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Phylogenetic Comparison of Endosymbionts with Their Host Ants Based on Molecular Evidence

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ABSTRACT—Some Formicid ants have symbiotic intracellular bacteria in the epithelial cells of their midgut. These endosymbionts are believed to be derived from a common ancestor. A recent study revealed that endosymbionts of the ant genus *Camponotus* are closely related to Enterobacteriaceae, but their relationship to endosymbionts of other genera of ants is unknown. In this study, the nucleotide sequences of 16S ribosomal RNA (rRNA) of endosymbionts and mitochondrial cytochrome oxidase subunit I (COI) of their host were determined in five genera of the subfamily Formicinae (Hymenoptera: Formicidae). Based on these molecular data, we constructed phylogenetic trees in order to characterize the systematic position of the symbionts were all connected with the Enterobacteriaceae but did not constitute a monophylitic group, while the three genera belonging to the tribe Camponotini, the endosymbionts and their hosts made a clade. The topologies of these trees were identical for the most part. These results suggest that the endosymbionts of ants have plural origins, and that in the Camponotini, ancestral symbionts have coevolved with their host ants, which are so divergent to several genera as to construct one tribe.

INTRODUCTION

Many insects, across a wide taxonomic spectrum, contain non-parasitic intracellular microorganisms (Buchner, 1965). There are two kinds of such microorganisms. "Guest microorganisms," most of which have no distinguishable effects on the survival, growth, and fecundity of the hosts, are present in various types of host cells. Guest microorganisms have been recorded in the majority of insect orders. The other one, "Mycetocyte endosymbionts" are restricted to host insect cells characterized by particular morphology and location, termed "mycetocytes." Mycetocyte endosymbionts are usually obviously advantageous to their host (Douglas, 1989). They are mainly observed on seven orders of insects: widespread in Blattaria, Hemiptera, Anoplura and Coleoptera, and a few host taxa from Mallophaga, Diptera, and Hymenoptera.

In the Hymenoptera, one of the taxan carrying mycetocyte endosymbionts is ants, the family Formicidae. The endosymbionts of ants were first reported in *Camponotus ligniperda*

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and *Formica fusca* (Blochmann, 1892). Thus far, several genera in the subfamily Formicinae are known as hosts: eight genera (including *Camponotus*) in the tribe Camponotini, the genus *Formica* in the tribe Formicini, and the genus *Plagiolepis* in the tribe Plagiolepidini (Buchner, 1965; Jungen, 1968; Dasch *et al.*, 1984). The tribe Camponotini is comprised of 14 genera, and mycetocyte symbiosis seems to be common among them (Dasch *et al.*, 1984). In *Formica*, the presence of symbionts varies with the species, the colony and the individual (Jungen, 1968), whereas the situation for *Plagiolepis* is unclear. Endosymbionts are observed in the mycetocytes in the midgut of workers, queens, males, larvae and pupae. Ovaries of workers and queens also contain symbiotic bacteria. The symbionts are vertically transmitted from the ovary to the eggs of the next generation (Lilienstein, 1932; Buchner, 1965).

Some morphological differences are known between the mycetocyte symbiosis of the Camponotini and two other host genera. In the Camponotini, mycetocytes of the midgut are localized between epithelial cells and rest on the basal membrane, filled with Gram-negative and rod-shaped symbionts arranged in an orderly fashion (Lilienstein, 1932; Buchner, 1965). In the adult midgut, they are straight rods, 1 μ m wide and 5–15 μ m long (Kolb, 1959). The symbionts lie freely in

the cytoplasm (Dasch *et al.*, 1984), like that of the tsetse fly *Glossina* (Reinhardet *et al.*, 1972) and the weevil *Sitophilus* (Nardon and Grenier, 1988), whereas in most insects the symbionts are individually enclosed by a membrane originating from the host cell. In *Formica* and *Plagiolepis*, mycetocytes are not in the midgut epithelium but arranged in an unicellular layer behind the midgut (Lilienstein, 1932; Buchner, 1965). The symbionts are Gram-negative, rod-shaped, and free in the cytoplasm like those of Camponotini; however, the cell length is shorter, ranging from 3 to 4 μ m.

