Wolbachia genome integrated in an insect chromosome: Evolution and fate of laterally transferred endosymbiont genes

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Recent accumulation of microbial genome data has demonstrated that lateral gene transfers constitute an important and universal evolutionary process in prokaryotes, while those in multicellular eukaryotes are still regarded as unusual, except for endosymbiotic gene transfers from mitochondria and plastids. Here we thoroughly investigated the bacterial genes derived from a Wolbachia endosymbiont on the nuclear genome of the beetle Callosobruchus chinensis. Exhaustive PCR detection and Southern blot analysis suggested that ~30% of Wolbachia genes, in terms of the gene repertoire of wMel, are present on the insect nuclear genome. Fluorescent in situ hybridization located the transferred genes on the proximal region of the basal short arm of the X chromosome. Molecular evolutionary and other lines of evidence indicated that the transferred genes are probably derived from a single lateral transfer event. The transferred genes were, for the length examined, structurally disrupted, freed from functional constraints, and transcriptionally inactive. Hence, most, if not all, of the transferred genes have been pseudogenized. Notwithstanding this, the transferred genes were ubiquitously detected from Japanese and Taiwanese populations of C. chinensis, while the number of the transferred genes detected differed between the populations. The transferred genes were not detected from congenic beetle species, indicating that the transfer event occurred after speciation of C. chinensis, which was estimated to be one or several million years ago. These features of the laterally transferred endosymbiont genes are compared with the evolutionary patterns of mitochondrial and plastid genome fragments acquired by nuclear genomes through recent endosymbiotic gene transfers.

[Supplemental material is available online at www.genome.org. The sequence data from this study have been submitted to GenBank under accession nos. EF534574–EF534700 and EF560581–EF560584.]

Lateral, or horizontal, gene transfer is the exchange of genetic materials between phylogenetically distant, and reproductively and genetically isolated, organisms. Conventionally, lateral gene transfer was regarded as a rather exceptional process with limited contributions to organismal evolution. However, recent accumulation of microbial genome data has revealed that, particularly in prokaryotes, lateral gene transfer is a much more universal evolutionary process than ever thought, contributing to reorganization of prokaryotic genomes and acquisition of novel biological functions (Ochman et al. 2000; Koonin et al. 2001).

Meanwhile, lateral gene transfer involved in eukaryotes is still regarded as unusual. However, as more and more genomic sequences from diverse eukaryotes become available, it has turned out that lateral gene transfers can also occur in eukaryotes. According to the comprehensive review by Andersson (2005), there are two types of gene transfers in eukaryotes: (1) transfer of genes from mitochondria and plastids, which are organelles of prokaryotic endosymbiont origin, hereafter called

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Article published online before print. Article and publication date are at http:// www.genome.org/cgi/doi/10.1101/gr.7144908. "endosymbiotic gene transfer," and (2) transfer of genes from unrelated organisms, hereafter called "lateral gene transfer."

In the nuclear genomes of diverse eukaryotes, in addition to the ancient endosymbiotic gene transfers dating back to the origin of eukaryotes, a large number of organellar genomic DNA fragments of much more recent origins are frequently found. It is now widely accepted that such endosymbiotic gene transfers are ubiquitous and currently ongoing in diverse eukaryotes (Bensasson et al. 2001; Leister 2003; Timmis et al. 2004).

By contrast, lateral gene transfers from unrelated organisms are thought to be uncommon among eukaryotes (Brown 2003; Andersson 2005). The majority of eukaryotic lateral gene transfers are found from unicellular eukaryotes (for review, see Andersson 2005). Since many unicellular eukaryotes are phagotrophic on bacteria and other micro-organisms, they have a constant intracellular exposure to prokaryotic DNA, which may predispose them to incorporate foreign genetic material into their genomes (Doolittle 1998). Meanwhile, lateral gene transfers are rarer in multicellular eukaryotes like animals (Doolittle 1998; Andersson 2005).

Here it is tempting to assume that endosymbiotic gene transfers of nonorganellar origin might have occurred in some multicellular animals that constantly harbor endocellular microorganisms. Insects may represent such an animal group, wherein microbial endocellular associates occur universally. In particular, rickettsia-like endocellular α -proteobacteria of the genus *Wolbachia* are found in a wide range of arthropods and nematodes, including at least 20% of insect species. The endosymbionts are intimately associated with germ line cells of their hosts and are maternally transmitted, like organelles, through the host generations (O'Neill et al. 1997; Bourtzis and Miller 2003; Veneti et al. 2004). Considering the high levels of host–symbiont developmental integrity and ecological ubiquity, it is conceivable that such lateral gene transfers from the vertically transmitted endosymbionts might have occasionally occurred, thereby contributing to the host nuclear DNA dynamics to some extent, as does organelle-derived DNA.

Such an insect–endosymbiont lateral gene transfer was first reported from the adzuki bean beetle *Callosobruchus chinensis*, in which genes originating from a *Wolbachia* endosymbiont were genetically identified on the X chromosome of the insect (Kondo et al. 2002a). Recently, other cases of *Wolbachia*-derived lateral gene transfers have been emerging from genome sequencing projects of insects and nematodes (Fenn et al. 2006; Dunning-Hotopp et al. 2007). However, the evolutionary process and consequences of such host–symbiont lateral gene transfers have not yet been investigated in detail and thus are poorly understood.

