**Rickettsia Symbiont in the Pea Aphid Acyrthosiphon pisum: Novel Cellular Tropism, Effect on Host Fitness, and Interaction with the Essential Symbiont Buchnera**

Makiko Sakurai, Ryuichi Koga, Tsutomu Tsuchida, Xian-Ying Meng, and Takema Fukatsu*

Institute for Biological Resources and Functions, National Institute of Advanced Industrial Science and Technology (AIST), Tsukuba 305-8566, Japan

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In natural populations of the pea aphid *Acyrthosiphon pisum*, a facultative bacterial symbiont of the genus *Rickettsia* has been detected at considerable infection frequencies worldwide. We investigated the effects of the *Rickettsia* symbiont on the host aphid and also on the coexisting essential symbiont *Buchnera*. In situ hybridization revealed that the *Rickettsia* symbiont was specifically localized in two types of host cells specialized for endosymbiosis: secondary mycetocytes and sheath cells. Electron microscopy identified bacterial rods, about 2 μm long and 0.5 μm thick, in sheath cells of *Rickettsia*-infected aphids. Virus-like particles were sometimes observed in association with the bacterial cells. By an antibiotic treatment, we generated *Rickettsia*-infected and *Rickettsia*-eliminated aphid strains with an identical genetic background. Comparison of these strains revealed that *Rickettsia* infection negatively affected some components of the host fitness. Quantitative PCR analysis of the bacterial population dynamics identified a remarkable interaction between the coexisting symbionts: *Buchnera* population was significantly suppressed in the presence of *Rickettsia*, particularly at the young adult stage, when the aphid most actively reproduces. On the basis of these results, we discussed the possible mechanisms that enable the prevalence of *Rickettsia* infection in natural host populations in spite of the negative fitness effects observed in the laboratory.

The genus *Rickettsia* is a phylogenetically distinct bacterial group that belongs to the order *Rickettsiales* in the α-subdivision of the class *Proteobacteria* (33, 37). All members of the genus are gram-negative bacteria in obligate association with eukaryotic cells (9, 44). This bacterial group includes notorious human and animal pathogens vectored by blood-sucking arthropods: *R. prowazekii*, vectored by lice and causing epidemic typhus; *R. typhi*, vectored by fleas and causing murine typhus; and *R. rickettsii* and allied species, vectored by ticks and causing spotted fever, etc. (9).

Although some *Rickettsia* species occasionally infect vertebrates with pathology, and a few species are obligatorily associated with vertebrates, it is believed that the primary hosts of most *Rickettsia* species are arthropods and other invertebrates (9, 21, 44). In these cases, the *Rickettsia* bacterium is inherited by ovaerial transmission through host generations with a high fidelity, is detected in host populations at a considerable frequency, and can be regarded as a facultative endosymbiotic associate (9, 44). Thus far, such *Rickettsia* symbionts have been identified from a wide variety of invertebrates such as ladybird beetles (19, 43, 46), an aphid (6), a leafhopper (10), a bruchid beetle (13), ticks (2, 30), leeches (21), and others.

Many obligate insect endosymbionts, like *Buchnera* in aphids and *Wigglesworthia* in tsetse flies, are harbored by specialized cells for endosymbiosis called mycetocytes (or bacteriocytes) (1, 3, 4). Some facultative insect endosymbionts, such as γ-proteobacterial secondary symbionts in aphids, are housed in different types of specialized cells called secondary mycetocytes and sheath cells (4, 14, 16, 22, 40). On the other hand, no such specific cellular localization has been reported for facultative symbionts that are widely found across diverse insect taxa, such as *Wolbachia*, *Spiroplasma*, and *Rickettsia* (8, 11, 15, 20, 21, 23), although concentrated distributions in ovary and germ plasm were recognized in some cases (8, 18, 20).

