

Epidemiology of asexuality induced by the endosymbiotic Wolbachia across phytophagous wasp species: host plant specialization matters.

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- 1 Title: Epidemiology of asexuality induced by the endosymbiotic Wolbachia across phytophagous wasp species: host plant specialization matters 2 3 Authors: Boivin T¹, Henri H², Vavre F², Gidoin C¹, Veber P², Candau J-N^{1,3}, Magnoux E⁴, 4 Roques A⁴ and Auger-Rozenberg M-A⁴ 5 6 **Addresses:** 7 1. INRA, UR629 Ecologie des Forêts Méditerranéennes, URFM, F-84914 Avignon, France 8 2. Université de Lyon, F-69000, Lyon ; Université Lyon 1 ; CNRS, UMR5558, Laboratoire de Biométrie et Biologie Evolutive, F-69622, Villeurbanne, France. 9 10 3. National Resources Canada, Canadian Forest Service, Great Lakes Forestry Ctr, Sault 11 Ste Marie, ON P6A 2E5, Canada 4. INRA, UR633 Unité de Recherche de Zoologie Forestière, 45075 Orléans, France 12 13 14 **Keywords**: parthenogenesis, ecological specialization, thelytoky, multilocus sequence typing, Megastigmus 15 16 17 Word count: 7770 18 Corresponding author: Thomas Boivin, INRA, UR629 Ecologie des Forêts Méditerranéennes, 19 URFM, Domaine Saint Paul, F-84914 Avignon, France, fax: +33 (0)4 32 72 29 02, 20
- 23 Running title: Spread of thelytoky in phytophagous wasps

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Abstract

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Among eukaryotes, sexual reproduction is by far the most predominant mode of reproduction. However, some systems maintaining sexuality appear particularly labile and raise intriguing questions on the evolutionary routes to asexuality. Thelytokous parthenogenesis is a form of spontaneous loss of sexuality leading to strong distortion of sex ratio towards females and resulting from mutation, hybridization or infection by bacterial endosymbionts. We investigated whether ecological specialization is a likely mechanism of spread of thelytoky within insect communities. Focusing on the highly-specialized genus Megastigmus (Hymenoptera: Torymidae), we first performed a large literature survey to examine the distribution of thelytoky in these wasps across their respective obligate host plant families. Second, we tested for thelytoky caused by endosymbionts by screening in 15 arrhenotokous and 10 thelytokous species for Wolbachia, Cardinium, Arsenophonus and Rickettsia endosymbionts and by performing antibiotic treatments. Finally, we performed phylogenetic reconstructions using multilocus sequence typing (MLST) to examine the evolution of endosymbiont-mediated thelytoky in Megastigmus and its possible connections to host plant specialization. We demonstrate that thelytoky evolved from ancestral arrhenotoky through the horizontal transmission and the fixation of the parthenogenesisinducing Wolbachia. We find that ecological specialization in Wolbachia's hosts was probably a critical driving force for Wolbachia infection and spread of thelytoky, but also a constraint. Our work further reinforces the hypothesis that community structure of insects is a major driver of the epidemiology of endosymbionts and that competitive interactions among closely related species may facilitate their horizontal transmission.

Introduction

Sex is by far the most predominant mode of reproduction among eukaryotes, such that only one out of a thousand animal species shows some type of asexual reproduction (Suomalainen *et al.*, 1987). Among the diverse mechanisms involved in asexual reproduction, parthenogenesis is defined *sensus stricto* as the development of an egg without fertilization. The origin of parthenogenesis is polyphyletic in both invertebrates and vertebrates, suggesting that systems maintaining sexuality are labile and stimulating elucidation of the evolutionary routes to parthenogenesis (reviewed in Simon *et al.*, 2003). Spontaneous loss of sexuality may occur through mutations or intra- and interspecific hybridization (Simon *et al.*, 2003; Dedryver *et al.*, 2001), but it can also result from infection by bacterial endosymbionts that are predominantly transmitted vertically through the female egg cytoplasm and distort the host sex ratio towards females to their own advantage (Engelstädter and Hurst, 2009).

In the insect order Hymenoptera, several forms of parthenogenesis have been defined according to the sex of offspring produced by a virgin female (Cook, 1993; Cook and Crozier, 1995). The dominant and ancestral form of parthenogenesis is arrhenotoky, in which fertilized eggs develop as diploid females and unfertilized eggs develop as haploid males. However, numerous species display thelytoky, which is a form of complete parthenogenesis where unfertilized eggs develop into diploid females. Thelytoky can be under the control of the insect itself or its endosymbionts (Stouthamer et al, 1990; Vavre et al., 2004). Parthenogenesis inducers include bacteria of the genera *Wolbachia* (Werren, 1997), Cardinium (Zchori-Fein and Perlman, 2004) and Rickettsia (Hagimori et al, 2006). While

induction of male production is impossible in genetically-based thelytoky, endosymbiotic thelytoky can be characterized by its reversibility, as antibiotic or heat treatment of females leads to bacteria elimination and production of males in their progeny (Stouthamer *et al.*, 1990). Generally, infection by parthenogenesis-inducing (PI) *Wolbachia* is fixed within species or populations and results in an obligate parthenogenesis associated with a loss of sexual function in females (Jeong and Stouthamer 2005; Kremer *et al.*, 2009), or in both sexes (Gottlieb and Zchori-Fein, 2001), making *Wolbachia* indispensable. Only in very rare cases, polymorphism of infected and uninfected females is maintained (REF).

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Despite vertical transmission via the egg cytoplasm being the predominant transmission mode of PI endosymbionts (Werren, 1997), phylogenetic incongruence between host and endosymbiont histories strongly indicates occasional horizontal transmission events (O'Neill et al., 1992; Stouthamer et al., 1993; Baldo et al. 2006; Perlman et al., 2010). Horizontal transmissions of Wolbachia have been successfully demonstrated experimentally by injection (Braig et al., 1994; Grenier et al, 1998), introgression (Jaenike, 2007), maintenance of close contacts between conspecifics (Rigaud and Juchault, 1995) or in host-parasitoid associations (Vavre et al. 1999). Effective horizontal transmission of endosymbionts depends (i) on intimate ecological associations which provide withincommunity horizontal transmission opportunities (Vavre et al., 1999; Sintupachee et al., 2006; Stahlhut et al., 2010) and (ii) on the phylogenetic similarity of donor and recipient host species because internal defence mechanisms against infections are likely to be more similar between closely related hosts (Stahlhut et al., 2010). Studies addressing how insect phylogeny and ecology affect patterns of similarity between strains of PI endosymbionts would help gain critical insights on how these two forces affect shifts in reproduction modes.

In particular, study systems involving several groups of closely related thelytokous species but displaying different levels of ecological proximity due to habitat specialization can be of critical interest to define the possible ecological boundaries to the evolutionary trajectory of thelytoky in insects.

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In this paper, we address the potential for phylogeny, ecology and infection by PI endosymbionts to drive the spread of asexuality within insects. For this purpose, we focused on the genus Megastigmus (Hymenoptera: Torymidae), which counts more than 125 arrhenotokous and thelytokous wasp species worldwide (Grissell, 1999). Half of these species are specialist seed feeders, the other half are presumed to be parasitoids, gallmakers or to have unknown hosts. Within the seed-specialized group, two thirds of species exploit gymnosperm hosts (Pinaceae and Cupressaceae families), whereas one third exploits preferentially angiosperms, especially the Rosaceae and Anacardiaceae families (Roques & Skrzypczynska, 2003). Being highly specialized in the seed resource, several wasp species exploiting the same host strictly require the same ecological niche, which is then particularly propitious to direct intra- or interspecific interactions within this insect group (Boivin et al., 2008; Auger-Rozenberg and Roques, 2012). In the genus Megastigmus, the dominant and ancestral form of parthenogenesis is by far arrhenotoky as thelytoky characterizes only a minority of species (Grissell, 1999), but both arrhenotokous and thelytokous species can occur in sympatry on gymnosperm or on angiosperm hosts (Boivin et al., 2008; Auger-Rozenberg and Roques, 2012). Thelytoky has only been established from records of highly female-biased sex ratios in field populations, but neither its determinism nor its evolutionary trajectory have been elucidated yet. We first performed a large literature survey to examine the distribution of thelytoky in *Megastigmus* across their known obligate host-plant families. Second, we tested for thelytoky caused by endosymbionts by screening in a large sample of arrhenotokous and thelytokous species for endosymbionts and by performing antibiotic treatments. Finally, we performed phylogenetic reconstructions to examine the evolution of endosymbiont-mediated thelytoky in *Megastigmus* and its possible connections to host-plant specialization. This study strongly suggests that thelytoky is caused by *Wolbachia* endosymbionts, and interestingly, we show that host plant specialization is a key determinant of *Wolbachia* infection and thelytoky.

