Unique clade of alphaproteobacterial endosymbionts induces complete cytoplasmic incompatibility in the coconut beetle

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Maternally inherited bacterial endosymbionts in arthropods manipulate host reproduction to increase the fitness of infected females. Cytoplasmic incompatibility (CI) is one such manipulation, in which uninfected females produce few or no offspring when they mate with infected males. To date, two bacterial endosymbionts, Wolbachia and Cardinium, have been reported as CI inducers. Only Wolbachia induces complete CI, which causes 100% offspring mortality in incompatible crosses. Here we report a third CI inducer that belongs to a unique clade of Alphaproteobacteria detected within the coconut beetle, Brontispa longissima. This beetle comprises two cryptic species, the Asian clade and the Pacific clade, which show incompatibility in hybrid crosses. Different bacterial endosymbionts, a unique clade of Alphaproteobacteria in the Pacific clade and Wolbachia in the Asian clade, induced bidirectional CI between hosts. The former induced complete CI (100% mortality), whereas the latter induced partial CI (70% mortality). Illumina MiSeq sequencing and denaturing gradient gel electrophoresis patterns showed that the predominant bacterial detected in the Pacific clade of B. longissima was this unique clade of Alphaproteobacteria alone, indicating that this endosymbiont was responsible for the complete CI. Sex distortion did not occur in any of the tested crosses. The 1,160 bp of 16S rRNA gene sequence obtained for this endosymbiont had only 89.3% identity with that of Wolbachia, indicating that it can be recognized as a distinct species. We discuss the potential use of this bacterium as a biological control agent. Biological control | reproductive isolation | speciation | symbiont | Wolbachia

Bacterial endosymbionts in arthropods influence host reproduction in various ways. Because bacterial endosymbionts are transmitted mainly through the female host, they manipulate host reproduction to increase the fitness of infected females. Such manipulation was first detected in the early 1970s in Culex pipiens infected with Wolbachia pipientis, a member of the Alphaproteobacteria class (1). W. pipientis induces cytoplasmic incompatibility (CI), in which uninfected females or those infected with a different strain than a male’s Wolbachia produce few or no offspring when they mate with infected males. Several other manipulation phenotypes have been identified in Wolbachia and other bacterial species, including partenogenesis induction, in which infected females produce daughters without fertilization by males in a haplodiplod system (Wolbachia, Ricketsia sp., Cardinium hertigi); feminization of genetic males, in which genetic male embryos develop phenotypically as females (Wolbachia, C. hertigi); male killing, in which male embryos are killed during development (Wolbachia, Ricketsia sp., Flavobacterium sp., Spiroplasma xoletis, Spiroplasma poulsonii); and oogenesis, in which uninfected females cannot produce mature oocytes (Wolbachia) (2). Although several bacterial species are known to manipulate host reproduction, only Wolbachia shows all of these phenotypes. CI is the most common phenotype of Wolbachia. Although the molecular mechanisms are unknown, CI can be explained by the “modification-rescue” system. Wolbachia “modifies” sperm in the testes, and the sperm develops abnormally, resulting in death of the embryo when appropriate Wolbachia is not present in the eggs to “rescue” the embryo from the modification (3). The other known CI inducer, Cardinium (Bacteroidetes), induces only a low level of CI and never causes 100% mortality in incompatible crosses (4–7). Although CI has important effects on host speciation (3, 8), and CI-inducing microbes might be useful for biological control (9), research focusing on such issues has been limited to Wolbachia.

Brontispa longissima (Gestro) (Coleoptera: Chrysomelidae) is a serious pest of coconut palm (Cocos nucifera L.) that is native to Indonesia and New Guinea (10). Around 2000, B. longissima was recorded in southeast and east Asia and has caused serious damage to coconut palms (11). Based on mitochondrial cytochrome oxidase I gene sequences, B. longissima comprises two cryptic species: the Asian clade, distributed over a wide area including Asia and the Pacific region, and is the Pacific clade.