The mycetocyte endosymbionts of insects are generally unculturable in vitro (Baumann and Moran, 1997) and are not amenable to traditional procedures for the taxonomy of microorganisms. Thus we have little information about the origin of the symbioses and the systematic relationships of the symbionts with other microorganisms. However, the development of polymerase chain reaction (PCR)-based techniques has provide a useful procedure for clarifying the phylogenetics of unculturable microorganisms. The nucleotide sequence data of the 16S rRNA gene provide the most useful information for the phylogeny reconstruction and systematic characterization (Carl, 1987). A number of insect endosymbionts have been the subject of molecular phylogenetic analyses and systematic characterization, such as the pea aphid (Unterman et al., 1989), whitefly (Clark et al., 1992) and tsetse fly (Beard et al., 1993, Askoy et al., 1995).

In ants, the systematic relationships of symbionts of the genus *Camponotus* were characterized based on the analy-

sis of the 16S rRNA gene, revealing that endosymbionts of *Camponotus* belong to the gamma-subdivision of the Proteobacteria and are placed contiguous to the Enterobacteriaceae (Schröder *et al.*, 1996). However, endosymbionts of the other taxa in the Formicinae were not included in that analysis. The origin of mycetocyte symbiosis of ants and the phylogeny among symbionts in each host taxon are thus still unclear. In this study, the nucleotide sequences of the endosymbionts' 16S rRNA gene were examined in 15 species belonging to *Camponotus, Colobopsis, Polyrhachys, Formica,* and *Plagiolepis.* A molecular phylogenetic analysis was carried out in order to characterize the systematic position of the symbionts among known prokaryotic microorganisms and to clarify the phylogeny among them.

MATERIALS AND METHODS

Insect species

The ant species investigated in this study are summarized in Table 1. The genus *Colobopsis* is often regarded as a subgenus of *Camponotus*. In this paper, it is treated as a separate genus. *Oecophylla smaragdina* was used as the outgroup in the phylogenetic analysis of the host ants. Based on a phylogenetical analysis of COI sequences, this genus is the most basal in the phylogenetic tree which contains 22 ant genera (Hasegawa, unpublished).

For Camponotus japonicus, Colobopsis nipponicus, Formica fusca and Plagiolepis pigmaea, the sample locations were not identical. A sample from a location (H) was used for the sequencing of the COI gene of a host and (S) for the 16S rRNA gene of a symbiont. In ants, as compared with interspecific variation, the intraspecific varia-

 Table 1.
 List of ants investigated in this study.

 H: sampled for sequencing of host COI, S: for symbiont 16S rRNA

Species		Sampling sites	GenBank accession numbers	
	Opecies	Camping sites	host COI	symbiont 16S rRNA
1.	Camponotus japonicus	Hachioji, Tokyo Met. Japan (H)	AB019411	AB018670
		Meguro, Tokyo Met. Japan (S)		
2.	C. kiusiuensis	Meguro, Tokyo Met. Japan	AB019412	AB018671
3.	C. nawai	Okinawa island, Japan	AB019413	AB018672
4.	C. quadrinotatus	Meguro, Tokyo Met. Japan	AB019414	AB018673
5.	C. vagus	Firenze, Italy	AB019415	AB018674
6.	C. vitiosus	Meguro, Tokyo Met. Japan	AB019416	AB018675
7.	Colobopsis nipponicus	Kamogawa, Chiba Pref. Japan (H)	AB019417	AB018676
		Meguro, Tokyo Met. Japan (S)		
8.	Colobopsis sp.	Okinawa island, Japan	AB019418	AB018677
9.	Formica fusca	Puszcza, Poland (H)	AB010925	-
		Firenze, Italy (S)		
10.	F. lemani	Gotenba, Shizuoka Pref. Japan	AB019425	AB018684
11.	F. yessensis	Ishikari, Hokkaido Pref. Japan	AB010928	-
12.	Plagiolepis manczshurica	Inchon, Korea	AB019423	AB018682
13.	PI. pigmaea	Sierra de Huetor, Spain (H)	AB019414	AB018683
		Firenze, Italy (S)		
14.	Polyrhachis dives	Okinawa island, Japan	AB019419	AB018678
15.	P. hippomanes	Okinawa island, Japan	AB019420	AB018679
16.	P. lamellidens	Machida, Tokyo Met. Japan	AB019421	AB018680
17.	P. ypsilon	Bako, Sabah, Malaysia	AB019422	AB018681
18.	Cataglyphis rosenhaueri	Sierra Nevada, Spain	AB010933	-
19.	Lasius spathepus	Kokubunji, Tokyo Met. Japan	AB007983	-
20.	Polyergus rufescens	Sierra de Huetor, Spain	AB010931	-
21.	Oecophylla smaragdina	Lambir, Salawak, Malaysia	AB019426	-
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tions are so small that sample locations do not affect the phylogeny reconstruction.