Material and Methods

Distribution of thelytoky among phytophagous Megastigmus species

Parthenogenesis, *i.e.* arrhenotoky or thelytoky, is the only mode of reproduction in the *Megastigmus* genus. It has never been formally studied and it was generally deduced from sex ratio estimates in field sampled wasp populations. Here, we used extensive catalogs of seed-feeding *Megastigmus* species to gather the current knowledge on the relative prevalence of arrhenotoky and thelytoky in the genus *Megastigmus* (Grissell, 1999; Roques & Skrzypczynska, 2003; Roques *et al.*, 2003; Auger-Rozenberg *et al.*, 2006). A bibliographic search of non-redundant articles in the Web of Science (2003-2012) and CAB abstracts (2003-2012) was also performed on key terms *'Megastigmus* and seed*' to check for potential new species descriptions or increase knowledge in the biology of the species described in the above catalogs. We considered only wasp species for which arrhenotoky or thelytoky were ascertained by authors, or for which male frequencies estimated at emergence from sampled seed lots provided unambiguous support for arrhenotoky (high male abundance) or thelytoky (absence or <1%). Species for which the parthenogenesis

mode was not mentioned or for which sample sizes were insufficient (<10) to estimate sex ratios were not included. The prevalence of thelytoky was assessed as the percentage of thelytokous species in all of the selected wasp species. The distributions of both arrhenotoky and thelytoky relative to host plant specialization was assessed by assorting both arrhenotokous and thelytokous wasp species according to their host plant group (gymnosperm or angiosperm), family and genus.

Wasp sampling

Megastigmus species were collected across the Northern Hemisphere (Nearctic and Palearctic) by seed samplings on diverse host plants. Insect-infested seeds were separated from non-infested ones by X-ray radiography (using Faxitron-43855*, 15 kV, 3 mA, 3'30" to 4'30" depending on seed species with X-ray-sensitive films Kodak 'Industrex M', and Faxitron-MX20*, 20 kV, 0.3 mA, 1'45" with an EZ20 digital scanner). The insect-infested seeds were placed in individual rearing boxes stored in outdoor insectaries located at INRA, Orléans, France and at INRA, Avignon, France. Adult emergence was recorded over the 3 years following seed maturation because of a possible prolonged diapause (Roques & Skrzypczynska, 2003). After emergence, insect species determination was ascertained using the morphological keys of Grissell (1999) and Roques and Skrzypczynska (2003). Emerged insects were then sorted by species and collection site and preserved in 100% alcohol at 20°C. A total of 25 Megastigmus species were used in this study, 10 were thelytokous and 15 were arrhenotokous. These species, their reproduction mode, their host-plants, collecting localities and distributions are summarized in Table 1.

DNA extraction

Total genomic DNA was isolated by crushing individually whole adult *Megastigmus* females using two different procedures, depending on sample size and further DNA use. Screening for PI endosymbiont infection, which involves large sample sizes, was performed using the Chelex method. This fast and cheap method consisted of digesting tissues kept for 2 h at 56°C in a 200 µl 10% Chelex solution (Biorad). After 30 min at 100°C, samples were centrifuged and supernatants were used as DNA sources. For sequencing and further phylogenetic analyses, which requires smaller sample sizes and long term conservation of DNA, we used the DNeasy extraction Tissue Kit (Qiagen). Total genomic DNA was eluted in 200 µl of AE buffer in this case. In addition, a PCR was systematically carried out on Qiagen extractions with *Wolbachia* primers in order to confirm the lack of bias in the detection of the symbiont due to different methods of DNA extraction.

Wasp sequences

We used Sigma RedTaq for PCR amplification. The forward and reverse primers used were I775-COI-F (Clyde, 5'-CGAATAAATAATATAAGATTTTG-3'), and 2773-COI-R (Bonnie, 5'-GGATAATCTCTATATCGACGAGGTAT-3') (Scheffer & Grissell 2003) for the segment of the cytochrome oxidase I (COI) gene. Cycling programs were as follows: a denaturation step at 94°C for I min, annealing for I min at 48°C, and extension at 72°C for I min with 30 cycles being performed. All PCR products were then purified with a QIAquick PCR purification kit (Qiagen) and were directly sequenced with the amplification primers. Sequencing was performed using the big-dye terminator sequencing kit (PE Applied Biosystem) and carried out with an ABI 3100 automatic sequencer. The gene fragment was sequenced on two to four individuals for each species, except for cryptic and rare species because of the small number of specimens available. When individuals were identified as

originating from a different biogeographic region (i.e. when invasive specimens were found), additional specimens from native areas were sequenced in order to ensure genetic similarity between native and introduced individuals. Sequences were aligned using Clustal W (Thompson et al. 1994) as implemented in BioEdit 7.05 (Hall 1999).

In addition to the COI gene, we studied a nuclear fragment, the D2 region of the 28S ribosomal subunit (rDNA), to build a phylogenetic tree of the studied *Megastigmus* species. Nuclear primers previously used for reconstructing a molecular phylogeny of *Megastigmus* spp. on conifers (Auger-Rozenberg *et al.*, 2006) were chosen due to their utility for molecular identification at intrageneric taxonomic level. However, the 28S data were not used further because their resolution was insufficient for the goal of this study (results not shown).

Population screenings for PI-endosymbiont infection

In a wide range of insect species, sex ratio distortion has been associated with infection by diverse bacterial symbionts such as *Wolbachia* (Werren, 1997), *Cardinium* (Zchori-Fein and Perlman, 2004), *Arsenophonus* (Thao and Baumann, 2004) and *Rickettsia* (Hagimori et al, 2006). As a first test for an association between such endosymbionts and thelytoky in the *Megastigmus* genus, we screened for their presence in both arrhenotokous and thelytokous species (listed in Table 1). Separate Polymerase Chain Reactions (PCR) were performed on each *Megastigmus* female using specific primers for *Wolbachia* surface protein (wsp) (Braig *et al.*, 1998), *Cardinium* 16S rRNA (Zchori-Fein and Perlman, 2004), *Arsenophonus* 23S rDNA (Thao and Baumann, 2004) and *Rickettsia* 16S rRNA (Hagimori et al, 2006) gene fragments.

PCR reactions were done in a 25-μl final volume reaction containing 200 μM dNTP, 10 pM primers, 0.5 IU HotStarTaq® DNA polymerase, and 1.5 μl DNA solution. All reactions were

run in a ABI thermal cycler (PE Applied Biosystems PCR System 9700) with an initial denaturing step at 94°C for 4 min, an annealing step for 1 min, an elongation step at 72°C for 1 min 30 s. Annealing temperatures and primer pairs used for *Wolbachia*, *Cardinium*, *Arsenophonus* and *Rickettsia* PCR screening are detailed in Table S2. All PCR reactions included a positive control of *Asobara tabida* (*Wolbachia* infected) and *Bemisia tabaci* biotype Q (*Cardinium*, *Arsenophonus* or *Rickettsia* infected) samples. For each bacterial gene, 5 µl of amplified reaction product was run on a 1% agarose gel stained with ethidium bromide after the PCRs in order to determine the presence of an amplified DNA fragment. If a sample of a *Megastigmus* species was negative for any bacterial-specific amplification, but COI amplification succeeded, the insect was considered uninfected. When amplification with a bacterial-specific primer yielded visible bands, the insect was considered infected, and samples were kept for sequencing and strain characterization.