In addition to the essential symbiont *Buchnera*, five facultative symbionts have been identified from natural populations of the pea aphid *Acyrthosiphon pisum*: PASS (pea aphid secondary symbiont or R-type symbiont), PAUS (pea aphid U-type symbiont or U-type symbiont), PABS (pea aphid *Bemisia*-type symbiont or T-type symbiont), *Rickettsia* symbiont (or PAR, after pea aphid *Rickettsia*), and *Spiroplasma* symbiont (35, 36, 39). In the host body, the three γ-proteobacterial symbionts PASS, PAUS, and PABS were reported to exhibit a remarkable cellular localization; they were detected in secondary mycetocytes and sheath cells and also in hemolymph (16, 22, 35, 40). By contrast, no such cellular tropism has been reported for the *Rickettsia* symbiont of *A. pisum*. Previous studies indicated that the *Rickettsia* symbiont is present in hemolymph (6) and confers negative fitness effects on the host aphid, while the intensity of the effects was dependent on environmental factors such as temperature and host plant species (5, 7, 28).

In this study, the *Rickettsia* symbiont of *A. pisum* was analyzed in detail by using molecular phylogenetic analysis, in situ hybridization, light and electron microscopy, a selective elimination technique, and quantitative PCR, which revealed previously unknown biological aspects of the *Rickettsia* symbiont, including highly specific cellular localization and interaction with the essential symbiont *Buchnera*.

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* Corresponding author. Mailing address: Institute for Biological Resources and Functions, National Institute of Advanced Industrial Science and Technology (AIST), Tsukuba 305-8566, Japan. Phone: 81-29-861-6087. Fax: 81-29-861-6080. E-mail: t-fukatsu@aist.go.jp.
MATERIALS AND METHODS

Materials. The following isofemale strains of A. pismum were used in this study. The *Rickettsia*-infected strain OY was collected at Tochigi, Japan, in 2000 and has been maintained in the laboratory with stable *Rickettsia* infection (39). The strain AIST, which is free of facultative symbionts, was collected at Tsukuba, Japan, in 1999 (16). Aphids were reared on seedlings of the broad bean *Vicia faba* at 20°C in a long-day regimen (16 h of light and 8 h of dark).

DNA extraction, cloning, and sequencing. Aphids of the strain OY were subjected to DNA extraction by using a QIAamp mini kit (QIAGEN). Genes of the 16S rRNA and *gltA* transcripts were PCR-amplified by using primers 16S1 (5′-AGATGGTTGATCTGTCG-3′) and 16SB1 (5′-TACGGYTACCTTGTTACGACT-3′) (16) and primers CS1 (5′-GAGCTTGCTCTTTGTCGGCAA-3′) and BuchGroEL-1967R (5′-AGCTCAAATGGTAAAAGAAGTTGCGTTTCAGATCCATTGGAT-3′) for the 16S rRNA gene and the *gltA* gene, respectively. The amplified PCR products were cloned and sequenced as described previously (16). More than three clones for each of the genes were sequenced to eliminate possible PCR errors.

Molecular phylogenetic analysis. Multiple alignments of nucleotide sequences were generated by using the program package MEGA, version 2.1 (26). The sequences were inspected and corrected manually. Neighbor-joining trees, with bootstrap resamplings, were constructed by using the program package Clustal W (38). The final alignments were used in a molecular phylogenetic analysis, with the program package MEGA, version 2.1 (26). The nucleotide sequences of the 16S rRNA gene and the *gltA* gene of the *Rickettsia* symbiont from *A. pismum* described previously (4) were used in the molecular phylogenetic analysis of the *Rickettsia* symbiont from *A. pismum* described previously (4).

RESULTS

Detection and characterization of *Rickettsia* symbiont in the aphid strain OY. The aphid strain OY has been maintained in the laboratory through 80 to 90 parthenogenetic generations since being collected in 2000. Periodical PCR checks always indicated that all individuals examined harbored *Rickettsia* symbiont in addition to the essential symbiont *Buchnera*. Other facultative symbionts, such as PASS, PAUS, PABS, and *Spiroplasma*, were not detected from the aphid strain (data not shown).