DNA extraction

The procedure used for the extraction of bacterial DNA was modified from Laird *et al.* (1991). The midguts and ovaries of ants were dissected and washed in 0.4% NaCl. The tissue sample was homogenized in a 1.5 ml micro-centrifuge tube containing 500 μ l lysis buffer (100 mM Tris-HCl, 5 mM EDTA, 0.2% SDS, 200 mM NaCl). The mixture was frozen in liquid nitrogen and defrosted at 50°C. Fifty μ g of Proteinase K (Wako Chemicals) was added to the mixture, and incubated overnight at 55°C. Then, 10 μ g of RNase A (Boehringer Mannheim) was added and incubated at 37°C for 30 min. Following a series of phenol-chloroform and chloroform extraction, DNA was precipitated with an equal volume of isopropanol. Pelleted DNA was dissolved in 40 μ l TE buffer (10 mM Tris-HCl, 1 mM EDTA). Genomic DNA of the host was also extracted from a mesosoma of an individual ant by the same method, which was uesd as the templates for the amplification of the COI gene.

Amplification and purification

A partial 16S rRNA region was amplified by PCR (Saiki et al., 1988). The primer sequences used in this study are shown in Table 2. The PCR was performed with universal primers that were designed to amplify a fragment of 16S rRNA of eubacteria approximately 570 base pair (bp) long (Lane et al., 1985). Reactions were performed in a GeneAmp 2400 thermal cycler (Perkin-Elmer) with the following conditions: 35 cycles of denaturation at 94°C for 30 sec, annealing at 54°C for 1 min and extension at 72°C for 2 min, followed by an additional extension at 72°C, 1 min. Final volume of the reaction mix was 40-µl composed of 30 µl of distilled water, 4 µl of 10xPCR buffer (100 mM Tris-HCl [pH8.3], 500 mM KCl, 15 mM MgCl₂, 0.01%[w/v] gelatin, Takara), 4 µl of dNTP mix (1 mM each of dATP, dGTP, dCTP and dTTP), 0.4 µl each of a pair of primers (100 mM), 0.7U of Taq polymerase (Takara TaKaRa Taq) and 1 µl of template DNA (10–200 ng). The PCR products were purified using Prep-A-Gene DNA Purification Kit (Bio Rad). The COI region was also amplified with PCR. Reactions were performed in the GeneAmp 2400 thermal cycler with the following conditions: 30 cycles of denaturation at 94°C for 1 min, annealing at 45°C for 1 min and extension at 60°C for 3 min, followed by an additional extension at 60°C for 1 min. The purification of the PCR products was performed by the same procedure. Purified products of the COI gene were subjected to the direct sequencing.