Wolbachia strain characterization

Only *Wolbachia* was detected in our samples (see Results section). We used multilocus sequence typing (MLST) (Baldo *et al.*, 2006) to characterize *Wolbachia* strain similarity among *Megastigmus* host species. A *wsp* sequence was also obtained from each infected insect. MLST was based on the methods of Baldo *et al.* (2006) using the standard primers that amplify the five conserved *Wolbachia* genes *hcpA*, *ftsZ*, *gatB*, *coxA* and *fbpA*. These are degenerate primers designed to amplify sequences from diverse *Wolbachia* strains. Despite repeated attempts, we could not amplify the fbpA gene. Thus, four MLST loci (*hcpA*, *ftsZ*, *gatB* and *coxA*), plus the *wsp* gene, were sequenced in both directions to provide double

coverage of the region of interest, with the exception of both *M. borriesi* and *M. rosae* var. alba, which could be sequenced for the wsp gene only.

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Phylogenetic analyses

Wolbachia. Two samples of each species were sequenced and revealed no polymorphism between individuals. A single sequence was therefore chosen for alignment to a set of database sequences for phylogenetic analysis. Our multi-locus gene phylogeny was based on the four successfully amplified loci (coxA, gatB, hcpA, ftsZ) used by Baldo et al. (2006) and includes many Wolbachia strains identified as from A and B supergroups. Sequences are accessible from the GenBank database under accession numbers KF531859 to KF531890. Wolbachia from Brugia malayi was used as an outgroup. As more sequences are available for wsp gene (in our sampling as well as in Genbank database) the analysis was conducted separately for wsp sequences for fine scale genotyping of Wolbachia harboured by Megastigmus species since this gene often evolves rapidly (Genbank accession numbers KF531891 to KF531900). For all datasets, alignments were initially generated using the MUSCLE software (Edgar, 2004) implemented in CLC Main Workbench v6.7.1 (CLC Bio). Phylogenetic analyses were performed using maximum likelihood (ML) inference with PhyML v3.0 (Guindon et al., 2010). The appropriate model of evolution was evaluated with jModeltest v0.1.1 (Posada, 2008). The models selected were TIM1+G for coxA and gatB, TIM3+G for ftsZ and hcpA and GTR+I+G for the concatenated MLST data set (coxA + qatB + ftsZ + hcpA) and for the wsp gene. The robustness of the nodes was assessed with 100 bootstrap replicates. Additionally, bayesian inferences were also used to reconstruct phylogenies with MrBayes v3.1.2

(Ronquist & Huelsenbeck, 2003) using appropriate settings leading to convergence between two independent runs. Finally, trees were edited with Figtree v1.4.0 (A. Rambaut, http://tree.bio.ed.ac.uk/software/figtree).

Megastigmus. The same procedure as for *Wolbachia* data was conducted on *COI* gene sequences. Sequences are accessible from the GenBank database under accession numbers KF531833 to KF531858. Models of substitution selected were GTR+I+G for the *COI* gene.

Analysis of congruence. The Shimodaira-Hasegawa (SH) and approximately unbiased (AU) tests were used to evaluate the significance of topological incongruence among trees. First, to check for some methodological artifacts, topologies obtained with the two methods (ML and bayesian inferences) were compared and showed no significant difference. Second, for Wolbachia phylogenies, we compared topologies obtained with the different loci and also with the concatenated data set. These comparisons were performed on the entire dataset (i.e. the Wolbachia from Megastigmus species and other sequences retrieved from Genbank) and for the Wolbachia infecting Megastigmus only. Differences in log likelihood between the best ML topology and an alternative topology were used as values of the AU test and the SH test and then p-values were calculated with Consel program (v0.1i) (Shimodaira & Hasegawa, 2001).

Analysis of host-symbiont associations

Parafit statistic was used to test whether hosts and symbionts evolution were independent or if there was a pattern of co-cladogenesis between them (Legendre *et al.*, 2002). The AxParafit program implemented in the Copycat software v2.00.02 (Meier-Kolthoff *et al.*, 2007) was used with patristic distances as input matrix and 999 permutations both for *COI*

sequences data and for 28S sequences compared with the Wolbachia sequences concatenated data set.

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Antibiotic treatments

When thelytoky is induced by symbionts, antibiotic curing of infected thelytokous females is expected to show reversion to male production with elimination of the bacteria (Stouthamer et al., 1990; Zchori-Fein et al, 1992). Because it was not possible to rear and treat all of the Wolbachia positive Megastigmus species due to the variety of their ecological requirements, we chose to focus antibiotic treatments on one model species, M. pinsapinis, to ascertain the association between thelytoky and Wolbachia infection. Antibiotic treatments were also performed on a closely related arrhenotokous species, M. schimitscheki, to exclude any toxic effects of antibiotics on *M. pinsapinis* reproduction. Two hundred M. pinsapinis females within 24h of emergence from seeds of Cedrus atlantica (Pinaceae) were split into two groups of 100, one dedicated to antibiotic treatment and the other used as a control. The 100 females to be treated with antibiotics were placed in ventilated plastic tubes and were provided with cotton soaked in a sugared solution containing tetracycline hydrochloride (Sigma) at a final 0.2% concentration in the diet. These females were referred to as being cured. The 100 control females were provided with sugared water only. Both cured and control females were left for 5 days at 19°C and natural daylight. Fifty females from each of the cured and control groups were then removed and stored at -20°C in 100% ethanol for subsequent DNA tests for the presence of Wolbachia with PCR tests using the wsp gene. The remaining females of each group were allowed to lay eggs on young conelets of *C. atlantica* after their introduction in insect-proof bags. Because emergence of a cohort of this species may require up to 5 years due to prolonged diapause and because sex is indistinguishable at the larval stage using morphological characters, proportions of males and females in the progeny of both cured and control females were estimated directly on larvae using flow cytometry (Boivin and Candau, 2007). Antibiotic treatments, *Wolbachia* detection and male proportion estimations in the *M. schimitscheki* progeny strictly followed the same procedure as that in *M. pinsapinis*, using 160 newly emerged females to constitute two groups of 80 for the establishment of both cured and control groups, and 30 and 50 females of each group for PCR and progeny production, respectively.

Results

Distribution of thelytoky among phytophagous Megastigmus species

A review of the current knowledge of parthenogenesis in the seed-feeding *Megastigmus* group showed a low prevalence of thelytoky (15%) as only 10 out of 69 species for which the reproduction mode could be unambiguously established are thelytokous (Table S1). Thelytoky tends to be more frequently associated with species developing on angiosperms (26%) than on gymnosperms (9%), but the difference was not statistically significant (Fisher's exact test, P=0.0729). Seed wasps of gymnosperms count 4 thelytokous species, all specialized on Pinaceae, specifically in the genus *Abies, Larix* and *Cedrus* (Table S1). Arrhenotokous species of *Megastigmus* are also found on the same host plant genus. Among seed wasp of angiosperms, thelytoky occurs almost exclusively in species specialized on the *Rosa* genus (Rosaceae) except for one on Anacardiaceae (Table S1).

In order to study the evolutionary history of thelytoky in Megastigmus, thelytokous species must be placed on the phylogenetic tree of the Megastigmus genus. COI reconstruction was performed on sequences obtained from the 25 species sampled in this study (Figure 1). There was an ancestral split between the species related to Anacardiaceae and the others. Wasp species related to Rosaceae form two clades, supported by high bootstrap values, with an initial split between the aculeatus group on the one hand and all others species associated with Rosaceae on the other hand. This latter clade is sister to a monophyletic clade comprising all species exploiting gymnosperms (posterior probability 0.73). In accordance with previous studies, Megastigmus developing on gymnosperms also show specialization, with for example, species exploiting the Pinaceae forming a monophyletic group (except for M. thyoides, the only species related to Cupressacae in the Nearctic region, which could rather reflect a secondary change of host family than an ancestral association). Replacing species reproducing through thelytoky on that tree does not show any strong phylogenetic signal: thelytokous species occur sporadically on the tree. While the ancestral mode of reproduction is difficult to assess, one clear result is that transitions from one mode of reproduction to the other have regularly occurred during the evolutionary history of seedspecialized *Megastigmus*.