Significance

Maternally inherited bacterial endosymbionts in arthropods manipulate host reproduction to increase the number of infected females. Cytoplasmic incompatibility (CI) is one such manipulation, in which infected females can produce offspring by mating with both infected and uninfected males, but uninfected females cannot or seldom produce offspring with infected males. Two bacterial endosymbionts, Wolbachia and Cardinium, are known CI inducers. Here we report a third CI inducer that belongs to a unique clade of Alphaproteobacteria. This bacterial clade was found to cause complete CI between two clades of the coconut beetle, a serious invasive pest of coconut palms. We discuss the potential use of this bacterium as a biological control agent and its effects on speciation of the coconut beetle.


The authors declare no conflict of interest.

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distributed in a limited area (12). Recent invasions and outbreaks have been reported only for the Asian clade. The two clades copulate with each other, and sperm are successfully delivered to the spermatheca, but hatchability is much lower than that of intraclade crosses, indicating that CI occurs between them (12); however, what induces this CI remains unknown.

Here we show that different bacterial endosymbionts, a unique clade of Alphaproteobacteria in the Pacific clade and Wolbachia in the Asian clade, induce bidirectional CI between the two B. longissima clades. The former induces complete CI (100% offspring mortality), whereas the latter induces partial CI (70% mortality).

**Results**

**Crossing Experiments.** To determine the effect of bacterial endosymbionts on CI between the two clades of *B. longissima*, we examined egg hatch rates among crosses of antibiotic-treated (indicated by −) or untreated (indicated by +) insects. Two populations were used: *B. longissima* collected in Lospalos, East Timor, which belongs to the Pacific clade (Lp), and *B. longissima* collected in Dili, East Timor, which belongs to the Asian clade (Di). Egg hatch rates in the untreated intraclade crosses were significantly lower than those in the untreated intraclade crosses (Fig. 1). Hatch rates were significantly lower in the DI* female × Lp* male crosses (0%; n = 9 crosses and 181 eggs) compared with the Lp* female × DI* male crosses (mean ± SEM, 30.8 ± 5.2%; n = 7 crosses and 75 eggs) (Fig. 1). When bacterial endosymbionts were eliminated by tetracycline treatment in both parental females and males, egg hatch rates in the interclade crosses increased dramatically, reaching levels similar to those of the intraclade crosses (Fig. 1). When parental males were treated with tetracycline but females were untreated, hatch rates also increased significantly in both the intraclade and interclade crosses, whereas when females were treated but males were untreated, hatch rates did not increase (Fig. 1). In the intraclade crosses, hatch rates decreased dramatically when females were treated with tetracycline but males were not. Antibiotic-treated females showed lower hatch rates in Lp* (0%; n = 6 crosses and 64 eggs) than in Di (mean ± SEM, 29.5 ± 5.1%; n = 7 crosses and 132 eggs) (Fig. 1). Dissection of females to examine the presence of sperm in the spermatheca showed that 4 of 117 pairs did not copulate during the 1-wk period: one DI* female × DI* male, one Lp* female × DI* male, one Lp* female × Lp* male, and one DI* female × Lp* male. Data for these pairs were omitted from our analysis. Sex ratios of offspring did not significantly differ from 1:1 in all tested pairs (P > 0.05, binomial test) (Fig. 1).

The number of eggs produced per female in 7 d did not differ between antibiotic-treated and untreated Lp females (Mann-Whitney test, U = 390.5, P = 0.451). Antibiotic-treated DI females produced fewer eggs than untreated females (U = 295.5, P = 0.022). Lp females produced fewer eggs than DI females (U = 786.5, P < 0.001 for untreated females vs. U = 598.5, P < 0.001 for antibiotic-treated females) (Fig. 2).