 Table 2.
 Primer sequences used for amplification of bacterial 16S

 rRNA gene and COI gene.
 COI gene.

region	name	sequence (5'-3')
16S rRNA	16S F-2	AACAAGATTAGATACCCTGGG
	UNI 1400 R	ACGGGCGGTGTGTA(AG)CAA
COI	COI 1-3	ATAATTTTTTTTATAGTTATACC
	COI 2-1	CTTTATCAACATTTATTTTGATTTTT
	COI 2-3	GAAGTTTATATTTTAATTTTACC
	COI 2-2	ACTCCAATAAATATTATAATAAATTGA
	COI 2-4	TCCTAAAAAATGTTGAGGAAA

Cloning of 16S rRNA

Purified products were ligated into pUC18 vector with a SureClone ligation kit (Pharmacia), and competent cells of *E. coli* (DH5) were transformed. They were inoculated on the LB ager plates containing ampiciline (50 mg/ml) and cultivated over night at 37°C. Single colonies were inoculated into 3 μ l of LB medium containing ampiciline (50 mg/ml), followed by the 20 hr of liquid cultivation at 37°C. Mini-preparation of vecter DNA was performed with boiling method. Length of insert in vecter DNA was checked with *Eco*RI-*HindI*II double diges-

tion. Clones with appropriate inserts were purified with a series of phenol/chloroform and diethylether extraction, after the digestion of RNA with RNase A. Purified DNA was dissolved in 10 μ l distilled water and used as the template for the sequencing reaction.

Nucleotide sequencing

Sequencing reactions were performed [on the GeneAmp 2400 thermal cycler] with the dideoxy-nucleotide cycle sequencing procedure using a Dye-Primer Cycle Sequencing Kit for the 16S rRNA clone and a Dye-Terminator Cycle Sequencing Kit (Perkin-Elmer) for the purified PCR products. Electrophoresis and data collection were performed using an automatic DNA sequencer (Perkin-Elmer model 373S) with 5.5% polyacrylamide gels (Super Reading DNA Sequence Solution, Toyobo).

Phylogenetic analysis

Sequences were analyzed by the neighbor-joining method (Saitou and Nei, 1987). The alignments and tree constructions were performed with the Clustal W program package (Thompson et al., 1994). Gap positions were included in the analysis. The numbers of the nucleotide substitutions were estimated according to Kimura's twoparameter method (Kimura, 1980). Bootstrap confidence intervals (Felsenstein, 1985) on each branching pattern were calculated from 1,000 resamplings. Parsimony analysis was also carried out with the PAUP 3.1.1 software program package (Swofford, 1993), with the heuristic search option with 100 random addition replicates. Gaps were treated as new states. Relative support for different nodes was assessed using 1,000 bootstrap replicates (Felsenstein, 1985) with 5 random addition sequence replicates for each bootstrap replicate. The following sequences of bacterial 16S rRNA were obtained from the GenBank database for the phylogenetic analysis of the endosymbionts; Escherichia coli (V00348), Ewingella americana (X88848), Glossina pallidipes S-endosymbiont (M99060), Melaphis rhois Pendosymbiont (M63255), Proteus vulgaris (X07652), Salmonella sofia (X80677), Schlectendalia chinensis P-endosymbiont (Z19056), Yersinia pestis (Z75317) and Xenorhabdus beddingii (D78006). Haemohilus ducreyi (M63900), H. influenzae (Z22806), Pasteurella haemolytica (U57072) and P. trehalosi (U57074), classified into the Pasteurellaceae, were set as the outgroup. The nucleotide sequence data of the 16S rRNA of the symbionts of the ants and the COI of the hosts reported in this paper will appear in the DDBJ/EMBL/GenBank nucleotide sequence databases with the accession numbers listed in Table 1.