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Screening for PI-endosymbiont infection

A total of 25 arrhenotokous and thelytokous *Megastigmus* species (15 and 10 species, respectively) were sampled and screened for the presence of *Wolbachia*, *Cardinium*, *Arsenophonus* and *Rickettsia* using specific primers. None of the arrhenotokous species was found infected by any of the targeted endosymbionts (Table 2). On the contrary, *Wolbachia*

was found fixed in all of the thelytokous samples tested, while *Cardinium, Arsenophonus* and *Rickettsia* were not detected in any of these species (Table 2). The correct amplification of both positive controls and arthropod COI indicated true lack of infection and not a failure in the set-up of the PCR reactions. These results demonstrate the strong association of *Wolbachia* infection with thelytokous parthenogenesis in *Megastigmus* (Fisher's Exact Test's p-value<0.001).

Antibiotic treatments

Tetracycline treatments led to an almost complete loss of *Wolbachia* infection in *M. pinsapinis* females and a significant recovery of male production, whereas control ones remained infected and produced only females (Table 3). As expected for arrhenotokous species, both control and treated *M. schimitscheki* females were not infected by *Wolbachia* and produced exclusively males. In addition, for each species, the mean brood size was not significantly altered by tetracycline treatment (Table 3). These data suggest that arrhenotoky restoration in treated *M. pinsapinis* was due to symbiont curing rather than direct toxicity of antibiotics to insects, and that *Wolbachia* is probably the causative agent of thelytoky.

Diversity and distribution of Wolbachia strains in Megastigmus

Topologies obtained with the four MLST genes were not congruent among each other on the entire dataset (34 taxa; all p-values <0.01 for SH test, <3.10⁻⁵ with AU test), as already shown (Baldo *et al.* 2006). In contrast, for the eight *Wolbachia* infecting *Megastigmus* for which sequences have been obtained on the four MLST genes, topologies were globally congruent as all SH tests and most AU were not significant. The only consistent exception was for the

coxA gene, for which topology was significantly different form the other genes based on AU tests. In relation to that, a closer look at the Wolbachia strain infecting M. pistaciae suggests this strain has undergone a recombination event between A and B Wolbachia. Indeed, this strain belongs to the B supergroup based on all sequenced genes, except for coxA for which it falls within the A supergroup (Figure S1). In summary, despite the complex history of Wolbachia strains at a global scale, the tree based on the concatenated dataset is a good mean representation of the history of Wolbachia infection in Megastigmus (Figure 2) and is also consistent with the tree obtained on the wsp gene (Figure S2). Three highly homogeneous lineages of Wolbachia infecting Megastigmus species were identified (namely WA1, WA2 and WB). We refer to these as lineages as opposed to strains, because although they are closely related, some substitutions were observed. A strict association between Wolbachia lineage and host plant is observed. The WB lineage is restricted to M. pistaciae, the only thelytokous species developing on Anacardiaceae. A second lineage, named WA1, was only found in thelytokous species developing on Rosaceae, i.e. M. brevicaudis, M. rosae (both subtypes) and M. aculeatus (both subtypes). The third lineage, named WA2, is closely related to WA1 and grouped the Wolbachia infecting M. borriesi, M. pictus, M. pinsapinis and M. suspectus, which all develop on Pinaceae. Importantly, and despite this strict association between host plant and Wolbachia lineage, no significant association was detected between Wolbachia and host lineages using Parafit test. In addition, Wolbachia (concatenated dataset) and Wolbachia-infected Megastigmus (COI) topologies were not congruent (SH test, p<0.014 and AU test, p<0.003). These analyses exclude co-cladogenesis, suggesting horizontal transfers have played a major role in the distribution of the Wolbachia strains.

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The distribution of Wolbachia strains among Megastigmus species suggests that horizontal transmission routes are constrained by the plant family, with only one plant family per strain. Our tree in Fig. 1 shows no exception to this rule, but this could nevertheless be obtained by chance, at least in principle. To evaluate how likely this is, we performed a simple test, stating as our null hypothesis that a strain can appear in an insect irrespective of the plant it parasites. Because infection with PI-Wolbachia rapidly leads to the irreversible loss of sexual reproduction, rendering Wolbachia indispensable for the insect, infection loss is highly improbable (Stouthamer et al. 2010). Under this hypothesis, from the 10 infected species, at least 7 independent Wolbachia acquisitions have occurred during Megastigmus evolution. Indeed, in three cases, it is not possible to distinguish between independent acquisition or ancestral infection (e.g. in the two M. rosae). In all other cases, the presence of an uninfected species within the clade excludes ancestral acquisition. Using the constraint solver clingo [1], we first enumerated all possible scenarios with 7 independent Wolbachia acquisitions (at least one by strain) possibly occurring at each node of the actual Megastigmus phylogeny and leading to 10 infected extant species. Among all these scenarios, we reported the proportion p of cases where a strain is strictly specific of a plant family and obtained a value for p of 0.0051. This result indicates that the probability of getting such a plant specific distribution by chance is exceptional, which constitutes a reasonable evidence that the events of horizontal transmission are not occurring randomly, but are rather influenced by the host plant.

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Discussion

Distribution and endosymbiotic origin of thelytoky in phytophagous Megastigmus species A few hundred species in all of the major animal groups are characterized by thelytokous parthenogenesis; its patchy taxonomic distribution in Hymenoptera is consistent with several independent evolutionary origins from ancestral arrhenotokous species (Cook, 1993). The relatively high prevalence of thelytoky in Hymenoptera might reflect that some mechanisms of their haplodiploid sex determination are rather easily redirected to thelytoky by mutations (Lattorff et al. 2005) or bacteria (Cordaux et al., 2011). Our literature review of parthenogenesis in the Megastigmus genus estimated the prevalence of thelytoky to 15%, which supported the assumption that arrhenotoky is the dominant and ancestral form of parthenogenesis in these phytophagous wasps. Thelytoky was unambiguously described in species exploiting diverse habitats, although it appears to occur in only three out of the ten host plant families currently known to host Megastigmus species in both gymnosperms and angiosperms (Pinaceae, and Rosaceae and Anacardiaceae, respectively). Interestingly, several thelytokous species share congeneric host plants (e.g. on the Rosa genus), but each groups of thelytokous species associated with Pinaceae, Rosaceae or Anacardiaceae remains confined at the host family taxonomic level. This observation supports the potential for intimate ecological connections within a given host plant family, but not between different host plant families. Thelytoky may be under the control of the insect itself or its endosymbionts (Stouthamer et al, 1990; Vavre et al., 2004). Strict association between thelytoky and Wolbachia infection and reversibility of thelytoky through antibiotic treatments provide a particularly strong support for an endosymbiotic origin of thelytoky in Megastigmus wasps. Within the Hymenoptera, Wolbachia-induced thelytoky is found in three super-families: the Cynipoidea

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(Rokas *et al.*, 2002), the Braconidae (Kremer *et al.*, 2009) and the Chalcidoidea (Stouthamer 1997), within which only entomophagous families were concerned. To our knowledge, we document here the first case of *Wolbachia*-induced thelytoky in phytophagous chalcids.