In this study, we thoroughly investigated the magnitude, process, and evolution of the *Wolbachia*–insect lateral gene transfer in *C. chinensis*. A previous study suggested that the original donor bacterium of the transferred genes is a member of the *Wolbachia* group A (Kondo et al. 2002b), to which the *Wolbachia* strain wMel from *Drosophila melanogaster* belongs. Hence, we analyzed the laterally transferred *Wolbachia* genes in *C. chinensis* by making use of the wMel genome data (Wu et al. 2004) as reference information.

Results and Discussion

Exhaustive PCR detection of *Wolbachia* genes in the insect genome

We designed 222 pairs of PCR primers for specific amplification of 222 *Wolbachia* genes that cover the whole genome of *w*Mel, consisting of 1195 open reading frames (ORFs) (Wu et al. 2004). When DNA samples of the *Wolbachia*-infected insect strain jC were subjected to PCR amplification with these primer sets, 205 pairs amplified a DNA segment of expected size while 17 pairs did not (Supplemental Table S1). When DNA samples of the tetracycline-treated, *Wolbachia*-free insect strain jC^{Aus} were subjected to PCR amplification with the 205 primer sets, 57 pairs amplified a DNA segment of expected size (Supplemental Table S1). DNA sequencing confirmed that all of these PCR products are certainly *Wolbachia* genes (Supplemental Table S1). These results suggested that a large number of *Wolbachia* genes are present in the insect genome.

Exhaustive Southern blot detection of *Wolbachia* genes in the insect genome

By using long PCR amplification with these primers, we prepared 143 partial genomic segments of *w*Mel. These segments had an average size of 8.4 kb (ranging from 960 to 18,030 bp), and neighboring segments shared an average overlap of 0.4 kb (ranging

from 180 to 3731 bp) (Supplemental Table S2). In total, these DNA segments covered 1145 kb, about 90% of the 1268-kb genome of *w*Mel, except for two prophage regions where repetitive sequences prevented reliable PCR amplification. By using these segments as probes, DNA samples of the *Wolbachia*-free insect strain jC^{Aus} were subjected to Southern blot analysis. Of the 143 probes, 52 probes produced significant hybridization signals (Supplemental Table S2), suggesting that large regions of the *Wolbachia* genome are present in the insect genome.

Mapping of transferred Wolbachia genes on the wMel genome

Figure 1 shows a mapping of the PCR-positive sites and the Southern-positive regions on the *w*Mel genome. In general, the Southern-positive regions showed good agreement with the PCR-positive sites. Namely, in each of the regions of the 43 positive probes, one or more PCR-positive sites were located. In the regions of the other nine positive probes, no PCR-positive sites were identified. Probably these regions contain only a short stretch of *Wolbachia* gene, or *Wolbachia* genes in these regions have accumulated too many mutations to be amplified by PCR. Both the PCR-positive and Southern-positive regions were remarkably concentrated on the left half of the *w*Mel genome (see Fig. 1). These patterns suggested that a large *Wolbachia* genome segment was integrated into the genome of the jC^{Aus} insect.

Estimating the magnitude of lateral transfer of Wolbachia genes

Of 205 *Wolbachia* genes inspected by PCR, 57 genes were successfully amplified (Supplemental Table S1). DNA sequencing confirmed that all these amplifications were true positives. There is no doubt that some of the other genes were not amplified due

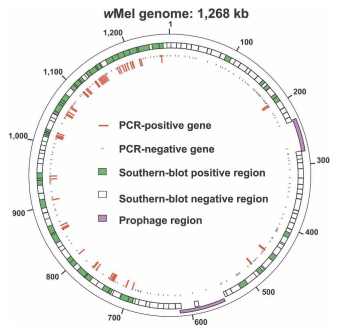


Figure 1. Wolbachia genes detected from the bean beetle *Callosobruchus chinensis* strain jC^{Aus} mapped on the whole genome of the *Wolbachia* strain wMel from *Drosophila melanogaster*. (*Inner* circle) PCR detection of *Wolbachia* genes. Red and black lines indicate PCR-positive and PCR-negative genes, respectively. (*Outer* circle) Southern blot detection of *Wolbachia* genome regions. Green and white blocks indicate Southern-positive and Southern-negative regions, respectively. Pink blocks show prophage regions.