16S rRNA gene and *gltA* gene of *Rickettsia* symbiont. A 1.5-kb segment of the eubacterial 16S rRNA gene was amplified by PCR from the DNA of the strain OY and was subjected to cloning and restriction fragment length polymorphism typing. Two restriction fragment length polymorphism types were identified among 25 clones examined, of which four clones from each of the types were sequenced. The sequences of one type, 1,470 bp in length, were identical to each other and exhibited 100% similarity to the sequences of *Buchnera* sp. from *A. pismum* (DDBJ accession number AB033774). The sequences of the other type, 1,422 bp in length, were identical to each other and exhibited 99.8%, 98.9%, and 99.4% similarity to the sequence of a *Rickettsia* sp. from *A. pismum* collected in California (accession number U42084), the sequence of a *Rickettsia* sp. from papaya leafhopper *Empoasca papayae* (accession number U76990), and the sequence of *Rickettsia bellii* (accession number U11014), respectively.

A 0.38-kb segment of the *rickettsia* *gltA* gene was also amplified by PCR from the DNA of the strain OY and was subjected to cloning and sequencing. The nucleotide sequences of six clones determined were completely identical (382 bp) and exhibited 95.5% and 91.9% similarity to the sequence of a *Rickettsia* sp. from papaya leafhopper *Empoasca papaya* (accession number U76990) and the sequence of *R. bellii* (accession number AY375161), respectively.

Molecular phylogenetic analysis of *Rickettsia* symbiont. Molecular phylogenetic analysis on the basis of 16S rRNA gene sequences from *Rickettsia* spp. revealed that our *Rickettsia* se-
sequence belonged to neither the “spotted fever group” nor the “typhus group” but was placed within the clade consisting of *R. bellii* and *Rickettsia* spp. from aphid and leafhopper (Fig. 1A). Molecular phylogenetic analysis on the basis of *gltA* gene sequences gave substantially the same result: our *Rickettsia* sequence was placed in the clade containing *R. bellii* and the *Rickettsia* sp. from leafhopper (Fig. 1B).

**In situ hybridization of Rickettsia symbiont.** Whole-mount fluorescence in situ hybridization revealed a remarkable cellular tropism of the *Rickettsia* symbiont (Fig. 2). In aphid embryos of the strain OY, strong signals of *Rickettsia*-specific probe were found in two types of cells: secondary mycetocytes (Fig. 2, arrows) and sheath cells (Fig. 2, arrowheads). The secondary mycetocytes were relatively large and intercalated between the primary mycetocytes harboring *Buchnera* (Fig. 2A and B). The sheath cells were small and flat, located on the periphery of the primary mycetocytes (Fig. 2A, C, and D). The primary mycetocytes, secondary mycetocytes, and sheath cells constituted a large symbiotic organ, mycetome, in the body cavity of the aphids. No-probe control, RNase digestion control, and competitive suppression control confirmed the specificity of the detected signals (data not shown). In situ hybridization of the *Rickettsia*-free aphid strain AIST detected signals of *Buchnera* but not those of *Rickettsia* (data not shown).

**Electron microscopy of Rickettsia symbiont.** Electron microscopy of embryos of the strain OY identified a dense population of rod-shaped bacteria in the cytoplasm of the sheath cells (Fig. 3A). The bacterial cells were slender rods, 1.84 ± 0.43 μm long and 0.48 ± 0.05 μm thick (mean ± standard deviation [n = 20]) and exhibited cell wall structure (Fig. 3B). The rod-shaped bacteria were not observed in the *Rickettsia*-free aphid strain AIST (data not shown). Virus-like particles were