**Sequences and Phylogenetic Relationships of Bacterial Endosymbionts in B. longissima.** To detect and analyze phylogenetic relationships of bacterial endosymbionts, total genomic DNA was extracted from the testes or ovaries of Di and Lp adults, and bacterial 16S rRNA gene sequences were analyzed using universal bacterial 16S rRNA primers. For Lp, 1,160-bp 16S rRNA gene sequences were obtained from 21 clones derived from three males and three females (LC177357–LC177361). The closest BLASTn match (for LC177357) was with a sequence of unidentified bacterial gut symbiont of a beetle (GU815124.1) with 97.9% (1,129 of 1,153) identity. For Di, 1,159-bp 16S rRNA gene sequences were analyzed using universal bacterial 16S rRNA primers. For Lp, 1,159-bp 16S rRNA gene sequences were obtained from 21 clones derived from three males and three females (LC177357–LC177361). The closest BLASTn match for LC177357 was with a sequence of unidentified bacterial gut symbiont of a beetle (GU815124.1) with 97.9% (1,129 of 1,153) identity. The closest BLASTn match with an identified sequence was with Wolbachia (EU499316.1), with 89.3% (1,038 of 1,162) identity. For Di, 1,159-bp 16S rRNA gene sequences were

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**Fig. 1.** Crossing tests among two populations in the absence or presence of antibiotic treatment in *B. longissima*. Lp, *B. longissima* collected in Lospalos, East Timor (Pacific clade); Di, *B. longissima* collected in Dili, East Timor (Asian clade); −, antibiotic-treated insects; +, antibiotic-untreated insects. Error bars indicate SEM. Means with the same letters do not differ significantly (Tukey–Kramer test, P > 0.05). Gray parts indicate percentage of females in each cross. In pairs, n = 9 for Di* Lp* (female × male, the same in the following), Di* Lp+, and n = 8 for Lp* Di+, Di* Lp+, and n = 6 for Di* Lp, Lp* Di+, Lp* Di+, n = 5 for Lp* Di, and n = 7 for the rest of the crosses. In the number of eggs, n = 165, 93, 75, 181, 131, 84, 57, 151, 131, 97, 76, 175, 132, 64, 89, and 161 for Di* Di+, Lp* Lp+, Lp* Di+, Di* Di+, Lp* Lp+, Lp* Di+, Di* Lp, Lp* Di+, Lp* Di+, Di* Di+, Lp* Lp+, Lp* Di+, and Di* Di+, respectively.

**Fig. 2.** Number of eggs produced every 7 d per female of *B. longissima*. Lp, *B. longissima* collected in Lospalos, East Timor (Pacific clade); Di, *B. longissima* collected in Dili, East Timor (Asian clade); −, antibiotic-treated insects; +, antibiotic-untreated insects. Error bars indicate SEM. Means with the same capital letters do not differ significantly among antibiotic-treated and untreated females in the same clade, and means with the same lowercase letters do not differ significantly between Lp and Di in the same treatment (P > 0.05, Mann–Whitney U test). n = 30 for Di* and Di*, n = 29 for Lp*, and n = 24 for Lp⁻.
obtained from 22 clones derived from three males and three females (LC177362). The closest BLASTn match was with a sequence of *Wolbachia* (KT273277.1), with 99.7% (1,155 of 1,159) identity. The phylogenetic tree showed that bacterial endosymbionts detected from *Lp* beetles formed a unique clade in Alphaproteobacteria with 100% bootstrap support, whereas endosymbionts detected from *Di* beetles belonged to *Wolbachia* (Fig. 3). A maximum likelihood tree for concatenated multilocus sequence typing (MLST) sequences (garB, coxA, hcpA, ftsZ, and fbpA) indicated that *Di* beetles were infected with *Wolbachia* endosymbionts belong to supergroup B (Fig. S1).

Elimination of the endosymbionts by antibiotic treatment was confirmed for eight adults from each cross using PCR with specific primers for *Lp* (designed in the present study) and primers for *Wolbachia* (13) for *Di* (Fig. S4).

Inheritance of *Lp* endosymbionts was examined by PCR with the specific primers for offspring of intraclace crosses. A strong band was detected from offspring larvae of the *Lp* female × *Lp* male crosses, whereas no band was detected in the offspring eggs from the *Lp* female × *Lp* male crosses (Fig. 4B).