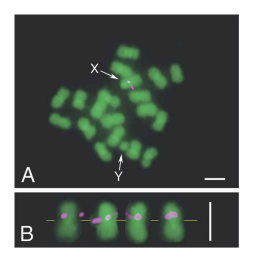


Figure 2. FISH detection of *Wolbachia* genes on the chromosomes of *Callosobruchus chinensis* strain C^{Aus} . (*A*) Metaphase chromosomes of a male insect. X and Y indicate X chromosome and Y chromosome, respectively. Note that *C. chinensis* is male-heterogametic with a karyotype of 2n = 20, XY. (*B*) Metaphase X chromosomes from different preparations, with yellow line indicating the position of centromeres, magenta, areas indicating *Wolbachia* gene signals; and green indicating insect chromosome some signals. Bars, 1 µm.

to primer mismatches, comprising pseudonegatives. Hence, the value of 28% (57/205) gene transfer in terms of the wMel genome must be an underestimate. Of 143 probes that covered nearly the whole genome of wMel, 52 probes gave positive signals in the Southern blot analysis of jC^{Aus} DNA (Supplemental Table S2). There is no doubt that, in many cases, only a partial region of these probes hybridized to the transferred *Wolbachia* genes located on the insect genome. Hence, the value of 36% (52/143) gene transfer in terms of the wMel genome must be an overestimate. On the basis of these data, we suggest that around 30% (28%–36%) of the wMel genes are present in the genome of the insect strain jC^{Aus} . Given that the wMel genome is 1268 kb in size, encoding 1195 ORFs (Wu et al. 2004), about 380 kb of the *Wolbachia* genomic region, encoding 360 genes, is expected to be located on the insect genome.

Physical mapping of transferred *Wolbachia* genes on the insect chromosomes

By using either the PCR-positive products or the Southernpositive regions as probes, we performed fluorescent in situ hybridization (FISH) analysis of the transferred *Wolbachia* genes on the chromosomes of the jC^{Aus} insects. A strong signal was consistently and specifically detected on the proximal region of the basal short arm of X chromosome (Fig. 2), confirming the previous report of X-linked inheritance of the *wsp* gene in the jC^{Aus} insects (Kondo et al. 2002a).

Phylogenetic relationship to other Wolbachia genomes

The *wsp* gene on the chromosome of *C. chinensis* was reported to belong to the *Wolbachia* group A (Kondo et al. 2002b). By using 56 genes commonly available for the *Wolbachia* strains *w*Mel (group A) (Wu et al. 2004), *w*Pip (group B) (Sanger Institute), and *w*Bm (group E) (Foster et al. 2005), we analyzed phylogenetic placement of the *Wolbachia* genes transferred to the *C. chinensis* chromosome. Molecular phylogenetic analysis of the total sequences indicated that the transferred genome is closely related

These results, namely, the concentrated distribution of the transferred genes on the *w*Mel genome (Fig. 1), their common physical location on the proximal region of the basal short arm of X chromosome (Fig. 2), and their consistent phylogenetic affin-

to the genome of *w*Mel rather than to the genome of *w*Pip (Supplemental Fig. S1). Distance analysis also showed that each

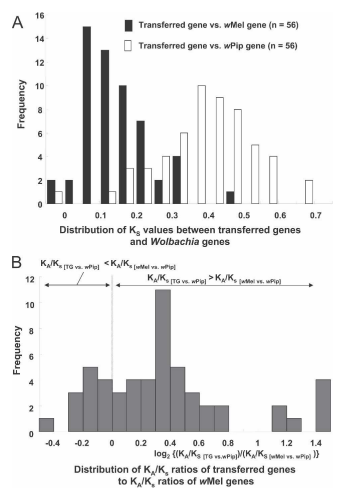


Figure 3. (*A*) Distribution of genetic distances from the transferred *Wolbachia* genes: *w*Mel vs. *w*Pip. In total, 56 partial gene sequences were subjected to the analysis (see Supplemental Table S1). Filled bars show the distribution of K_s values between transferred *Wolbachia* genes and wMel genes. Open bars indicate the distribution of K_s values between transferred *Wolbachia* genes and *w*Pip genes. (*B*) Comparison of molecular evolutionary rate between the transferred *Wolbachia* genes and the wMel genes. In total, 54 partial gene sequences were subjected to the analysis (see Supplemental Table S1). K_A/K_s values of transferred genes vs. wPip genes, and the distribution of the log ratios was plotted. In the plot, values less than zero indicate that the transferred genes exhibit lower K_A/K_s values than the corresponding *w*Mel genes, while values larger than zero imply that the transferred genes show higher K_A/K_s values than the corresponding *w*Mel genes. Note that the transferred genes generally exhibit higher K_A/K_s values than the corresponding *w*Mel genes.

ity to the *w*Mel genes (Fig. 3A), suggest that the *Wolbachia* genes present on the *C. chinensis* chromosome are not acquired through multiple lateral transfer events but are of single evolutionary origin from a *Wolbachia* strain that belongs to group A. Of course, however, the possibility of multiple acquisitions from the same or genetically close *Wolbachia* strains cannot be ruled out.

Structural disruptions in transferred Wolbachia genes

It is of evolutionary interest whether the transferred *Wolbachia* genes on the insect chromosome are functional. Of 57 transferred genes whose sequence was determined, 27 genes contained stop codon(s) and/or frame shift(s) (Supplemental Fig. S2; Supplemental Table S1). Hence, many of the transferred *Wolbachia* genes have been inactivated by structural disruptions.