![FIG. 1. Molecular phylogenetic analysis of the *Rickettsia* symbiont from *A. pisum* strain OY. (A) Neighbor-joining tree on the basis of 16S rRNA gene sequences. A total of 1,417 unambiguously aligned nucleotide sites were subjected to the analysis. (B) Neighbor-joining tree on the basis of *gltA* gene sequences. A total of 353 unambiguously aligned nucleotide sites were subjected to the analysis. The percent bootstrap values are shown at the nodes, although values less than 70% are not indicated. The numbers in brackets are accession numbers. Three major groups in the genus *Rickettsia*, “spotted fever group,” “typhus group,” and “bellii group,” are indicated on the right side.

![FIG. 2. Whole-mount in situ hybridization of aphid embryos targeting *Rickettsia* (red) and *Buchnera* (green). (A) Mycetomes of an aphid embryo in which secondary mycetocytes and sheath cells harboring *Rickettsia* are seen in addition to a number of primary mycetocytes harboring *Buchnera*. (B to D) Magnified images of mycetome. Arrows, secondary mycetocytes; arrowheads, sheath cells; bars, 50 μm.](http://aem.asm.org/)

sometimes observed in association with the bacterial cells (Fig. 3C).

Selective elimination of *Rickettsia* symbiont by antibiotic treatment. Fifteen adult aphids of the *Rickettsia*-infected strain OY were injected with ampicillin solution. In the G1 offspring, both *Buchnera* and *Rickettsia* were detected by diagnostic PCR. In the G2 offspring, *Rickettsia* infection disappeared in all the lines, while *Buchnera* infection was maintained. After maintenance for 10 generations and on, *Rickettsia* infection did not recover in the treated lines (data not shown), whereby we established a *Rickettsia*-eliminated aphid strain, OYamp. A control aphid strain, OYdw, was generated in the same way, but distilled water was injected instead of the antibiotic.

**Fitness effects of *Rickettsia* symbiont.** Fitness parameters were compared between the *Rickettsia*-infected aphid strain OYdw and the *Rickettsia*-eliminated aphid strain OYamp with identical genetic backgrounds. The *Rickettsia*-infected aphids exhibited lower fresh body weight and total number of offspring than the *Rickettsia*-eliminated aphids, whereas there were no significant differences with regard to age of first reproduction and longevity between the aphid strains (Table 1).

**Population dynamics of *Rickettsia* and *Buchnera* in the developmental course of the host aphid.** By using a quantitative PCR technique, we investigated the population dynamics of the symbiotic bacteria in the *Rickettsia*-infected aphids and the *Rickettsia*-eliminated aphids with identical genetic backgrounds. The population of *Rickettsia* increased during nymphal growth, reaching a plateau in 10-day-old adults, and the infection level was maintained in older insects (Fig. 4A). When normalized by the titers of a host gene, the density of *Rickettsia* exhibited similar dynamics but a transient drop around the 5-day-old stage and a resurgence at the 15-day-old and later stages (Fig. 4B).

The population dynamics of *Buchnera* were different between the *Rickettsia*-infected aphids and the *Rickettsia*-eliminated aphids. In the absence of *Rickettsia*, the population of *Buchnera* increased during nymphal growth, reaching a peak in actively reproducing young adults, and declined in older insects. Notably, the titers of *Buchnera* were significantly lower in the presence of *Rickettsia* (at 50 to 60% of those in the absence of *Rickettsia*) during the young adult period, when the aphids most actively reproduce. At later stages, in contrast, the titers of *Buchnera* became lower in the absence of *Rickettsia* (Fig. 4C). Even when normalized by the titers of a host gene, the density of *Buchnera* exhibited qualitatively similar dynamics: significantly higher densities in the absence of *Rickettsia* during the young adult period but lower densities at later stages (Fig. 4D).

