs with variations on the Type IV domain architecture for \textit{CifA} (supplementary fig. S4, Supplementary Material online). Many of the \textit{CifB} homologs are disrupted Type I variants that contain the deubiquitylase-like domain, but not the nuclease-like domains (supplementary fig. S4, Supplementary Material online).

**Discussion**

We explored three key features of \textit{cif} evolution: 1) The operon hypothesis, 2) potential novel functions across the \textit{cifA} and \textit{cifB} phylogenies, and 3) the conservation and diversity of \textit{cif} alleles across strains with different host-manipulation phenotypes. We provide multiple lines of evidence that although \textit{cifA} and \textit{cifB} can be cotranscribed, they are divergently transcribed in \textit{w}Mel during host development, suggesting a more complex regulation of gene expression than found in classical operons. Indeed, \textit{cifB} transcription has a significant negative correlation with \textit{cifA}. In \textit{Escherichia coli}, operons frequently have internal promoters and terminators that result in different units of transcription, which are preferentially used during certain conditions (Conway et al. 2014). Although \textit{cifB} is
expressed at about 1/10 the level of cifA across all life cycle stages, the significant negative correlation in their levels suggests that the same factor(s) could upregulate cifA while downregulating cifB, or a cifA encoded RNA or protein may inhibit cifB expression or vice versa. The new annotations for cifA alleles, including a Puf family RNA-binding-like domain and STE transcription factor, could theoretically play a role in inhibiting cifB expression. More detailed analyses from a variety of strains, cif Types, and conditions would help develop a comprehensive understanding of the factors regulating expression of these genes and the CI phenotype.

It is especially interesting that cifA and cifB synteny is maintained across prophage WO regions, despite the high level of recombination and rearrangements in prophage WO and Wolbachia genomes (Baldo, Bordenstein, et al. 2006; Kent, Funkhouser, et al. 2011; Ellegaard et al. 2013). Although it is not yet clear why cifA and cifB homologs maintain their syntenic orientation, given that they have very different absolute and relative expression levels, we hypothesize that this feature can be attributed to 1) their location within prophage WO and/or 2) functions associated with the ability of cifA and cifB to act synergistically to induce CI (LePage et al. 2017), or 3) with the potential antagonism of cifA on cifB transcripts or transcription. For example, they could share the same promoter, but in addition to the Rho-independent terminator, cifA may further inhibit cifB expression by binding to the intergenic region between them, causing the polymerase to terminate. Such a model would be consistent with both the absolute expression differences and the negative correlation. Alternatively, cifA and cifB may have different promoters, but a bicistronic message occurs because of an imperfect hairpin terminator between the genes. We conclude that cifA and cifB are cotranscribed but not coregulated as in a classical operon, and do not act strictly as a toxin–antitoxin system due to the requirement of both Cif proteins for the induction of CI in arthropods. Determining how cifA and cifB expression is regulated in the insect host will advance an understanding of both the basic biology of CI and vector control programs that deploy CI to control disease transmission.

Despite the conservation of gene order, Cif proteins showed extensive amounts of divergence and differences in domain structure as previously reported (LePage et al. 2017). Here, the levels of amino acid conservation in the Cifs are lower than FtsZ and Wsp, the latter of which is known to lower than FtsZ and Wsp, the latter of which is known to

Based on what is known about Wolbachia biology, some of the protein domains may be especially good candidates for further study and in vivo functional characterization. Predicted PDDEXK-like nuclease domains are present in all four CifB Types. Given the predicted interaction of these domains with DNA (Kosinski et al. 2005), and the presence of these domains across CifB proteins, determining whether and how these regions interact with host (Wolbachia or insect) DNA, and whether or not they contribute to CI function would be useful in understanding the consistent presence of this module. Mutating the predicted catalytic site of the nuclease region in wPip’s Type IV CifB (aka CinB) reduces toxicity in yeast (Beckmann et al. 2017). However, this catalytic residue is not conserved, so further exploration of nuclease function across more divergent alleles will be useful. As aforementioned, many of the CifA alleles encode Puf family RNA-binding-like domains, which have previously been implicated in mRNA localization and transcriptional regulation (Quenault et al. 2011). This RNA binding-like domain is found upstream of an STE transcription factor-like domain and could provide a promising direction for understanding the complicated transcriptional dynamics of the cif genes.