Accelerated molecular evolution in transferred Wolbachia genes

If, as suggested above, most of the transferred *Wolbachia* genes have been disrupted and lost their function, it is expected that accelerated molecular evolution should be observed in these genes due to relaxed functional constraints. In order to test this idea, we compared K_A/K_S values of the transferred genes with those of the corresponding wMel genes, by using the corresponding genes of wPip as outgroup. The distribution of K_A/K_S ratios clearly indicated that the values for the transferred genes are generally higher than the values for the wMel genes (Fig. 3B). The tendency was highly significant statistically (randomization test, P < 0.00001).

RT-PCR assay of transcriptional activity in transferred *Wolbachia* genes

If some of the transferred *Wolbachia* genes are actually functional, then such genes must be transcribed into messenger RNAs. Hence, 57 transferred *Wolbachia* genes were subjected to RT-PCR detection (Supplemental Table S1). While no PCR products were obtained for 23 genes, amplified products of expected size were detected for the other 34 genes (Fig. 4A). Negative controls without reverse transcription did not exhibit these products, and DNA sequencing revealed that all these RT-PCR products certainly represent the target genes (data not shown). At a glance, these results indicated that, strikingly, more than half of the transferred *Wolbachia* genes are transcriptionally active on the insect chromosome.

Quantitative RT-PCR assay of transcriptional activity in transferred *Wolbachia* genes

Thus, we estimated the transcriptional activity of some of these genes by a quantitative RT-PCR technique. As internal controls for the quantitative assay, exon and intron regions of elongation factor 1α (*ef1* α) gene of the host insect were also quantified. We measured 26 RT-PCR–positive genes and nine RT-PCR–negative genes and found that expression levels of these genes were consistently low. Most of the genes showed expression levels of 10^{2} – 10^{3} copies per microgram of total RNA; the highest level was observed with the *ftsH* gene at about 10^{4} copies per microgram of total RNA, but all of these values were lower than the expression level of the *ef1* α intron (Fig. 4B). Since negative controls without reverse transcription always resulted in substantially zero values, transcripts of these genes may certainly exist, but their expression levels are very low. At this stage, it is obscure whether these

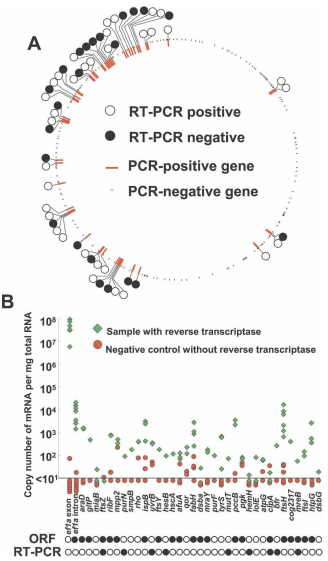


Figure 4. Transcriptional activity of the transferred *Wolbachia* genes. (*A*) RT-PCR assay of 57 transferred *Wolbachia* genes. The circle is identical to the inner circle of Figure 1. Open dots indicate genes with amplified products, while filled dots show genes without amplification. (*B*) Quantitative RT-PCR assay of 35 transferred *Wolbachia* genes, of which 26 were RT-PCR positive and nine were RT-PCR negative. An exon region and an intron region of the insect *ef1* α gene were also quantified as positive and negative controls. Each of the green diamonds shows an RNA sample from an insect treated with reverse transcriptase, while each of the red dots indicates an RNA sample from an insect without reverse transcriptase. In the ORF line, open dots indicate structurally intact ORFs, while dots show disrupted ORFs. In the RT-PCR line, open dots and filled dots indicate RT-PCR–positive and RT-PCR–negative, respectively.

very low expression levels of the transferred *Wolbachia* genes have biological significance. Considering that bacterial gene promoters usually do not work in eukaryotic cells, the possibility should be considered that these transcripts are not produced by a normal promoter-mediated transcriptional process but rather by a nonspecific leaky transcription at background levels. Future studies should focus on structure of the transcripts revealed by 5' and 3' RACE techniques and on spatial and temporal dynamics of the transcripts analyzed by in situ hybridization and tissuelevel quantitative RT-PCR.

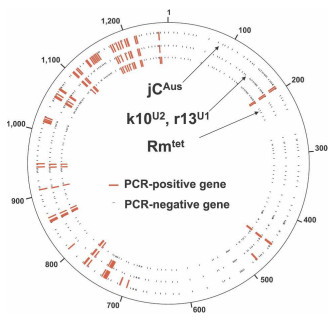


Figure 5. PCR detection of the *Wolbachia* genes from different strains of *Callosobruchus chinensis*. (*Outermost* circle) Genome map of *w*Mel as shown in Figure 1; (*second* circle) PCR detection of *Wolbachia* genes in the strain jC^{Aus}, as shown in Figure 1; (third circle) PCR detection of *Wolbachia* genes in the strains k10^{U2} and r13^{U1}; (*innermost* circle) PCR detection of *Wolbachia* genes in the strain Rm^{let}. Red and black lines indicate PCR-positive and PCR-negative genes, respectively.