**TABLE 1. Effect of *Rickettsia* infection on fitness parameters of the host aphid**

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>OY_{dw} (+Rickettsia)</th>
<th>OY_{amp} (−Rickettsia)</th>
</tr>
</thead>
<tbody>
<tr>
<td>No. of aphids measured</td>
<td>17</td>
<td>20</td>
</tr>
<tr>
<td>Body wt* (mg) at 12 days old</td>
<td>3.78 ± 0.48</td>
<td>4.38 ± 0.40</td>
</tr>
<tr>
<td>Total no. of offspring*</td>
<td>113 ± 19.5</td>
<td>134 ± 12.3</td>
</tr>
<tr>
<td>Age at first reproduction (days after birth)</td>
<td>8.35 ± 0.49</td>
<td>8.41 ± 0.50</td>
</tr>
<tr>
<td>Longevity (days)</td>
<td>28 ± 4.0</td>
<td>28 ± 3.8</td>
</tr>
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*Means ± standard deviations are shown. Asterisks indicate the fitness parameters that exhibited statistically significant differences (Mann-Whitney U test, \(P < 0.001\)). The measurements were conducted 15 generations after the injection treatments.*

In the G2 offspring, *Rickettsia* infection disappeared in all the lines, while *Buchnera* infection was maintained. After maintenance for 10 generations and on, *Rickettsia* infection did not recover in the treated lines (data not shown), whereby we established a *Rickettsia*-eliminated aphid strain, OYamp. A control aphid strain, OYdw, was generated in the same way, but distilled water was injected instead of the antibiotic.

**FIG. 3.** Electron microscopy of the *Rickettsia* symbiont. (A) A sheath cell harboring rod-shaped cells of the *Rickettsia* symbiont (arrows) surrounded by primary mycetocytes harboring *Buchnera* (asterisks). (B) A magnified image of the *Rickettsia* cells in a sheath cell. (C) Virus-like particles (arrowheads) in association with the *Rickettsia* cells. Abbreviations: ShC, sheath cell; PM, primary mycetocyte.
DISCUSSION

Members of the genus *Rickettsia* are found from a wide variety of arthropods and are obligatorily associated with the host cells (9, 44). However, no specific cells for harboring *Rickettsia* have been reported, although concentrated infections in ovaries for vertical transmission and/or in salivary glands for horizontal transmission might be presumed implicitly (9, 30). Previous studies suggested that the *Rickettsia* biont of *A. pisum* is present in the hemolymph, on the grounds that hemolymph transfer from infected aphids to uninfected ones established *Rickettsia* infection in the recipients (5), and rod-shaped bacteria were observed in the hemolymph of *Rickettsia*-infected aphids (6). In addition to the hemolymph infection, we identified the localization of the *Rickettsia* symbiont in two types of cells specialized for endosymbiosis: secondary mycetocytes and sheath cells (Fig. 2). To our knowledge, this study provides the first report of specific host cells that harbor *Rickettsia*. The cellular tropism suggests that elaborate molecular and cellular interactions are involved in the infection and maintenance of the *Rickettsia* symbiont in the host aphid.

The in vivo localization of the *Rickettsia* symbiont in secondary mycetocytes, sheath cells, and hemolymph was quite similar to the localization of other secondary symbionts of *A. pisum* such as PASS, PAUS, and PABS (16, 22, 35, 40). It should be noted that these facultative symbionts belong to phylogenetically distinct bacterial lineages: the *Rickettsia* symbiont is a member of the α-Proteobacteria, whereas PASS, PAUS, and PABS constitute distinct clades in the γ-Proteobacteria (34, 35). The similar localization patterns of PASS, PAUS, PABS, and the *Rickettsia* raise the possibility that although speculative, these facultative symbionts might share some common molecular and cellular mechanisms for their infection and maintenance in the host aphid.