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Host plant specialization matters in the spread of thelytoky

The phylogenetic tree of the Megastigmus genus including all main ecological groups (angiosperm- and gymnosperm-specialized seed wasps) indicated that thelytokous Megastigmus species do not form a monophyletic group, but occur sporadically on the COI phylogenetic tree (Figure 1). This pattern is consistent with the hypothesis of independent evolutionary origins of thelytokous species from ancestral arrhenotokous species (Cook, 1993), and thus that transitions from arrhenotoky to thelytoky have repeatedly occurred. While transition from arrhenotoky to thelytoky is possible, the reverse is much more constrained and even impossible after sexual traits have decayed (Jeong and Stouthamer 2005; Kremer *et al.*, 2009). The phylogenetic tree obtained on the concatenated MLST dataset provided critical insights on the history of Wolbachia infection in Megastigmus. Three homogeneous lineages of Wolbachia (WA1, WA2 and WB) were identified and shown to be associated with the host plant families of their hosts (Figure 1). Although both lineages were found closely related, WA1 infected exclusively thelytokous wasp species exploiting the Rosaceae while WA2 was detected only in those exploiting the Pinaceae. The third lineage WB was restricted to the only currently known thelytokous wasp species exploiting Anacardiaceae (M. pistaciae). More interestingly, there was no statistical support for an association between Wolbachia and host lineages. For example, WA1 is found in five species developing on Rosaceae, but belonging to highly divergent clades (10% COI). Similarly, WA2 is only found in four species exploiting Pinaceae, despite a divergence of as high as 5-6% on COI among these species. On the opposite, we found a non-random association between Wolbachia strains and the host plant used by the insects. Altogether, this led us to postulate on the one hand that thelytoky may have spread across the Megastigmus genus through horizontal transmission of Wolbachia among wasp species, and on the other hand that these transfers occurred preferentially between species exploiting similar host plants. Within-community horizontal transmission of Wolbachia is likely to result from ecologically mediated pathways such as host-parasitoid associations (Vavre et al. 1999; Huigens et al., 2004). Shared feeding and breeding sites may also act as a route through which Wolbachia can be transmitted from one host species to another (Sintupachee et al. 2006). Primary infection by Wolbachia has to face two main filters (Vavre et al. 2003). First, Wolbachia must come into contact with the recipient species (i.e. has to pass the encounter filter). Second, Wolbachia has to evade the host immune system and to develop in the new host (i.e. has to pass the compatibility filter), which is facilitated by the relatedness of the donor and recipient species. Community structure may affect both filters and thus impact the epidemiology of Wolbachia across species, as recently showed theoretically using smallworld networks (Zug et al., 2012). Whether communities are composed of generalist or specialist species, and of distantly or closely related species will clearly impact these two filters. On the one hand, interspecific horizontal transmission of Wolbachia may be favoured in particular habitats that are attractive for a wide set of generalist host species, which will simultaneously feed and develop on it and/or which can be connected by shared parasitoids (Stahlhut et al., 2010). The situation encountered in Trichogramma may be representative of

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such case with the existence of a specialized PI-Wolbachia clade infecting these generalist wasps, but within which horizontal transmission is rampant (Schilthuizen and Stouthamer, 1997). On the other hand, intimate interspecific connections between host species may also arise from host plant specialization, which may favour Wolbachia transmission due to narrow shared ecological niches. This situation opens widely the encounter filter, but also restricts the host spectrum to the few species exploiting this particular resource. When species within the community are moreover closely related species exploiting the same resource, the compatibility filter will also be open and facilitates symbiont infection. This latter situation is probably the one encountered in the seed-feeding Megastigmus analysed here, revealing two essential features of the spread of thelytoky in the context of insect ecological specialization. First, the COI reconstruction of the present 25 Megastigmus species strongly confirmed host plant specialization in this genus (Figure 1). Within the species on angiosperms, the clade of species attacking Anacardiaceae first diverged. Within one of the sister groups, wasp species exploiting gymnosperms form a monophyletic group and even show specialization, as depicted by the monophyletic group developing on the Pinaceae (except for M. thyoides). The existence of a common Wolbachia lineage in all wasps exploiting the Rosaceae (WA1) and another one in all those developing on the Pinaceae (WA2) suggests that host ecological specialization can promote the spread of thelytoky through seed use. All Megastigmus females oviposit during a rather narrow period of the development of the host reproductive structures and progeny develop exclusively within seeds at a final density of one larva per seed, while several eggs can be found within the same seed (Turgeon et al., 1994: Rouault et al. 2004; T. Boivin, personal observation). On both wild roses and Pinaceous trees, many

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Megastigmus species show some restricted behavioral plasticity allowing shifts onto different congeneric host species or even onto a new host genus (Grissell 1999; Auger-Rozenberg & Roques 2012). The possibility of short-range host shifts may have favoured novel wasp assemblages and novel opportunities for them to interact and potentially exchange symbionts. Although shared parasitoids between phytophagous Megastigmus spp. can not be ruled out (Mailleux et al., 2008; M.-A. Auger-Rozenberg, personal observation), host-parasitoid associations in this genus remain too poorly documented to formally invoke this Wolbachia horizontal route at this point. We rather suggest that if Wolbachia infection occurs mainly via ingestion during the early larval stages, both seed tissues and/or cannibalism by larvae competing for the seed tissues may facilitate interspecific horizontal transfers of the endosymbiont. A similar pattern of endosymbionts sharing among sibling species competing for a common resource have recently been shown in weevils (Merville et al. 2013). Another key result of this study is that Megastigmus ecological specialization at the host plant family level probably constrained the invasion of Wolbachia lineages throughout the whole host genus. Indeed, diverging strategies in the use of angiosperms and gymnosperms may be an essential feature of the strict host radiation depicted here in the evolutionary history of Megastigmus. For this reason, intimate ecological connections promoting horizontal transfers of Wolbachia (see above) are thus unlikely to arise at this community scale. More interestingly, thelytoky appears to have still not invaded wasps developing on both the Cupressaceae (Fig. 1, Table S1). According to Auger-Rozenberg et al. (2006), wasps developing on the Cupressaceae exhibit a higher level of host specificity, because they seem host species specific, while wasps on Pinaceae or Rosaceae are specialized to particular host

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genera, but frequently attack several congeneric species if they occur in sympatry. This is consistent with a taxonomic radiation following initial host adaptation, which might have constituted an efficient barrier to the spread of thelytoky in this group, due to unlikely opportunities for interspecific horizontal transfers of *Wolbachia*.

Altogether, our results show that ecological specialization can be a driving force of the spread of endosymbiotic thelytoky, but also a constraint. This further reinforces the hypothesis that community structure of insects is a major driver of the epidemiology of endosymbionts (Ferrari and Vavre, 2011) and that competitive interactions among closely related species may facilitate horizontal transmission (Merville *et al.*, 2013).

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708 709 **Data accessibility** 710 Wolbachia and wasp DNA sequences: GenBank accession numbers KF531859-KF531890 and 711 KF531891-KF531900, and KF531833-KF531858, respectively. 712 713 714 **Author contributions** 715 M.-A.A.R., F.V. and T.B. conceived and designed the study. M.-A.A.R, A.R. and T.B. acquired 716 the samples and provided funding support. E.M., C.G., J.-N., M.-A.A.R and T.B. performed the 717 experiments and produced the data. H.H., M.-A.A.R, F.V. and T.B. analyzed the data and 718 wrote the paper. All authors have checked and approved the final version of the manuscript. 719 720 **Figure Legends** 721 Figure 1. Bayesian likelihood inference phylogeny based on COI sequences in seed-722 specialized wasps of the Megastigmus genus (26 taxa, 962 bp). Torymus azureus was used as 723 an outgroup. Posterior probability values are indicated at each node. Host plant families are 724 indicated at each branch forming a monophyletic group (excepted for the Cupressacae). 725 Anarcadiaceae and Rosaceae are angiosperms, and Pinaceae, Cupressaceae and Taxodiaceae are gymnosperms. Megastigmus species infected by Wolbachia are indicated by arrowheads 726 727 followed by the name of the Wolbachia lineage. 728 729 Figure 2. Phylogenetic placement of the Wolbachia strains infecting the seed-specialized

wasps Megastigmus spp. among other Wolbachia belonging to A and B supergroups.

Bayesian likelihood inference phylogenies are shown, while maximum likelihood analyses gave substantially the same results. *Wolbachia* sequences are labelled with the name of their host. *Wolbachia* of *Brugia malayi* was used as an outgroup. Posterior probability values are indicated at each node. This phylogeny is based on concatenated sequences data set for the four MLST loci *coxA*, *gatB*, *ftsZ* and *hcpA* (34 strains, 1650 bp).

Supplementary material

Figure S1. Phylogenetic placement of *Wolbachia* infecting the seed-specialized wasps *Megastigmus* spp., among other *Wolbachia* strains belonging to A and B supergroups, based on the sequences of each of the four MLST genes used in this study (*coxA*, *gatB*, *ftsZ* and *hcpA*).