**Analysis of Microbial Composition.** We performed Illumina MiSeq sequencing to check for the possible presence of other bacteria not detected by the foregoing cloning procedure. We obtained a total of 215,058 (mean ± SEM, 35,843 ± 5,393 reads per sample) high-quality pair-end 16S rRNA reads from the testes or ovaries of *B. longissima*. Operational taxonomic units (OTUs) were generated using USEARCH (14) with identity ≥97%. After excluding the OTUs containing 18S rRNA genes of the host beetles, a total of 1,504 OTUs were generated. The predominant OTU in both the OTUs containing 18S rRNA genes of the host beetles, a total of 1,504 OTUs were generated. The predominant OTU in both female and male *Lp* was S001909539, uncultured bacterium alone (Fig. 5). Although one predominant OTU, S004255337, a *Wolbachia* endosymbiont of *Coelostomidia zealandica*, was detected in *Di*; other Alphaproteobacteria were found as well (Fig. 5). Phylogenetic analysis revealed that major Alphaproteobacteria detected in *Di* belonged to different clusters within *Wolbachia* (Fig. S2). The number of detected OTUs was rather small in antibiotic-treated insects (Fig. 5).

**PCR and Denaturing Gradient Gel Electrophoresis.** In addition to the Illumina MiSeq sequencing, we performed PCR and denaturing gradient gel electrophoresis (DGGE) analyses to check for the possible presence of other bacteria. DGGE patterns of the variable V3 region of 16S rRNA genes showed that one band disappeared due to the antibiotic treatment in both males and a female of *Lp*, and two closely spaced bands disappeared in *Di* (Fig. 6).

**Discussion**

In this study, we used crossing tests between different clades of *B. longissima* (*Di* and *Lp*) and analyses of the 16S rRNA gene sequences to demonstrate that bidirectional CI between the two clades is induced by different endosymbionts. We found that the complete CI (100% mortality) between *Di* females and *Lp* males was induced by a unique clade of Alphaproteobacteria, and that the partial CI (70% mortality) between *Lp* females and *Di* males was induced by *Wolbachia* (Figs. 1, 3, and 4A). With antibiotic treatment, the hatch rates in interclade crosses became as high as those in intraclade crosses (Fig. 1).

Several factors may cause reproductive isolation among sympatric populations, including a host shift (15), a change in breeding time (16), and infection with CI inducers (3, 8). *Nasonia* parasitoid wasps have three sibling species (*Nasonia vitripennis*, *Nasonia longicornis*, and *Nasonia giraulti*) and show bidirectional CI among the species, caused by different strains of *Wolbachia* (17, 18). In this case, *Wolbachia* may play a causal role in speciation (3, 19). Our findings regarding *B. longissima* suggest that this may be another case in which bidirectional CI causes speciation. Although we used *B. longissima* obtained from East Timor alone in this study, CI between the two clades occurs in coconut beetles from different locations; individuals from Papua New Guinea belonging to the Pacific clade show complete CI when crossed with those from Japan belonging to the Asian clade (12). In the Papua New Guinea beetles, we detected the same endosymbionts as in the beetles from *Lp* (Fig. S3). We also detected the same endosymbionts in beetles of the Pacific clade collected from different places, including Australia and Indonesia (Sumba) (Fig. S3).

No effects of this bacterium on host reproduction other than CI were found in the present study. Sex ratio of the two clades was maintained at ~50% in all crosses (Fig. 1), indicating that neither feminization nor male killing was caused by this bacterium in *B. longissima*. Because the same *Wolbachia* species may induce different phenotypes in different host species (20, 21), further research on the phenotypes of this Alphaproteobacterium in other insect species is warranted.
Our phylogenetic tree indicates that bacterial symbionts in Lp form a clear monophyletic group in class Alphaproteobacteria, order Rickettsiales (Fig. 3). In bacteriology, similarities of ≥97% (22) or >98.7% (23) in 16S rRNA gene sequences are required for individuals to be classified as the same species. Sequences of bacterial symbionts of Lp have only 89.3% identity with a sequence of *Wolbachia* (EU499316.1), the closest BLASTn match of identified bacteria. Although further studies on morphologic or anatomic characteristics are needed to determine the taxonomic status of this Alphaproteobacterium, we can recognize it as a distinct species based on the sequences of 16S rRNA genes. This distinct species may elucidate the origin and relationships among supergroups of *Wolbachia* owing to its similarity. So far, the outgroups used for this purpose are too divergent, and this distinct species could serve as a suitable outgroup in a phylogenetic study of *Wolbachia* (24, 25).