Are there any host insects without transferred Wolbachia genes?

In Japanese populations of C. chinensis, diagnostic PCR survey revealed that 2.9% (12/410) insects did not possess the wsp gene originating from the transferred Wolbachia genome (Kondo et al. 2002b). We established two isofemale lines from such insectsk10 and r13-of different geographic origins. By tetracycline treatment and subsequent maintenance in the laboratory, we generated two derivative lines of the insects-k10^{U2} and r13^{U1}from which no wsp gene was detectable by PCR. Recently, in 80 individuals of C. chinensis from six Taiwanese populations, we identified some insects from which wsp gene of the transferred Wolbachia type was not detected (Kondo et al., unpubl.). We established an insect line named Rm and similarly generated a tetracycline-treated derivative line Rm^{tet} from which no wsp gene was detected by PCR (Supplemental Table S3). It was of evolutionary interest whether these insect strains are actually free of the transferred Wolbachia genes.

Variations in transferred *Wolbachia* genes in host insect populations

These insect strains $k10^{U2}$, $r13^{U1}$, and Rm^{tet} were subjected to PCR amplification with the 205 primer sets for *Wolbachia* genes. It turned out that these insects certainly possessed the transferred *Wolbachia* genes, but the gene repertoire was remarkably different between them (Fig. 5; Supplemental Fig. S3; Supplemental Table S1). Of the 205 genes inspected, 40 genes were detected from strains $k10^{U2}$ and $r13^{U1}$, while only 31 genes were found from strain Rm^{tet} . These numbers were remarkably smaller than 57 genes identified from the strain jC^{Aus} . These results indicated that the transferred *Wolbachia* genome fragment is universally found among natural populations of *C. chinensis* and that con-

siderable variations in the transferred *Wolbachia* gene repertoire are found within and between the populations.

Parsimonious estimation of evolutionary process leading to the variations in transferred *Wolbachia* genes

An interesting pattern was seen in these transferred Wolbachia gene repertoires. Namely, 57 Wolbachia genes were detected in the insect strain jCAus; 40 Wolbachia genes, which comprised a subset of the 57 genes, were present in the strains $k10^{U2}$ and r13^{U1}; and 31 Wolbachia genes, 29 of which constituted a subset of the 40 genes, were found in the strain Rm^{tet} (Supplemental Fig. S3). On the assumption that acquisition of Wolbachia genes is difficult while loss of Wolbachia genes is easy, the nested gene possession pattern suggested the following evolutionary scenario: (1) among the insect strains, jC^{Aus} diverged first; (2) 16 genes were lost in the lineage leading to the strains $k10^{U2}$, $r13^{U1}$, and Rm^{tet}; (3) Rm^{tet} diverged from the lineage of k10^{U2} and r13^{U1}; (4) 11 genes were lost in the lineage of Rm^{tet}; (5) a gene, vdcC, was lost in the lineages of jC^{Aus} and k10^{U2}/r13^{U1} independently, and a gene, *pyrD*, was lost in the lineage of $k10^{U2}$ and r13^{U1} (Fig. 6A).

Phylogenetic relationship between the insect strains

Indeed the parsimonious reconstruction of the evolutionary process of the transferred *Wolbachia* genes was simple and beautiful, but several lines of evidence raised an objection to the scenario. When we constructed phylogenetic relationships between the insect strains, both mitochondrial and nuclear genes indicated that not jC^{Aus} but Rm^{tet} constituted the basal lineage (Fig. 6B,C). Phy-

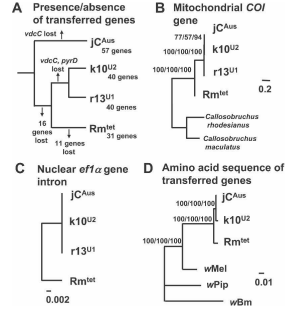


Figure 6. Phylogenetic relationships between the insect strains jC^{Aus} , $k10^{U2}$, $r13^{U1}$, and Rm^{tet} of *Callosobruchus chinensis*. (*A*) Relationship of the insect strains deduced from their possession patterns of the transferred *Wolbachia* genes. (*B*) Relationship inferred from 522 aligned nucleotide sites of mitochondrial cytochrome oxidase I gene. (*C*) Relationship inferred from 590 aligned nucleotide sites of nuclear elongation factor 1 α intron. (*D*) Relationship of the concatenated amino acid sequence of the 27 transferred *Wolbachia* genes based on 2929 aligned amino acid sites. NJ trees are shown, whereas ML and BA trees exhibit the same topologies. Support values >50% are indicated at the nodes.

logenetic analysis of the 27 transferred *Wolbachia* genes also supported the relationship (Fig. 6D). These results consistently refuted the above evolutionary hypothesis Fig. 6A).