Figure S2. Phylogenetic placement of the *Wolbachia* strains infecting the seed-specialized wasps *Megastigmus* spp., among other *Wolbachia* strains belonging to A and B supergroups, based on wsp sequences of 49 strains (530 bp).

Table S1. Literature review of the distribution of arrhenotoky, thelytoky and host-plant specialization among the seed-feeding wasps of the *Megastigmus* genus.

Table S2. The primer pairs used for endosymbiont PCR screening in *Megastigmus* spp.

Table 1. Collection data for the specimens used in the study of 15 arrhenotokous and all the currently known thelytokous species (N=10) in the *Megastigmus* genus. A: arrhenotokous parthenogenesis. T: thelytokous parthenogenesis. Note that *Torymus azureus* was used as an outgroup for building the COI phylogenetic tree of the *Megastigmus* genus presented in Fig. 1. Specimens collected outside of their native range belonged to invasive populations.

Species name	Reproduction	Host-plant group	Host-plant family	Host-plant species	Collection site	Native range
M. aculeatus	Т	Angiosperm	Rosaceae	Rosa rugosa	Krasnoyarsk, Russia	Palearctic
M. aculeatus nigroflavus	Т	Angiosperm	Rosaceae	Rosa sp.	Iowa, USA	Nearctic
M. borriesi	Т	Gymnosperm	Pinaceae	Abies koreana	Rold skov, Denmark	East-Asia
M. brevicaudis	Т	Angiosperm	Rosaceae	Sorbus sp.	Ojcow, Poland	Palearctic
M. pictus	Т	Gymnosperm	Pinaceae	Larix gmelini	Loiret, France	Palearctic
M. pinsapinis	Т	Gymnosperm	Pinaceae	Cedrus atlantica	Vaucluse, France	Palearctic
M. pistaciae	Т	Angiosperm	Anacardiaceae	Pistacia terebenthus	Exocori, Greece	Palearctic
M. rosae	Т	Angiosperm	Rosaceae	Rosa sp.	Hautes-Alpes, France	Palearctic
M. rosae alba	Т	Angiosperm	Rosaceae	Rosa majalis	Vilnius, Lithuania	Palearctic
M. suspectus	Т	Gymnosperm	Pinaceae	Abies alba	Gard, France	Palearctic
M. amicorum	А	Gymnosperm	Cupressaceae	Juniperus phoenicea	Ericera, Portugal	Palearctic
M. atedius	Α	Gymnosperm	Pinaceae	Picea sp.	British Columbia, Canada	Nearctic
M. atlanticus	Α	Gymnosperm	Cupressaceae	Cupressus atlantica	Idni, Morocco	Afro-tropical
M. bipunctatus	Α	Gymnosperm	Cupressaceae	Juniperus communis	Hautes-Alpes, France	Palearctic
M. cryptomeriae	Α	Gymnosperm	Cupressaceae	Cryptomeria fortunei	Zeihjang, China	East-Asia
M. hoffmeyeri	Α	Gymnosperm	Pinaceae	Tsuga canadensis	Ontario, Canada	Nearctic
M. lasiocarpae	Α	Gymnosperm	Pinaceae	Abies amabilis	British Columbia, Canada	Nearctic
M. nigrovariegatus	Α	Angiosperm	Rosaceae	Rosa sp.	British Columbia, Canada	Nearctic
M. pinus	Α	Gymnosperm	Pinaceae	Abies procera	California, USA	Nearctic
M. rafni	Α	Gymnosperm	Pinaceae	Abies alba	Aude, France	Nearctic
M. schimitscheki	Α	Gymnosperm	Pinaceae	Cedrus atlantica	Luberon, France	Palearctic
M. spermotrophus	Α	Gymnosperm	Pinaceae	Pseudotsuga mensiezii	California, USA	Nearctic

M. thyoides	Α	Gymnosperm	Cupressaceae	Chamaecyparis sp.	North Carolina, USA	Nearctic
M. transvaalensis	Α	Angiosperm	Anacardiaceae	Schinus molle	Marrakech, Morocco	Afro-tropical
M. tsugae	Α	Gymnosperm	Pinaceae	Tsuga heterophylla	British Columbia, Canada	Nearctic
Torymus azureus	Α	Gymnosperm	Pinaceae	Picea abies	Suchora, Poland	Palearctic

Table 2. *Megastigmus* species screened for bacterial infection (+) or non-infection (-) using PCR with *Wolbachia, Cardinium, Arsenophonus* and *Rickettsia*-specific primers. Each female was screened for all target endosymbionts. The mitochondrial cytochrome oxidase-I (COI) gene of the *Megastigmus* host was amplified when no bacterial-specific primers yielded PCR products to test for correct DNA extraction in the procedure. N: number of females tested for infection. +: amplification of the primer. -: no amplification of the primer tested. NA: non available data.

Parthenogenesis and species (N)	Wolbachia (infection %)	Cardinium	Arsenophonus	Rickettsia	Arthropod COI
Thelytoky					
M. aculeatus (43)	+ (100)	-	-	-	
M. aculeatus nigroflavus (21)	+ (100)	-	-	-	
M. brevicaudis (1)	+ (100)	NA	NA	NA	
M. borriesi (1)	+ (100)	NA	NA	NA	
M. pictus (30)	+ (100)	-	-	-	
M. pinsapinis (60)	+ (100)	-	-	-	
M. pistaciae (30)	+ (100)	-	-	-	
M. rosae (30)	+ (100)	-	-	-	
M. rosae alba (5)	+ (100)	-	-	-	
M. suspectus (65)	+ (100)	-	-	-	
Arrhenotoky					
M. amicorum (20)	-	-	-	-	+
M. atlanticus (20)	-	-	-	-	+
M. cryptomeriae (20)	-	-	-	-	+
M. pinus (20)	-	-	-	-	+
M. rafni (30)	-	-	-	-	+
M. spermotrophus (30)	-	-	-	-	+
M. schimitscheki (30)	-	-	-	-	+
M. transvaalensis (20)	-	-	-	-	+

Table 3. Wolbachia infection rates in control and tetracycline-treated females of M. pinsapinis (thelytokous) and M. schimitscheki (arrhenotokous), and brood size and proportions of males produced by control and tetracycline-treated females of these species.

Wolbachia infection			Progeny produced by treated females			
Species	Treatment (N*)	% of <i>wsp</i> positives**	Number of mothers	Mean brood size***	Mean % of males	
M. pinsapinis	control (44)	100	44	21.4 ± 5.3^{a}	0	
	tetracycline (42)	4.7	42	18.7 ± 7.8°	92 ± 1.1	
M. schimitscheki	control (30)	0	30	30.3 ± 4.5^{a}	100	
	tetracycline (28)	0	28	32.1 ± 7.5^{a}	100	

^{*}Number of treated females

^{***}Infection rates were determined by PCR using *Wolbachia*-specific primers of the *wsp* gene.

***For each species, similar letters indicate that the mean number of larvae (<u>+</u>SE) produced do not differ significantly using a Kruskal-Wallis test (P<0.05).