Detection of bacterial symbionts using universal bacterial 16S rRNA primers for PCR and cloning identified only two endosymbionts, a unique clade of Alphaproteobacteria in Lp and *Wolbachia* in Di. Our findings indicate that only these bacterial endosymbionts are responsible for CI in the two clades, which is consistent with Illumina MiSeq sequencing and DGGE patterns showing only one predominant bacterium at the strain level and only one band in Lp (Figs. 5 and 6). For Di, however, Illumina MiSeq sequencing showed that major OTUs belong to different clusters within *Wolbachia* (Fig. S2). DGGE patterns also showed two closely spaced bands that disappeared with antibiotic treatment. Thus, we infer that Di is infected by more than one *Wolbachia* strain. Although this finding is inconsistent with the Di *Wolbachia* cloning and MLST sequence results (garB, coxA, hcpA, ftsZ, and fipA), the limited number of analyses (i.e., 22 clones from six beetles in the cloning and direct sequencing from two beetles in MLST) may explain the detection of only a single strain.

The use of the bacterial symbiont *Wolbachia* as a biological control agent against insect pests has been attempted. A simple technique is to release males infected with CI-inducing *Wolbachia* into target uninfected populations to suppress population growth of the target, as in the sterile insect technique (SIT) (9). Guaranteeing the sterility of males is essential in SIT, and strong CI expression can effectively suppress a target population. Various intensities of CI expression have been reported in the CI inducers *Wolbachia* and *P. longissima*.
Cardinium. In Wolbachia, 0–100% CI has been reported in different host species (26–28). In Cardinium, 70% hatchability has been reported in incompatible crosses in snake mites (5), and incompatible crosses in the parasitoid wasp Encarsia pergandiella produced only 7% and 38% of the number of offspring in compatible crosses and showed an 8% pupation rate (6, 7). Therefore, the previously unidentified alphaproteobacterial endosymbiont, which induces complete CI, is a strong candidate agent for biological control of the coconut beetle.

Different effects from CI caused by Wolbachia may contribute to pest control. Infection by Wolbachia decreases the fecundity of host insects, and releasing transinfected insects with low fecundity into the field may suppress the pest population (29–31). In B. longissima, egg production is higher in the Asian clade than in the Pacific clade (32). In this study, elimination of the previously unidentified alphaproteobacterial endosymbiont, which induces complete CI, is a strong candidate agent for biological control of the coconut beetle.

Materials and Methods

Insects. Two colonies of B. longissima maintained in our laboratory were used for this study. One colony was initiated from insects collected in Dili, East Timor in July 2014 (referred to as Di), and the other was initiated from insects collected in Lospalos, East Timor in July 2014 (referred to as Lp). Di belongs to the Asian clade, and Lp belongs to the Pacific clade (34). Insects were reared on fresh leaves of Phoenix canariensis Chabaud as described previously (12).

Antibiotic Treatment. To eliminate bacterial endosymbionts, newly hatched larvae were reared on an artificial diet, Diet C, as described previously (35) as a control or Diet C with tetracycline. After the artificial diet was autoclaved, larvae were reared on the artificial diet with tetracycline and X-Gal was used. At least four insects from each colony were subjected to colony PCR with primers pUC/M13 (forward, 5′-CGC CAG GGT TTT CCC AGT CAC GAC-3′; reverse, 5′-ACA CAG AAA TAA AAA TTC CTA C3′) and primers (13) for Di (forward wsp_F1, 5′-GTC CAA TAR STG ATG ARG AAA C3′; reverse wsp_R1, 5′-CGY CAC CAA YAG YRC TRT AAAA-3′), with annealing temperatures of 58 °C. After crossing tests, the elimination was examined in eight antibiotic-treated adults of each clade. DNA extraction was conducted as described above.