RESULTS

Identity of the clones of the 16S rRNA gene

Preliminary sequencing of eight clones of the gene encoding the 16S rRNA derived from the midgut of Camponotus vitiosus, and of six clones derived from the ovaries showed that 11 of the sequences were identical, and that the other three clones differed by only 1 or 2 bps. It is likely that these small differences were due to ether reproduction errors of the Tag polymerase or point mutations. Therefore we concluded that all of the clones originated from the same endosymbiont. Eight clones from the midgut and six clones from the ovaries of Plagiolepis pigmaea were also sequenced; all 14 clones were identical. The PCR products of both C. vitiosus and P. pigmaea were also sequenced directly and the sequences were identical to those of the majority of the clones. Therefore, direct sequencing was used to identify the endosymbionts of the rest of the ant species. The length of the 16S rRNA gene analyzed in this study ranged from 567 to 575 bps.

Insertions of 5 bps in *C. vitiosus* and 4 bps in *C. nawai* were present at the same position. This insertion region was excluded from the analysis.

Phylogenetic analysis of the endosymbionts

Homology searches of the GenBank database showed that the 16SrRNA gene of the endosymbionts were more than 90% homologous to those of the Enterobacteriaceae. Pre-

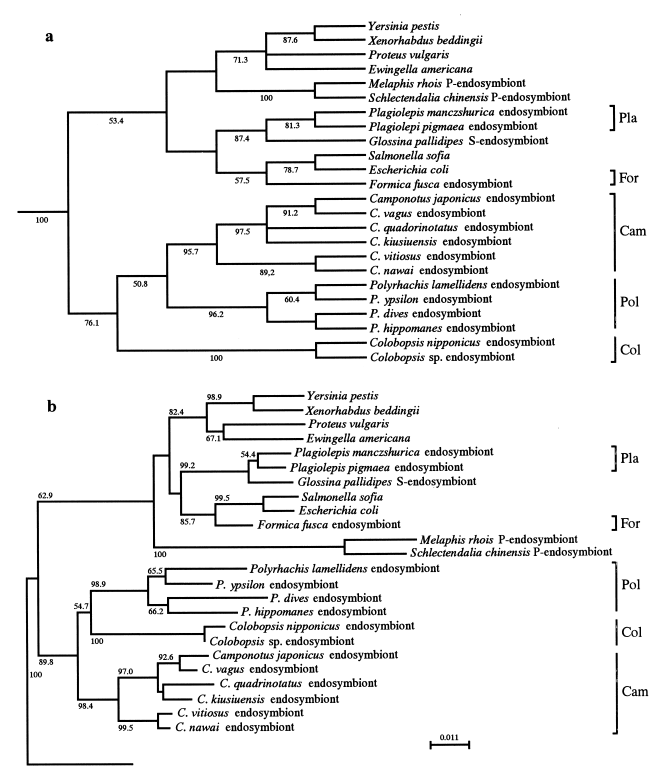


Fig. 1. Phylogenetic trees of the endosymbionts of ants based on 570 bp of the 16S rRNA gene sequence. a: Strict consensus tree of parsimony analysis. Numbers on the branches indicate the percentage of 1,000 bootstrap replicates. b: Bootstrapped tree of neighbor-joining analysis. Numbers on the branches indicate the percentage of 1,000 bootstrap replicates. Cam; *Camponotus*, Col; *Colobopsis*, Pol; *Polyrhachis*, For; *Formica*, Pla; *Plagiorepis*.

liminary phylogenetic analysis of eight groups of the gamma subdivision of the Enterobacteriaceae (Alteromonadaceae, Enterobacteriase, Methylococcaceae, Halomonadaceae, Pasteurellaceae, Pseudomonas, Vibrionaceae, Xanthomonas) rooted with the Alcaligenaceae and the Neisseriaceae classified into the beta-subdivision (Olsen *et al.*, 1994) showed that the endosymbionts of the Formicinae form a clade with the Enterobacteriase. Therefore, four species in the Pasteurellaceae were assigned to the outgroup in the following analysis (data not shown). The species names and the accession numbers of the outgroup are described in the materials and methods section.