Figure 1

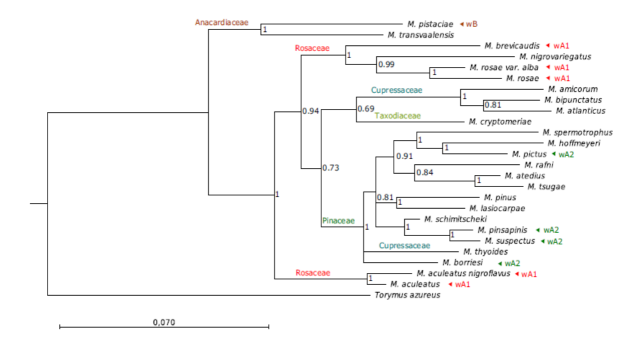


Figure 2

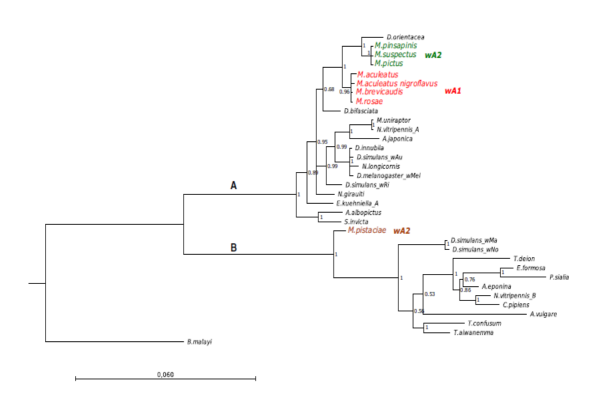


Table S1. Distribution of arrhenotoky, thelytoky and host-plant specialization among the seed-feeding wasps of the *Megastigmus* genus. Data were compiled from reviews of Grissell (1999), Roques and Skrzypczynska (2003), Roques et al. (2003) and Auger-Rozenberg et al. (2006). Species in bold were specifically studied in this paper. A: arrhenotokous parthenogenesis. T: thelytokous parthenogenesis. * Ratio of males to females; balanced: consideration of similar frequencies of females and males by authors; NA: no specific mention on sex ratio by authors but numbers of males mentioned in samples (thelytoky being associated with "males unknown" or "males are scarce" specific mentions).

Host-plant taxon	Species	Reproduction	Sex ratio*	Main host-plant genus	Native area
Gymnosperms					
Cupressaceae	M. chamaecyparidis	A	balanced	Chamaecyparis	Palaearctic
	M. thyoides	A	balanced	Chamaecyparis	Nearctic
	M. atlanticus	A	balanced	Cupressus	Palaearctic
	M. carinus	A	NA	Cupressus	Palaearctic
	M. cupressi	A	NA	Cupressus	Oriental
	M. duclouxiana	A	NA	Cupressus	Palaearctic
	M. watchli	A	0.5-1.7	Cupressus	Palaearctic
	M. amicorum	A	balanced	Juniperus	Palaearctic
	M. bipunctatus	A	balanced	Juniperus	Palaearctic
	M. certus	A	NA	Juniperus	Palaearctic
	M. formosana	A	balanced	Juniperus	East-Asia
	M. fidus	A	balanced	Juniperus	Palaearctic
	M. gravis	A	balanced	Juniperus	Palaearctic
	M. juniperi	A	NA	Juniperus	Palaearctic
	M. pingii	A	NA	Juniperus	East-Asia
	M. procerae	A	NA	Juniperus	Palaearctic
i i i	M. rigidae	A	NA	Juniperus	Palaearctic
	M. sabinae	A	balanced	Juniperus	Palaearctic
	M. somaliensis	A	NA	Juniperus	Afrotropical
	M. thuriferana	A	balanced	Juniperus	Palaearctic
	M. validus	A	balanced	Juniperus	Palaearctic
	M. thuyopsis	A	NA	Thuyopsis	Palaearctic
<u>Pinaceae</u>	M. firmae	A	NA	Abies	Palaearctic
	M. lasiocarpae	A	balanced	Abies	Nearctic
	M. milleri	A	balanced	Abies	Nearctic
	M. pinus	A	0.4-0.5	Abies	Nearctic
	M. rafni	A	0.4	Abies	Nearctic
	M. specularis	A	balanced	Abies	Nearctic
	M. schimitscheki	A	0.4-0.5	Cedrus	Palaearctic
	M. laricis	A	balanced	Larix	Nearctic
	M. atedius	A	0.19-0.66	Picea	Palaearctic
	M. ezomatsuanus	A	0.5	Picea	Palaearctic
	M. likiangensis	A	NA	Picea	East-Asia
	M. strobilobius	A	0.4-0.6	Picea	Palaearctic
	M. albifrons	A	balanced	Pinus	Nearctic
	M. strobiformis	A	NA	Pinus	Nearctic

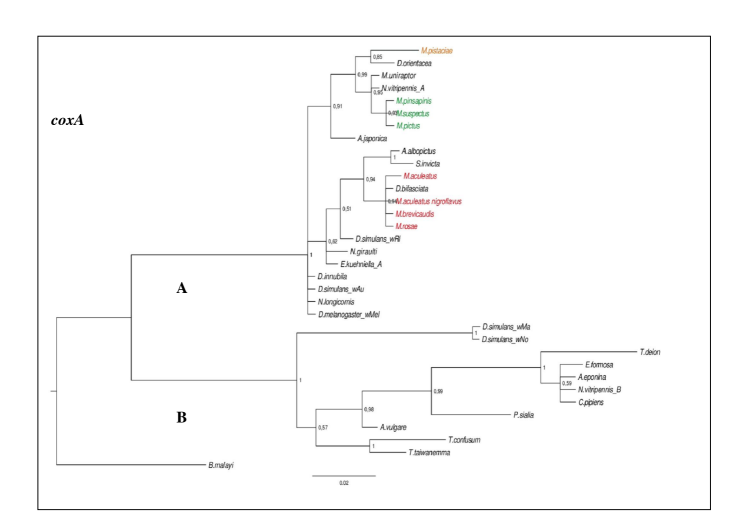
	M. pseudotsugae	A	NA	Pseudotsuga	East-Asia
	M. spermotrophus	A	balanced	Pseudotsuga	Nearctic
	M. hoffmeyeri	A	balanced	Tsuga	Nearctic
	M. tsugae	A	balanced	Tsuga	Nearctic
	M. tsugaphilus	A	NA	Tsuga	Palaearctic
	M. borriesi	T	♂<0.1%	Abies	East-Asia
	M. suspectus	T	♂<0.2%	Abies	Palaearctic
	M. pinsapinis	T	♂<0.1%	Cedrus	Palaearctic
	M. pictus	T	♂<1%	Larix	Palaearctic
<u>Taxodiaceae</u>	M. cryptomeriae	A	balanced	Cryptomeria	East-Asia
Angiosperms					
Anacardiaceae	M. thomseni	A	NA	Heeria	Afrotropical
	M. ozoroae	A	NA	Ozoroae	Afrotropical
	M. rhusi	A	NA	Rhus	Afrotropical
	M. transvaalensis	A	0.5	Schinus	Afrotropical
	M. pistaciae	T	♂<0.1%	Pistacia	Palaearctic
<u>Fabaceae</u>	$M.\ albizziae$	A	NA	Albizia	Oriental
<u>Hamameliaceae</u>	M. distylii	A	NA	Distylium	Palaearctic
Myrtaceae	$M.\ ophelinii$	A	NA	Eucalyptus	Australasia
Proteaceae	M. hakeae	A	NA	Hakea	Australasia
Rhamnaceae	M. helianthae	A	NA	Helinius	Afrotropical
Rosaceae	$M.\ amelanchier is$	A	NA	Amelanchier	Nearctic
	M. cotoneastri	A	NA	Cotoneaster	Palaearctic
	M. mali	A	NA	Malus	Palaearctic
	M. pourthiaceae	A	0.6	Photinia	Palaearctic
	M. physocarpi	A	NA	Physocarpus	Nearctic
	M.fangii	A	NA	Rosa	East-Asia
	M. nigrovariegatus	A	balanced	Rosa	Nearctic
	M. yunnanensis	A	NA	Rosa	East-Asia
	M. aculeatus	T	∂<4%	Rosa	Palaearctic
	M. aculeatus nigroflavus	T	♂<1%	Rosa	Palaearctic
	M. rosae	T	♂<1%	Rosa	Palaearctic
	M. rosae var. alba	T	♂<1%	Rosa	Palaearctic
-	M. brevicaudis	T	♂<1%	Sorbus	Palaearctic

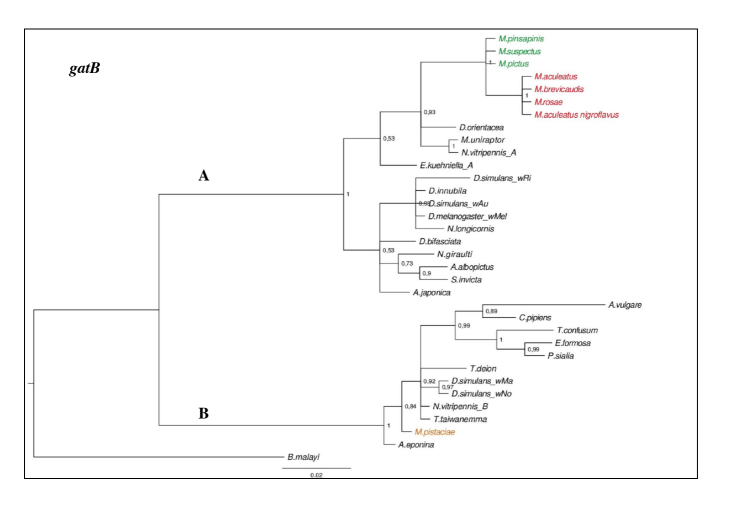
Table S2. Primer pairs used for endosymbiont PCR screening in Megastigmus spp.