In addition, inheritance of a unique clade of Alphaproteobacteria was examined in the offspring of crosses of Lp− females × Lp− males and Lp− females × Lp+ males. DNA was extracted from the whole body of first instar larvae in the former crosses and from whole eggs in the latter crosses, because no larvae hatched in the latter crosses. Ten larvae or eggs were analyzed for each cross.

Analysis of Microbial Composition. To check for the presence of other bacteria not detected by the foregoing cloning procedure, microbial communities in testes and ovaries were analyzed by Illumina MiSeq 16S rRNA gene sequencing. One male and one female reared on the artificial diet without tetracycline, and one male reared on the artificial diet with tetracycline were used for each clade. To amplify the variable V3 region of 16S rRNA gene, PCR was conducted with primers designed in the present study for Lp (forward L355F, 5′-GTC CAA TAR STG ATG ARG AAA C3′; reverse L749R, 5′-CGC CAG GGT TTT CCC AGT CAC GAC-3′; r e v e r s e, 5′-ACA CAG AAA TAA AAA TTC CTA C3′) (42) under the following conditions: an initial cycle at 94 °C for 1 min, followed by 35 cycles at 94 °C for 10 s, 55 °C for 15 s, and 72 °C for 10 s. The 20-μL PCR mixture contained 0.2 mM dNTP mixture, 0.5 μM each primer, 0.5 μL of TaKaRa PrimeSTAR GXL DNA polymerase, and 2.0 μL of DNA. PCR products were purified with NucleoSpin Gel and PCR Clean-up (Macherey-Nagel) and were inserted into the TA cloning vector (pMD20-T; TaKaRa Bio), which was used to transform Escherichia coli DH5α competent cells (Competent Quick; Toyobo). The blue-white colonies system with ampicillin and X-Gal was used. At least four insects from each individual were subjected to colony PCR with primers pUC/M13 (forward, 5′-CGC CAG GGT TTT CCC AGT CAC GAC-3′; reverse, 5′-TCA CAC AGG AAA CAG CTA TGA C3′). PCR products of the expected length (~1.4 kb) were sent to FASMAC (Atsugi, Japan) for sequencing. The obtained 16S rRNA gene sequences were aligned using CLUSTAL X (38). The phylogenetic tree was constructed by the neighbor-joining method with MEGA software (39); sequences from representatives of all alphaproteobacteria were included in the analysis. The evolutionary distances were computed by the Kimura two-parameter method, which accounts for differing rates of transitions vs. transversions (40). Nodal support was evaluated with 1,000 bootstrap resamplings (41). The sequence data were deposited in the DDBJ/EMBL/GenBank database (accession nos. LC177357–LC177362).

MLST Analysis. MLST analysis was conducted to determine the supergroup of Wolbachia infecting Di. Five ubiquitous genes (parB, coxA, fbpA, ftsZ, and rpoA) were analyzed in one male and one female (Fig. 51) (13). The sequence data were deposited in the DDBJ/EMBL/GenBank database (accession nos. LC164018–LC164022).

Confirmation of the Elimination of Bacterial Endosymbionts by the Antibiotic Treatment. Bacterial endosymbionts were detected using primers designed in the present study for Lp (forward L355F, 5′-GCT ATG CCG GGT GAG TGA TT3′; reverse L749R, 5′-ACA CAG AAA TAA AAA TTC CTA C3′) and primers (13) for Di (forward wsp_F1, 5′-GTC CAA TAR STG ATG ARG AAA C3′; reverse wsp_R1, 5′-CYG CAC CAA YAG YRC TRT AAAA-3′), with annealing temperatures of 58 °C. After crossin g tests, the elimination was examined in eight antibiotic-treated adults of each clade. DNA extraction was conducted as described above.