To evaluate the biological effects of the *Rickettsia* symbiont, it was pivotal to generate aphid strains that are genetically identical and differ only in *Rickettsia* infection. Recently, a novel selective elimination technique was developed for the γ-proteobacterial facultative symbionts PASS and PAUS: microinjection of ampicillin, an antibiotic that inhibits bacterial cell wall synthesis, resulted in complete elimination of these symbionts in the offspring of injected aphids, while the coexisting essential symbiont *Buchnera* was not affected (22, 41). The technique was also effective for selective elimination of the α-proteobacterial *Rickettsia* symbiont, suggesting potential applicability of the technique to a wider variety of bacterial symbionts. Previous therapeutic studies showed that penicillin and allied antibiotics, including ampicillin, are generally not effective in *Rickettsia*-infected patients (9, 44). However, in vitro studies demonstrated that penicillin is potentially able to eradicate *Rickettsia*, suggesting that inefficient permeation of the antibiotic to the intracellular location of the bacterium may be responsible for the apparent in vivo resistance (47, 48). It should be noted that *Buchnera* is always intracellular, whereas PASS, PAUS, and *Rickettsia* have an extracellular phase in the hemolymph (6, 22, 40). The susceptibility of the facultative symbionts to the antibiotic could be understood in this context.

By using a quantitative PCR technique, we monitored the dynamics of population and density of *Buchnera* and *Rickettsia* during the host development (Fig. 4), which provided some insights into the nature of the symbiotic associations. The pro-

FIG. 4. Infection dynamics of *Rickettsia* and *Buchnera* in the developmental course of the host aphid. (A) *Rickettsia* titers in terms of *gldA* gene copies per insect. (B) *Rickettsia* densities in terms of *gldA* gene copies per *ef1α* gene copy. (C) *Buchnera* titers in terms of *groEL* gene copies per insect. (D) *Buchnera* densities in terms of *groEL* gene copies per *ef1α* gene copy. Open circles, the *Rickettsia*-infected aphid strain OY<sup>+</sup>; closed squares, the *Rickettsia*-eliminated aphid strain OY<sup>−</sup>. Means and standard deviations are shown (n = 12, respectively). Asterisks indicate statistically significant differences (Mann-Whitney U test; *, P < 0.05; **, P < 0.01; ***, P < 0.001). The measurements were conducted at 18 to 20 generations after the injection treatments.
liferation of Buchnera was coincident with the reproductively active stage of the host aphid (Fig. 4C and D), which probably reflects the principal biological role of the symbiont: the provision of essential amino acids and other nutrients to support the rapid development of a large number of embryos (12). The proliferation pattern also suggests that the host aphid may have a control mechanism over the proliferation of Buchnera, which must have evolved during the long history of the host-symbiont coevolution (29). By contrast, the population of Rickettsia exhibited simple growth-and-plateau dynamics (Fig. 4A). The density of Rickettsia exhibited similar dynamics but with a transient drop around the 5-day-old stage and a resurgence in old insects (Fig. 4B). The transient drop probably reflects the condition that the proliferation of Rickettsia cannot catch up with the rapid growth of young aphids. The resurgence might be related to the normalization by the titers of a host gene: young aphids produce embryos very actively, but the activity drastically declines in old insects, which results in reduced titers of the host gene and consequently overestimated densities of the symbiont. These patterns are suggestive of less strict control over the proliferation of Rickettsia in the host aphid, which may be relevant to the facultative nature of the symbiont.

The secondary mycetocytes and sheath cells harboring Rickettsia are in close association with the primary mycetocytes harboring Buchnera, constituting a symbiotic organ called the mycetome (Fig. 2). The spatial proximity suggests that various biological interactions may take place between the facultative and essential symbionts. The dynamics of Buchnera in the presence and absence of Rickettsia revealed an aspect of such symbiont-symbiont interactions: coinfection with Rickettsia significantly suppressed the population density of Buchnera (Fig. 4C and D). In the endosymbiotic ecosystem of the host body, the facultative symbiont Rickettsia negatively affected the essential symbiont Buchnera, probably via competition for limited resources, energy, and/or an endocellular niche. It was reported that the presence of PASS similarly suppressed the population of Buchnera (22).