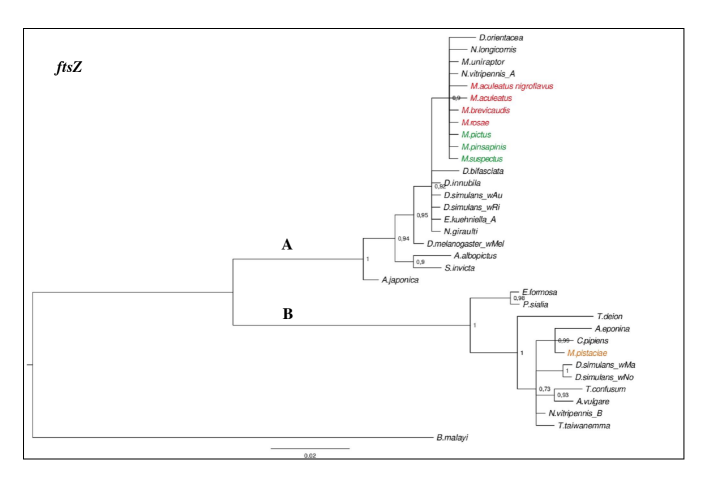
Primer pair (5'-3')	Target group	Target gene	Annealing	
		(fragment size)	temperature (°C)	
wsp81F TGGTCCAATAAGTGATGAAGAAAC ^a	Wolbachia pipientis	wsp		
wsp691R AAAAATTAAACGCTACTCCA ^a		(610 bp)	59	
Ch-F TACTGTAAGAATAAGCACCGGC ^b	Cardinium sp.	16S rRNA	57	
Ch-R GTGGATCACTTAACGCTTTCG ^b		(~900 bp)	31	
Ars23S-1 CGTTTGATGAATTCATAG TCAAA ^c	Arsenophonus nasoniae	23S rDNA	60	
Ars23S-2 GGTCCTCCAGTTAGTGTTACCCAAC ^c		(650 bp)	00	
27f GAGAGTTTGATCCTGGCTCAG ^d	Rickettsia sp.	16S rRNA	50	
1495r CTACGGCTACCTTGTTACGA ^d		(1500 bp)	50	
LCO1490 GGTCAACAAATCATAAAGATATTGG ^e	Arthropods	COI	48	
HCO2198 TAAACTTCAGGGTGACCAAAAAATCA ^e		(658 bp)	40	

^aBraig *et al*. (1998) ^bZchori-Fein and Perlman (2004) ^cThao and Baumann (2004) ^dHagimori *et al*. (2006) ^eFolmer *et al*. (1994)

Figure S1. Phylogenetic placement of *Wolbachia* infecting the seed-specialized wasps *Megastigmus* spp., among other *Wolbachia* strains belonging to A and B supergroups, based on the sequences of each of the four MLST genes used in this study (*coxA*, *gatB*, *ftsZ* and *hcpA*). Bayesian likelihood inference phylogenies are shown, while maximum likelihood analyses gave substantially the same results. *Wolbachia* sequences are labelled with the name of their host. *Wolbachia* of *Brugia malayi* was used as an outgroup. Posterior probability values are indicated at each node. Each tree represents phylogenetic reconstruction based on 34 Wolbachia strains for *coxA* (402 bp), *gatB* (369 bp), *ftsZ* (435 bp) and *hcpA* (444 bp). Three lineages are revealed (green, red and light brown), two belonging to the A supergroup while the position of the *Wolbachia* infecting *M. pistaciae* suggests a recombination event between A and B *Wolbachia* supergroups. Indeed, this strain belongs to the B supergroup based on all sequences genes except for *coxA* for which it is placed in the B supergroup.







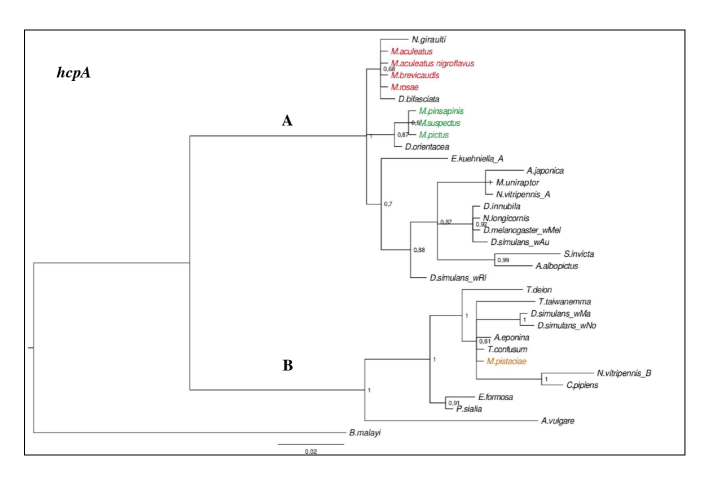
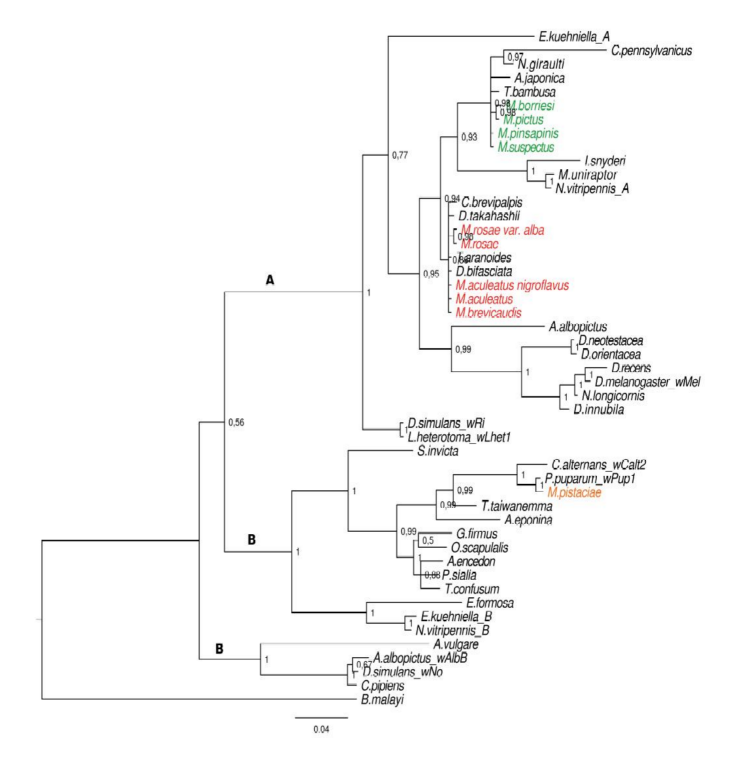


Figure S2. Phylogenetic placement of the *Wolbachia* strains infecting the seed-specialized wasps *Megastigmus* spp., among other *Wolbachia* strains belonging to A and B supergroups, based on wsp sequences of 49 strains (530 bp). Bayesian likelihood inference phylogenies are shown, while maximum likelihood analyses gave substantially the same results. *Wolbachia* sequences are labelled with the name of their host. *Wolbachia* of *Brugia malayi* was used as an outgroup. Posterior probability values are indicated at each node.



References

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