DNA Extraction, PCR, Cloning, Sequencing, and Phylogenetic Analysis of 16S rRNA. Total genomic DNA was extracted from the testes or ovaries of Di and Lp adults obtained from the colonies (three males and three females from each clade). Bees were dissected in distilled water to remove the testes or ovaries, which were placed on a clean bench under UV lighting for 5 min to eliminate possible surface contamination from other organs. DNA was extracted using the Qiagen DNeasy Blood and Tissue Kit. To amplify 16S rRNA genes, PCR was conducted with universal bacterial 16S rRNA primers (forward, 5′-GCT TAA CAC ATG GAA C3′; reverse, 5′-CCA TTG TAG CAC GTG T-3) (37), under the following conditions: an initial cycle at 94 °C for 1 min, followed by 30 cycles at 94 °C for 10 s, 55 °C for 15 s, and 72 °C for 90 s. The 20-μL PCR mixture contained 0.2 mM dNTP mixture, 0.5 μM each primer, 0.5 μL of TaKaRa PrimeSTAR GXL DNA polymerase, and 2.0 μL of DNA. PCR products were purified with NucleoSpin Gel and PCR Clean-up (Macherey-Nagel) and were inserted into the TA cloning vector (pMD20-T; TaKaRa Bio), which was used to transform Escherichia coli DH5α competent cells (Competent Quick; Toyobo). The blue-white colonies system with ampicillin and X-Gal was used. At least four insects from each individual were subjected to colony PCR with primers pUC/M13 (forward, 5′-CGC CAG GGT TTT CCC AGT CAC GAC-3′; reverse, 5′-TCA CAC AGG AAA CAG CTA TGA C3′). PCR products of the expected length (~1.4 kb) were sent to FASMAC (Atsugi, Japan) for sequencing. The obtained 16S rRNA gene sequences were aligned using CLUSTAL X (38).
DNA. PCR products were used for library production using the NEBNext Ultra DNA Library Prep Kit for Illumina (New England Biolabs Japan). The nucleotide sequence of the PCR products was determined using the Illumina MiSeq Reagent Kit v3.

We developed a custom pipeline for analysis of the 16S rRNA gene V3 region sequences. First, we excluded 16S reads that did not have primer sequences. Second, we trimmed the reads to maintain high quality of the first 100 bases using the FASTX-Toolkit (hannonlab.cshl.edu/fastx_toolkit). Third, we merged and filtered the pair-end reads using MeFiT (github.com/ckishimoto/ MeFiT) with an average quality value ≥ 25. We used all merged reads of the 16S rRNA gene V3 region to generate OTUs using the USEARCH program (www.drive5.com) (14) with ≥ 97% pairwise identity cutoff. The OTUs including 18S rRNA genes of the host beetles were detected by BLASTN match against the National Center for Biotechnology Information nucleotide database, and were excluded from the analysis. We constructed our own 16S rRNA gene database publicly available on Ribosomal Database Project release 11 (RDP) with ≥ 1200 bases. Each OTU was assigned to bacterial 16S rRNA by a BLASTN search with coverage ≥ 90% and pairwise identity ≥ 90%. All sequences of the 16S rRNA V3 region used in this study were deposited in the DNA Data Bank of Japan with accession numbers DRA005752.

PCR and DGGE. Along with the Illumina MiSeq sequencing, PCR and DGGE were performed to examine the possible presence of other bacteria. Two males and one female reared on the artificial diet with or without tetracycline were performed to examine the possible presence of other bacteria. Two release 11 (RDP) with base, and were excluded from the analysis. We constructed our own 16S rRNA gene database against the National Center for Biotechnology Information nucleotide database, and were excluded from the analysis. We constructed our own 16S rRNA gene database publicly available on Ribosomal Database Project release 11 (RDP) with ≥ 1200 bases. Each OTU was assigned to bacterial 16S rRNA by a BLASTN search with coverage ≥ 90% and pairwise identity ≥ 90%.