Wolbachia strains that have lost CI have a strong signature of cif gene degradation and loss, consistent with their role in CI. The two parthenogenesis-inducing strains (wTpre and wUnin) appear to be at different places in this process of gene loss, with divergent Cif amino acid sequences recovered for wUnin, but no modules identified in wTpre. There are several explanations for this. wUnin is likely a more recent transition to parthenogenesis, as it is closely related to a CI strain (wVitA) (Baldo, Dunning Hotopp, et al. 2006; Newton et al. 2016). In comparison, wTpre is part of a unique clade of Wolbachia that all induce parthenogenesis in Trichogramma wasps (Rousset et al. 1992; Werren et al. 1995; Schiltzuizen and Stouthamer 1997). This strain has lost its WO phage association and only has relics of WO phage genes (Gavotte et al. 2007; Lindsey et al. 2016). Additionally, the two strains that independently transitioned to the parthenogenesis phenotype have evolved separate mechanisms for doing so (Stouthamer and Kazmer 1994; Gottlieb et al. 2002). Differences in time since transition to the parthenogenesis phenotype, phage WO associations, and mechanisms of parthenogenesis induction likely all play a role in the rate of cif gene degradation.

Although cifA and cifB are prophage WO genes, not all CI-inducing strains have a complete prophage. Indeed, the wRec strain of Wolbachia in Drosophila recens is one such example where approximately three quarters of prophage WO genes were eliminated (Metcalf et al. 2014), previously resulting in failed detection of WO presence (Bordenstein and Wernegreen 2004). However, genomic analyses of phage WO particles from wasps and moths revealed that several
genes packed in the genome of phage WO particles (Bordenstein SR and Bordenstein SR 2016) are in fact retained in prophage WO of wRec (Metcalf et al. 2014), including cifA and cifB. Genes in prophage WO relics are apparently a source of host-manipulation across Wolbachia genomes. Additionally, there is considerable variation in the strength of CI across different Wolbachia strains (Veneti et al. 2003). CifA and CifB have an additive effect on the strength of CI (LePage et al. 2017), so it is possible that the level of cif expression, or the ratio of cifA and cifB transcripts across development, are ways in which CI strength is adjusted. The rapidly evolving nature of the Cif proteins may affect other ways in which they function in the host. For example, in Type I CifB proteins that have the essential deubiquitylase residue, other sequence variation may affect the ability to bind with CifA, locations of posttranslational modifications, or the ability to be efficiently localized to the host nucleus. Additionally, the level of CI is often host-dependent (Bordenstein and Werren 1998; McGraw et al. 2001), possibly a result of how well Wolbachia replicate in the host, and/or the specificity of Cif proteins with the host target, which is currently unknown. There are also environmental conditions that affect the strength of CI, and they likely do so by affecting Wolbachia titers and resulting Cif expression in the host (Clancy and Hoffmann 1998; Yamada et al. 2007).

On the basis of our analyses, we propose three avenues of research on the function of the Cif proteins. First, functional confirmation of the newly annotated modules will be important in understanding how these genes function enzymatically. In total, we predict six modules in the Cif protein sequence homologs, with varying degrees of confidence (supplementary tables S1 and S2, Supplementary Material online). For some of these modules, straightforward experiments can be designed in model systems (such as Saccharomyces) to determine whether their predicted function is correct, as has been done for the deubiquitylase domain of CidB (Beckmann et al. 2017) and countless other bacterial effectors (Kramer et al. 2007; Siggers and Lesser 2008; Archuleta 2011). The necessity and importance of these modules to the CI phenotype can be assessed in the Drosophila model, where the induction of the phenotype and rescue is straightforward (LePage et al. 2017). Second, detailed characterization of cif gene regulation will be important for understanding CI expression and penetrance, thus informing vector control programs that rely on proper expression of the CI phenotype, often in a transfected host. Finally, we suggest that although the discovery of these genes is fundamental, it is clear from this analysis that we have not comprehensively evaluated or identified the mechanisms behind CI and other reproductive manipulations. The gene characterization analyses described here reveal new and relevant annotations, but with many regions of unknown function across all of the phylogenetic Types, missing deubiquitylase domains in particular CI strains, and a coevolving, phylogenetic relationship across the Cif trees. Importantly, the locus, presumably expressed in the female insect infected with a compatible Wolbachia, and mechanism behind rescuing CI are still unknown, as is the exact mechanism by which all Cif proteins induce CI. Therefore, the recent discovery of these CI genes and their sequence characterization described here pave the way for investigating key mechanisms of the Wolbachia–host symbiosis.

**Supplementary Material**

Supplementary data are available at Genome Biology and Evolution online.

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**Literature Cited**


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