Nucleotide sequence polymorphisms in transferred Wolbachia genes

Furthermore, we found an interesting pattern in the 57 transferred *Wolbachia* genes identified in the insect strain jC^{Aus} . When PCR products of these genes were directly sequenced, it turned out that 24 of the 57 genes contained nucleotide sequence polymorphisms. These polymorphisms were also confirmed by cloning and sequencing of the PCR products (Supplemental Table S4). Strikingly, 23 of the 24 genes, except for *pyrD*, were a subset of the 29 genes that were conserved among all the insect strains jC^{Aus} , $k10^{U2}$, $r13^{U1}$, and Rm^{tet} (Supplemental Fig. S3).

Putative duplication of transferred *Wolbachia* genome fragment on the insect chromosome

These patterns suggested that (1) the transferred *Wolbachia* genome is, at least partially, duplicated; (2) the conserved *Wolbachia* genes tend to be duplicated while the other genes are likely to be single-copied; and (3) the gene duplication probably constitutes a reason why some of the transferred genes are conserved whereas others are not. The process and mechanism of the duplication are at present obscure. However, considering that the FISH signal was restricted to the proximal region of the basal short arm of X chromosome (Fig. 2), the duplicated genome fragments are likely to be located in the physical vicinity, possibly constituting tandem repeats. If so, unequal crossing over between synapsing X chromosomes is suspected as a candidate process involved in the gene duplication. To verify this idea, the genomic structure of the transferred *Wolbachia* genes should be determined.

Absence of transferred *Wolbachia* genes in bean beetles allied to *C. chinensis*

In Japanese populations of *C. chinensis*, 97.1% (398/410) insects possessed *wsp* gene of lateral transfer origin (Kondo et al. 2002b). In representatives of the remaining 2.9% (12 individuals)— namely, the strains $k10^{U2}$ and $r13^{U1}$ —we found that the transferred *Wolbachia* genes were actually present, although *wsp* and some other genes were not detected. In Taiwanese populations of *C. chinensis*, even in the absence of *wsp* gene, many of the transferred *Wolbachia* genes were also identified (Fig. 5; Supplemental Fig. S3; Supplemental Table S1). By contrast, in *Callosobruchus maculatus* and *Callosobruchus rhodesianus*, which are congenic relatives of *C. chinensis*, no *Wolbachia* genes were detected (Supplemental Table S1). Hence, it appears likely that the *Wolbachia* genome fragment was transferred to the insect chromosome after speciation of *C. chinensis*.

When did the Wolbachia genome transfer occur?

The *Wolbachia* genome fragment must have been present in the ancestor of the Japanese and Taiwanese strains of *C. chinensis*. The evolutionary distance values between the strains jC^{Aus} and Rm^{tet} were nearly comparable to the values between *D. melanogaster* and *Drosophila simulans*. By applying the evolutionary rates estimated from biogeography and molecular clocks of multiple loci of the fruit flies (Lachaise et al. 1988; Li et al. 1999), we estimated the divergence time of the *C. chinensis* strains as 2.5

MYA and 0.74 MYA on the basis of mitochondrial *COI* gene and nuclear *ef1* α intron gene sequences, respectively (Supplemental Table S5). Hence, although the estimate is quite rough, the *Wolbachia* genome fragment was probably transferred to the insect chromosome about one or a few million years ago in the ancestor of *C. chinensis*.

Possible mechanisms for prevalence of transferred *Wolbachia* genes in host insect populations

All of these results suggested that the Wolbachia genome fragment integrated in the X chromosome of C. chinensis is nearly fixed in the insect populations and has undergone different levels of evolution and mutational degradation in different lineages. The mechanism whereby the prevalence has been established is currently not understood. Fixation through genetic drift, possibly associated with a population bottleneck that occurred in the ancestor of C. chinensis, is a candidate mechanism. In the process of spreading in the insect populations, hitchhiking with coexisting Wolbachia endosymbionts-which are universally found in current C. chinensis populations and cause cytoplasmic incompatibility (Kondo et al. 2002b)-might have played some roles. Alternatively, the Wolbachia genome fragment might behave like Medea, a maternal-effect chromosomal factor known from the flour beetle Tribolium castaneum that is capable of spreading in host populations in a selfish manner (Beeman et al. 1992). Meaningfully, Medea phenotype and Wolbachia-induced cytoplasmic incompatibility can be explained by the same "poison-antidote" or "modification-rescue" mechanism (Poinsot et al. 2003).

Evolution and fate of laterally transferred endosymbiont genes

All of the data obtained in this study consistently indicate that the *Wolbachia* genes transferred to the host insect chromosome tend to be structurally disrupted (Supplemental Fig. S2), eroded and lost (Fig. 5), freed from functional constraints (Fig. 3), and transcriptionally inactive (Fig. 4). These results strongly suggest that most, if not all, of the laterally transferred endosymbiont genes have been pseudogenized and, although currently prevalent in the host insect populations, are destined to be DNA junk in the insect genome.