In previous studies, fitness effects of Rickettsia infection were investigated by using Rickettsia-infected and uninfected aphids with identical genetic backgrounds that were generated by hemolymph transfer. These studies indicated that Rickettsia infection generally showed negative effects on the host aphids, while the intensity of the effects was dependent on environmental factors such as temperature and host plant species (5, 7, 28). In this study, by using the antibiotic-treated aphids, we identified reduced body weight and fecundity of the Rickettsia-infected host aphids (Table 1), reinforcing the notion that Rickettsia infection generally has negative effects on A. pisum, at least under the laboratory rearing conditions.

In this study, we demonstrated that Rickettsia infection negatively affects both the population density of Buchnera (Fig. 4C and D) and the host fitness (Table 1). The following hypotheses are conceivable to account for the action of the Rickettsia symbiont: (i) Rickettsia negatively affects the host aphid, and as a result, the population density of Buchnera is indirectly suppressed; and (ii) Rickettsia suppresses the essential symbiont Buchnera, and this leads to reduced performance of the host aphid. Of course, these hypotheses are not necessarily mutually exclusive and should be tested in future studies. Even the possibility that a third party is involved cannot be ruled out; the ampicillin treatment might have suppressed unidentified microbial associates of the aphid and thus led to the improved fitness of the host aphid and the increased population density of Buchnera.

Electron microscopy detected the occurrence of virus-like particles associated with the Rickettsia cells (Fig. 2C). At present, without molecular data, the nature of the particles is obscure. Even the possibility that the particle is merely an artifact cannot be excluded. It should be noted, however, that a bacteriophage of the Podoviridae, referred to as APSE-1, has been identified from A. pisum and other aphids, whose occurrence was associated with PABS and allied secondary symbionts (35, 42). Although speculative, such elements might be involved in the negative effects of the facultative symbionts on Buchnera and/or host aphids. These elements might also contribute to evolutionary processes such as symbiont-symbiont and host-symbiont horizontal gene transfer and genetic recombination (25, 45).

Rickettsia infection has been detected at considerable frequencies in natural populations of A. pisum: 48% (59/122) in California (6); 22% (17/77) in another survey in California (27); 4% (35/858) in Japan (39); 30 to 37% on pea and 0 to 10% on alfalfa in France (36), etc. The prevalence in natural populations appears contradictory to the negative fitness effects of Rickettsia infection observed in the laboratory. At present, we do not understand what mechanisms maintain the Rickettsia symbiont in the aphid populations. In a simple model, the infection frequency of an endosymbiont in a host population is determined mainly by three parameters: (i) fidelity of vertical transmission, (ii) fitness effect on the host, and (iii) frequency of horizontal transmission (16, 17). Therefore, for example, sufficiently frequent horizontal transfers between aphid individuals may be able to counter the negative fitness effects for maintaining Rickettsia infection in host populations. More plausibly, the negative fitness effects of Rickettsia infection may be cancelled out by positive fitness effects on some aspects of the physiology and ecology of the host aphid. Previous studies have demonstrated that other facultative symbionts of A. pisum conferred context-dependent positive fitness effects on the host aphid. PASS infection conferred tolerance of the aphid to high temperature (28). PABS infection induced resistance of the aphid against parasitoid wasps (31). PAUS infection improved fecundity of the aphid on an unsuitable host plant (41). PASS infection could support reproduction of the aphid deprived of the essential symbiont Buchnera (22). In this context, it is of great interest to investigate the effects of Rickettsia infection on various life history traits of A. pisum under different environmental conditions. In Japan, the Rickettsia symbiont tended to be detected in southwestern regions (39). In France, the Rickettsia symbiont was frequently found on pea but not on red clover (36). These observations might be relevant to the biological significance of the facultative symbiont for the host aphid.

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