Trees were constructed using the 570-bp 16S rRNA gene sequences by the maximum parsimony (MP) method (Fig. 1a) and by the neighbor-joining (NJ) method (Fig. 1b). The parsimony analysis resulted in four most parsimonious trees with 454 steps, and a strict consensus tree. The results indicated

that the endosymbionts of five genera of the Formicinae were the Enterobacteriaceae or were closely related to the Enterobacteriaceae.

The endosymbionts of the ants did not constitute a monophyletic group. The endosymbionts of *Plagiolepis* spp. and *Formica fusca* were both in the clade of the Enterobacteriaceae. The endosymbionts of *Plagiolepis* spp. were the closest relatives of the S-endosymbiont of the tsetse fly, *Glossina pallidipes*. The endosymbionts of *Formica fusca* formed a monophyletic group with *Salmonera* and *Escherichia*. The endosymbionts of three host genera in the Camponotini (*Camponotus, Colobopsis* and *Polyrhachys*) constituted a single clade, which was the sister group of the Enterobacteriaceae. The monophyly of the clade was supported by relatively high bootstrap values (MP, 76.1%; and NJ, 89.8%). In this clade, the endosymbionts of each host genera formed monophyletic groups, supported by bootstrap values of more

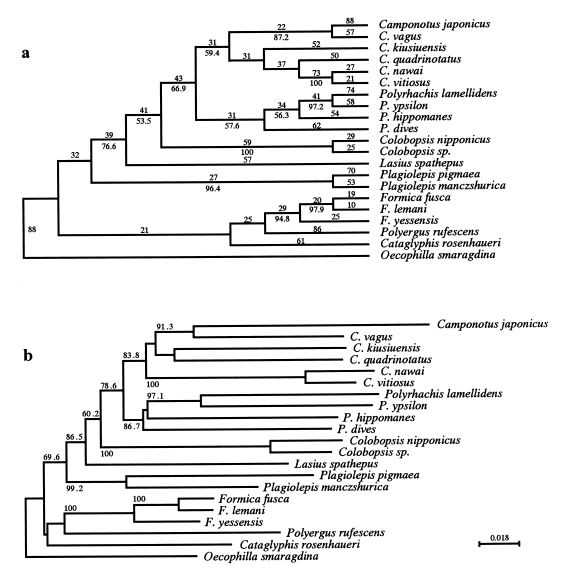


Fig. 2. Phylogenetic trees of the host ants based on 974 bp of the COI gene sequence. a: The single most parsimonious tree of the parsimony analysis is shown. The numbers above the branches indicate the branch length, and the numbers below the percentage of 1,000 bootstrap replicates. b: Bootstrapped tree of neighbor-joining analysis. Numbers on the branches indicate the percentage of 1,000 bootstrap replicates.

than 90%. The MP tree indicated that the relationships of the symbionts of the three genera in the Camponotini were (*Colobopsis* + (*Polyrhachys* + *Camponotus*)), whereas the NJ tree supported ((*Colobopsis* + *Polyrhachys*) + *Camponotus*), however statistical support was not significant.

The phylogeny of the host ants

The COI gene was sequenced in 21 species in Formicinae including five genera and 15 species of hosts. The length of the sequence used for the analysis in this study was 974 bps.

The parsimony analysis of the ant species resulted in a single most-parsimonious tree with 1702 steps (Fig. 2a). The members of each of the five host genera formed clades. The monophyly of each host genus was also shown by NJ analysis (Fig. 2b). In both analyses, the five host genera did not form a monophyletic group, although three of these genera, the members of the Camponotini, were monophyletic. These relations reflected those between the endosymbiont groups, however the branching patterns of the endosymbionts were not the same as those of the hosts. The analysis of the hosts showed that *Formica* was in the basal position and the Camponotini was at the apex.

Our analyses by the MP and NJ methods both resulted in the same topology for the three genera of the Camponotini with bootstrap values of more than 50% in each case: (*Colobopsis* + (*Polyrhachys* + *Camponotus*)). Three species of *Formica* formed a monophyletic group with *Polyergus rufescens* and *Cataglyphis rosenhaueri*; they all belong to the Formicini, but the relationships among the three genera were supported by less than 50% of the bootstrap replicates.