Similarity to organelle genomes recently transferred to host nucleus

Interestingly, these evolutionary patterns are prevalent among remnant mitochondrial and plastid genomes located on the nuclear genomes of various plants, fungi, and animals, including humans (Bensasson et al. 2001; Timmis et al. 2004). These organelles are evolutionarily of bacterial endosymbiont origins (Margulis 1981), acquired by ancestral eukaryotes very anciently (Hedges et al. 2004); most of their original gene repertoire has been lost in the evolutionary course, leading to drastically reduced organellar genomes (Lang et al. 1999; Simpson and Stern 2002), and some of the genes are transferred to the nuclear genome where such genes are transcribed, translated, and transported to the organelles for functioning (Leister 2003; Brown 2003). Aside from the ancient endosymbiotic gene transfers, a large number of organellar genomic DNA fragments of much more recent origins are present in the nuclear genomes of diverse eukaryotes, indicating that endosymbiotic gene transfer is a ubiquitous and currently ongoing process that pervades nuclear DNA dynamics (Bensasson et al. 2001; Timmis et al. 2004). The transferred organelle DNA can be amplified once inserted (Woischnik and Moraes 2002; Hazkani-Covo et al. 2003), and some of them are organized as tandem repeats (Tourmen et al. 2002). These nuclear copies of organellar genes are totally pseudo-genized, like the *Wolbachia* genes transferred to the X chromosome of *C. chinensis*.

How frequent are lateral gene transfers from *Wolbachia* endosymbionts to their hosts?

Here it should be noted that *Wolbachia* endosymbionts are intimately associated with germ-line cells and are maternally transmitted, like organelles, via ovarial passage through host generations (Veneti et al. 2004). Worldwide PCR surveys of *Wolbachia* infection in diverse insect taxa have consistently revealed high infection frequencies, >20% at the species level (Werren and Windsor 2000). On account of such host–symbiont developmental integrity and ecological ubiquity, the following questions may arise: Why not more cases of *Wolbachia*–host gene transfers? Is the case of *C. chinensis* an orphan exception, or are other cases to be found? How frequent have such lateral gene transfers occurred?

Actually, other cases are emerging from recent genome sequencing projects of insects and nematodes. Genome fragments of *Wolbachia* endosymbionts, whose sizes vary from single-gene level to nearly whole-genome level, have been detected from the fruit fly *Drosophila ananassae*, the parasitoid wasps *Nasonia giraulti*, *Nasonia vitripennis*, and *Nasonia longicornis*, and the filarial nematodes *Onchocerca volvulus*, *Onchocerca ochengi*, *Brugia malayi*, *Brugia timori*, *Brugia pahangi*, and *Dirofilaria immitis* (Fenn et al. 2006; Dunning-Hotopp et al. 2007). These recent findings suggest that such symbiont–host gene transfers might have been occurring more frequently than previously envisioned, like the gene transfers from organelles to nucleus.

Conclusion and perspective

In this study, we comprehensively investigated the magnitude, process and evolution of the lateral genome transfer from *Wolbachia* endosymbiont to *C. chinensis* with reference to the *w*Mel genome information. Currently we are making efforts to completely determine the whole genomic region of *C. chinensis* containing all the transferred *Wolbachia* genome fragments by fosmid/BAC genomic library screening.

Once all the transferred *Wolbachia* genes are identified, we will be able to evaluate the transcriptional activities of all the symbiont-derived genes on the insect chromosome. Certainly it is plausible that all the transferred genes will finally turn out to be molecular fossils of an evolutionary episode in the past, but if some of the genes would be identified as transcriptionally active and functional, such genes would provide novel insights into how host–symbiont gene transfers could contribute to organismal evolution.

The evolutionary origin of the transferred *Wolbachia* genes in natural populations of *C. chinensis* is still an enigma. The current worldwide distribution of the pest of stored beans is probably mediated by human activities (Borowiec 1987). Mitochondrial DNA analysis of Asian populations of *C. chinensis* certainly showed that insect samples collected from bean field/storage/ market exhibited specific genotypes, while insect samples collected from wild leguminous plants were genetically diverse (Tuda et al. 2004). Hence, it is expected that a worldwide survey of wild *C. chinensis* populations not infesting agricultural beans but feeding on wild legumes would lead to key discoveries, such as insect populations without the transferred *Wolbachia* genes or insect populations infected with a relict bacterial *Wolbachia* endosymbiont closely related to the transferred *Wolbachia* genome.

Methods

Materials

Strains of *C. chinensis* and allied bean beetles used in this study are listed in Supplemental Table S3. These insects were maintained on seeds of the adzuki bean *Vigna angularis* at 30°C in a long-day regimen of 16 h light and 8 h dark, except for the strains Rm and Rm^{tet} fed on seeds of the wild legume *Rhynchosia minima*.

DNA extraction

Female adults of *C. chinensis* were subjected to DNA extraction by using the QIAamp DNA Mini kit (Qiagen) for PCR amplification or by a standard proteinase K/SDS/phenol/chloroform extraction procedure for Southern blotting.

PCR detection of Wolbachia genes

Specific primer sets for amplifying each of the *Wolbachia* genes were designed on the basis of published and ongoing genome sequences of three *Wolbachia* strains, *w*Mel (Wu et al. 2004), *w*Bm (Foster et al. 2005), and *w*Pip (Sanger Institute) (Supplemental Table S1). Primer pairs that failed to amplify PCR products of expected size from the positive DNA samples—the *D. melanogaster* strain OGS-1 infected with *w*Mel and the *Wolbachia*-infected *C. chinensis* strain jC—were not used for further analyses. The amplified products were directly sequenced, and the sequences were subjected to BLAST search to confirm their sequence identity to *Wolbachia* genes.