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Supporting Information

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Materials and Methods

Multilocus Sequence Typing Analysis. To determine the supergroup of Wolbachia infecting Di, multilocus sequence typing (MLST) analysis was conducted (13). DNA was extracted from one male and one female as described in the analysis of 16S rRNA gene. Five ubiquitous genes (gatB, coxA, hcpA, ftsZ, and fbpA) were amplified by using specific primers (13) under the following conditions: an initial cycle of 94 °C for 2 min; 37 cycles of 94 °C for 30 s, 54 °C (59 °C for ftsZ) for 45 s, and 72 °C for 1 min 30 s; and a final cycle of 72 °C for 10 min. The 20-μL PCR mixture contained 10× ExTaq Buffer (TaKaRa Bio), 0.2 mM dNTP mixture, 0.5 μM each primer, 0.5 U of TaKaRa ExTaq polymerase, and 2.0 μL of DNA. PCR products were purified with NucleoSpin Gel and PCR Clean-up (Macherey-Nagel) and were sent to FASMAC (Atsugi, Japan) for sequencing. Obtained sequences were aligned by using CLUSTAL X (37). The phylogenetic tree was constructed by the maximum likelihood method for concatenated MLST sequences by using MEGA6 software (38); sequences from other Wolbachia representatives were included. The nucleotide substitution model GTR + G was selected by using jModelTest 2.1.10. Nodal support was evaluated with 1,000 bootstrap resamplings (40). The sequence data were deposited in the DDBJ/EMBL/GenBank database (accession nos. LC164018–LC164022).

Phylogenetic Tree Construction. The full-length 16S rRNA gene sequences of the OTUs that belonged to Anaplasma in Fig. 5 were aligned using the multiple sequence alignment program MUSCLE in MEGA6 (38, 44). The phylogenetic tree was constructed using neighbor-joining method with 100 bootstrap replications. Wolbachia supergroups B, D, and F were distributed within Cluster 1. Supergroups A and E were distributed within cluster 2. The major species, S004255337 (Wolbachia endosymbiont of C. zealandica) in male and female B. longissima, belonged to cluster 1 (Fig. S2).

Presence of the Previously Unidentified Alphaproteobacterial Endosymbiont in B. longissima from Different Locations. Presence of the bacterial endosymbiont was tested as described in the methods section under “Confirmation of the elimination of bacterial endosymbionts by the antibiotic treatment” for the specimens from the following locations: Australia, Papua New Guinea, Indonesia (Sumba), and East Timor. Details of the specimens have been described previously (12). A total of 29 specimens that belong to the Pacific clade and two specimens that belong to the Asian clade were analyzed. DNA extracted from the thorax muscle was used (12) and a single band was detected in all specimens belonging to the Pacific clade (Fig. S3), indicating that all were infected with the previously unidentified alphaproteobacterial endosymbiont.

Fig. S1. Maximum likelihood phylogenetic tree of the concatenated MLST data (2,073 or 2,079 bp). Letters next to the species names indicate the supergroups. The ID codes and the sequence type numbers are obtained from the MLST database. Maximum likelihood bootstrap values based on 1,000 replicates (>50%) are given at the nodes.
Fig. S2. Phylogenetic distance tree of *Anaplasma* obtained from OTUs of male and female *Brontispa longissima* collected in Dili, East Timor (Asian clade). Circle colors indicate the top five numbers of OTUs in Fig. 5. The number at each branch indicates the bootstrap values (>50%) (100 replications). The phylogenetic tree was constructed in MEGA6 using neighbor-joining method, and based on the full length 16S rRNA sequences of the mapped OTUs that belonged to *Anaplasma* in Fig. 5.
Fig. S3. Detection of the previously unidentified bacterial endosymbiont in *B. longissima* from different locations: 1–3, Australia; 4–6, Papua New Guinea; 7–9, Indonesia (Sumba); 10 and 11, East Timor (Puno); 12–14, East Timor (Home); 15–17, East Timor (Raca); 18 and 19, East Timor (Dili). Specimens in 1–17 belong to the Pacific clade; specimens in 18 and 19, to the Asian clade.