DISCUSSION

It is usually impossible to extract enough morphological characters for phylogenetic analysis from prokaryotic microorganisms; ribosomal genes provide the most useful information for phylogeny reconstruction and systematic characterization of microorganisms, and are the most commonly used molecular markers for phylogenetics (Carl, 1987). However, careful interpretation is necessary to align the sequences, because insertions and deletions are more frequent than in protein-coding genes. In this study, 5 bps of insertions found in Camponotus vitiosus and 4 bps in C. nawai were excluded from the analysis. They seemed to originate from a single insertion event, which took place on a common ancestor of the subgenus Myrmamblys. Other deletions, which were all single base deletions, were included in the analysis to maximize the available information. Gap-exclusive analysis was also performed and gave the same topologies as gap-inclusive analysis both by the MP and NJ methods (data not shown).

To establish the phylogenetic tree of the host ants, we choose to study the COI gene because it has excellent properties as a molecular marker; it is the largest protein-coding mitochondrial gene and detailed information is available about its protein product, a respiratory enzyme, including variability of each domain (Lunt *et al.*, 1995). These features enable us

to perform multiple factor analysis based on information about codon positions or domains. We estimated the saturation of nucleotide substitutions to plot the number of transitions and transversions against the uncorrected genetic distance for each codon position (Reo *et al.*, 1997) and we found that saturation of transitions may have occurred at the third codon. Therefore, weighted parsimony analysis was performed using four weighting schemes: weighting transversions 2, 4, and 8 times of transitions at the third codon and excluding transitions at the third codon (Reo *et al.*, 1997). However, the topologies we obtained using these schemes were all identical to that of the non-weighted MP tree (data not shown).

It has been proposed that the endosymbionts of the Formicinae have a common origin for two reasons. First, the distribution of the endosymbionts is restricted to the subfamily Formicinae. Second, the morphology and the pattern of transmission of the endosymbionts is similar among the Formicinae (Jungen, 1968; Dasch *et al.*, 1984). However, the molecular phylogeny obtained in this study did not support this hypothesis of a common origin for both the endosymbionts and the hosts. Instead, the results of the present study suggested that the endosymbionts have been independently acquired by the Formicinae at least three times: by the common ancestor of the Camponotini, by the ancestor of *Plagiolepis*, and by the ancestor of *Formica*.

Within the Camponotini, the phylogeny of the endosymbionts showed strong, if not perfect, congruence with that of the hosts. In our topological comparison among the symbionts and the hosts within the Camponotini (Fig. 3) clades supported by bootstrap values of less than 50% were treated as polytomy. Camponotus, Polyrhachys and Colobopsis formed a single clade in all cases, and the topology (Colobopsis + (Polyrhachys + Camponotus)) was supported in three cases (Fig. 3a, 3c and 3d). Within Camponotus, the branching patterns of four analyses had nearly identical topologies. The situation was the same within the genus Polyrhachys with the exception of P. dives, which formed a subcluster with P. hippomanes in the analysis of the symbionts by the NJ method (Fig. 3b), although it was in the basal position of the clade of Polyrhachys in the analysis of the hosts by the MP method (Fig. 3c). Dasch et al. (1984) studied the endosymbionts of 8 of 14 known genera of the Camponotini and concluded that mycetocyte symbiosis is common in this tribe. We suggest that a single bacterial infection occurred in a common ancestor of the Camponotini, followed by co-speciation of the hosts and the endosymbionts. Improved data matching may be seen in future analyses after more data has been collected, in particular on Colobopsis.