Probe construction for Southern blotting

DNA probes for Southern blotting were synthesized by long-PCR amplification from the *w*Mel-infected *D. melanogaster* DNA as template. The *Wolbachia*-specific gene primers used are listed in Supplemental Table S2. The amplified products were electrophoresed in agarose gels, excised and extracted from the gels, and digoxigenin-labeled by using the DIG Random Primed DNA Labeling Kit (Roche).

Southern blotting

Genomic DNA preparations of the *C. chinensis* strain jC^{Aus} were digested with restriction endonuclease *EcoRI*, *BamHI*, or *HindIII*. The digested DNA samples, 5–10 µg in each lane, were electrophoresed in agarose gels. Genomic DNA of the *w*Mel-infected *D. melanogaster* was used as positive control. The separated DNA fragments were transferred to nylon membranes by a standard capillary blotting procedure, and fixed by UV-cross-linking. Hybridization and detection of each of the probes were performed by using the DIG Detection Kit (Roche) according to the manufacturer's instructions. Specific hybridization of all the probes was confirmed by using the genomic DNA of *w*Mel-infected *D. melanogaster*, wherein the banding patterns of the signals agreed with the restriction patterns inferred from the genomic sequence of *w*Mel deposited in the DNA database.

DNA sequencing

The PCR products were sequenced by direct sequencing and/or sequencing of plasmid-cloned DNA. In the former, the PCR products were treated with Exo-SAP-IT (Amersham) and sequenced by using amplifying primers. In the latter, the PCR products were

cloned into pGEM T-vector (Promega) or pCR Blunt-vector (Invitrogen). The cloned inserts were amplified by single-colony PCR with vector-specific primers and subjected to sequencing with the primers.

FISH

Testes were dissected from adult individuals of C. chinensis in a phosphate-buffered saline, swollen for 30 min in a hypotonic solution (75 mM KCl), and fixed, crushed, and suspended in methanol-acetic acid (3:1) by repeated pipetting; the suspended cells were then spread and dried on clean glass slides. Biotinlabeled DNA probes were prepared by using the Biotin Nick Translation Mix (Roche). Two sets of labeled probes were synthesized. The short-probes set was produced by labeling short PCR products (180-2500 bp) of the 36 transferred Wolbachia genes (Supplemental Table S1). The long-probes set consisted of four long PCR products (3.5, 4.0, 5.5, and 8.8 kb in size) amplified by the primer pairs as described in Supplemental Table S6. The labeled probes were dissolved in a hybridization buffer (50% formamide, 10% dextran sulfate, 2× SSC). Chromosomal DNA preparations were denatured at 76°C for 2 min in a denaturing buffer (70% formamide, $2 \times$ SSC), and were incubated overnight with the probe-containing hybridization buffer. After being thoroughly washed, the hybridized probes on the chromosomes were detected by using AlexaFluor488-labeled streptavidin (Invitrogen), and the signals were amplified by using antistreptavidin antibody biotin conjugate (Vector). The whole chromosomes were counterstained with 4,6-diamino-2-phenylidole, and the hybridization signals were observed under a fluorescent microscope (BX-50, Olympus) and recorded by a chilled CCD camera (Sensys 1400, Photometrics). Pseudocolor images were generated by IPLab (Scanalytics).

RT-PCR

Total-RNA samples were individually prepared from adult females of *C. chinensis* by using the RNA Mini Kit (QIAGEN). The RNA samples were treated with DNase, reverse-transcribed with SuperScriptIII (Invitrogen), and subjected to PCR using the primers listed in Supplemental Table S1. All of the RNA samples were tested for genomic DNA contamination by performing PCR without reverse transcription.

Quantitative RT-PCR

The reverse-transcribed RNA samples were subjected to real-time fluorescent quantitative RT-PCR by using ABI Prism 7300 Sequence detection system (Applied Biosystems), with SYBR Green I (Molecular Probes) as detection dye and ROX (Invitrogen) as reference dye, as described elsewhere (Narita et al. 2007). The PCR primers are listed in Supplemental Table S6.

Molecular phylogenetic and evolutionary analyses

Phylogenetic analyses were conducted by the neighbor-joining (NJ) method, the maximum-likelihood (ML) method, and Bayesian analysis (BA). NJ trees were constructed by using the program package ClustalW (Thompson et al. 1994) or MEGA3.1 (Kumar et al. 2004). Bootstrap probabilities were calculated by generating 1000 bootstrap replicates. ML trees were estimated by using the program package TREE-PUZZLE 5.2 (Schmidt et al. 2002). In BA analysis, we used the program package MrBayes 3.1.2 (Ronquist and Huelsenbeck 2003). The numbers of synonymous and nonsynonymous substitutions per site were calculated as described (Miyata and Yasunaga 1980).

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