Schröder *et al.* (1996) compared the 16S rRNA gene of the endosymbionts of four *Camponotus* spp. with the endosymbionts of aphids (*Buchnera*), P-endosymbionts of the tsetse fly and other members of the Enterobacteriaceae. Their results indicated that the *Camponotus* symbionts form a clade ((*Camponotus* + tsetse fly P-symbionts) + aphid) that is adjacent to the other members of Enterobacteriaceae. However, we obtained a different result: the endosymbionts of

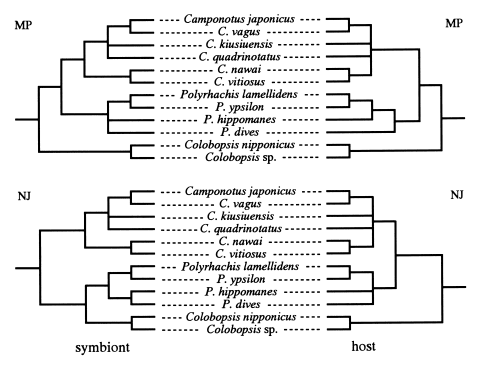


Fig. 3. Topological comparison between the symbionts and the hosts within the Camponotini. a, c: MP method; b, d: NJ method

Camponotus formed a clade with those of *Colobopsis* and *Polyrhachis*, and this clade was adjacent to the cluster of the Enterobacteriaceae, that contained *Buchnera*. The bootstrap values that supported this clade as a sister clade of the Enterobacteriaceae were low (54.3% in MP, and 62.9% in NJ), however the clade consisting of the symbionts of *Camponotus*, tsetse fly and aphid was not supported statistically (Schröder *et al.* 1996). The relationships among the endosymbionts of these insects, in either case, is not certain.

Two taxonomic treatments have been proposed for "*Colobopsis*"; an independent genus (Brown, 1973; Sneling, 1981; Hölldobler and Wilson, 1990) and a subgenus in the genus *Camponotus* (Bolton, 1996). The molecular phylogeny we present here supported the former viewpoint. Our result reflected the current subgeneric classification of the genus *Camponotus* and *Polyrhachis* (Hung, 1967; Bolton, 1996). The species *C. japonicus* and *C. vagus*, classified into the subgenus *Camponotus*, formed the subcluster (Figs. 1 and 2). Similarly the following species formed subclusters; *C. vitiosus* and *C. nawai*, into the subgenus *Myrmamblys*, *P. lamellidens* and *P. ypsilon*, into the subgenus *Polyrhachis*.

The presence of mycetocyte endosymbionts in Formicinae has not been surveyed comprehensively. Thus, new ant species that possess endosymbionts may be discovered in other genera or subfamilies. In the case of the genus *Formica*, endosymbiosis has been reported for only a small fraction of the known species (Jungen, 1968). Prior to this study, mycetocyte symbionts were preliminarily surveyed in several *Formica* species (*F. japonica*, *F. lemani*, *F. yessensis*) in Japan. However no mycetocytes were found in these species. Jungen (1968) reported that the infection rate by endosymbionts in several colonies of *Formica lemani* was not 100% and that some colonies lack the symbionts. Considering these circumstances, more data is needed to clarify the biology of the endosymbionts of *Formica*. Phylogenetic analyses of the symbionts in other genera in the Formicini and in the Plagiolepidini will also be indispensable to establishing a detailed scenario of the establishment of the symbiosis.

Little is known about the biological relationships between ants and their endosymbionts, although many mycetocyte symbionts are known to have nutritional interactions with their hosts (Noggi, 1981; Cochran, 1985; Lai, 1994). It seems suggestive that the P-endosymbionts of the tsetse flies and aphids, which provide the host with essential nutrients, are related to the endosymbionts of the Formicinae. Although Smith (1944) discussed the relation between nutritional condition and infection by endosymbionts in *Camponotus*, there has been no experimental study of the contribution of endosymbionts to growth or survival of host ants. To understand the functions and biological significance of the endosymbionts of ants, experimental investigations, such as the removal of the symbionts from their hosts and feeding experiments, are essential. Detailed information regarding the phylogeny of the endosymbionts and their hosts should be useful for comparative studies of the relationships among various ant symbioses, and of the metabolism of the symbionts and their free-living relatives.

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