Host genotype-specific microbiota do not influence the susceptibility of *D. magna* to a bacterial pathogen

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Host-associated microbiota have been claimed to play a role in hosts’ responses to parasitic infections, often protecting the hosts from infection. We tested for such a role in the crustacean *Daphnia* and the parasitic bacterium *Pasteuria ramosa*, a widely used model system for host-parasite interactions. We first determined the infection phenotype (i.e., resistotype) of eight clonal *D. magna* genotypes against four strains of *P. ramosa* by attachment test, followed by 16 S rDNA amplicon sequencing to determine if their genotype or their parasite resistotype influences the composition of their microbiome. We then reciprocally transplanted the microbiota of two host genotypes with opposite resistotypes to four *P. ramosa* isolates, followed by a reassessment of their resistotype after transplantation. We found significant differences in microbiome composition and structure between *Daphnia* genotypes and between *Daphnia* resistotypes to specific *P. ramosa* strains. Reciprocal microbiota exchange or making the *Daphnia* hosts bacteria-free, however, did not influence the resistotypes of the hosts. Thus, in contrary to what has been observed in some taxa, our results suggest that *D. magna* susceptibility to *P. ramosa* is strongly dictated by the genetic differences of the hosts and is still dependent on *Daphnia*’s first line of immune defense against the esophageal attachment of *P. ramosa*, which appears to be uninfluenced by the host’s microbiota.

Until a few years ago, conventional wisdom stated that differences among host populations in resistance to pathogens are largely explained by host genetic differences1–3 with environmental factors playing a subordinate role. Recently, microbiota have been added to the list of non-genetic factors that contribute to host resistance against parasites4–7. The specificity of infection from a gut pathogen in honeybees, for instance, is purportedly influenced by microbiome composition, and hints at the protective role of the microbiome in counteracting parasite infection8,9. Contrastingly, reports that the microbiome can also mediate host susceptibility to diseases have been gaining a lot of attraction in the biomedical sciences9–11. The beneficial role of the microbiome to host health has been overtly emphasized in recent years, but their possible influence on host pathogen resistance comes from a handful of investigations from a few host taxa so far. Therefore, this emerging concept requires more evidence before a solid generalization across taxa can be made.

Next generation sequencing has allowed researchers to characterize the microbiome composition of many organisms, resulting in increasingly more complex patterns. Environmental conditions such as diet, can explain variation in microbiome composition between individuals12,13, but species-specificity of microbiome composition is still observed, and differences between groups of organisms can still be explained in part by their phylogenetic position14,15. Recently, the host genotype in mice was reported to influence gut microbial composition13,16, with some members of microbiome being more heritable than others17. Moreover, genetic variation in immune-related pathways between human hosts is found to be associated with microbiome composition, lending support to the idea that aside from environmental factors, immune traits can influence microbiome composition15,18,19. However, it is not known whether it is the differences in overall host genetic background or the differences in specific immune traits that influences the microbiome composition in a given host taxa. Teasing apart these factors would allow to understand the role that the microbiome plays on immune traits and pathogen resistance. This would be especially interesting among invertebrates that lack adaptive immunity.

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We used the *Daphnia-Pasteuria* host–parasite system to address these questions. Genetic variation is observed among natural genotypes of *D. magna*, and quantitative trait locus (QTL) approaches have been developed to address questions on the genetic basis of parasite resistance\(^{20,21}\). Many microbial parasites infect *Daphnia*, and one of them, *P. ramosa*, a gram-positive bacterium of the class *Bacilli*, was shown to display a co-evolutionary arms race with its hosts\(^{22,23}\). Furthermore, the role of microbiome on *Daphnia* fitness has been established and the experimental system can be manipulated such that the production of gnotobiotic *Daphnia* is possible\(^{24}\). These characteristics make *Daphnia* a good model to disentangle the contribution of host genetic background and microbiome composition, and to determine if the microbiome plays a role in *Daphnia-P. ramosa* interactions.

The interaction of *P. ramosa* with *D. magna* reveals a very high genetic specificity, visible as strong host genotype by parasite genotype interaction with an almost perfect binary infection outcome\(^{25,26}\). When a potential host encounters the parasite spores, the spores are activated followed by attachment to the esophagus wall. After successful attachment, the spores penetrate the foregut epithelium into the host body cavity and start proliferating in the haemolymph. Eventually, the host dies and releases *Pasteuria* spores in the environment\(^{27}\). Infections are highly virulent with severe reduction in host fecundity and lifespan. Most variation in infection success is explained by the first line of defense, i.e., the ability of the host to prevent attachment to the esophagus, the entry to *Daphnia’s* foregut\(^{25}\). Susceptibility to the host’s first line of defense can be assessed with an esophagus attachment test, during which fluorescently labeled *P. ramosa* spores are observed while attaching to the esophagus wall of the host. No esophageal attachment indicates resistance to the parasite\(^{25}\).

The main goal of this study is to determine if the microbiome plays a role in the resistance of *D. magna* to *P. ramosa*. We evaluated this by determining the infection phenotype of 8 *Daphnia* genotypes (=clones) to four *P. ramosa* strains, followed by comparison of their microbiome structure and composition based on genotype and resistance phenotype (termed resistotype in this study) to four *Pasteuria* strains. We then performed a reciprocal transplant experiment where we swapped the microbiome of resistant and susceptible *D. magna* genotypes and tested for resistance to one *P. ramosa* strain. Furthermore, we compared the resistotype of these *D. magna* genotypes when they were bacteria-free. We found that *D. magna* genotypes vary in microbiome composition even after years of culturing them under the same environmental conditions, and that statistical analysis indicated that their microbiome composition can potentially be structured by their resistotype. However, replacing the microbiome of the hosts or making the hosts bacteria-free have not altered their resistance or susceptibility to a *P. ramosa* pathogen.

### Results

#### Resistotype of *D. magna* genotypes.

Four *Pasteuria* strains were tested for attachment in the esophagus to identify *Daphnia* resistotypes (*n* = 31) (Table 1). Most *Daphnia* genotypes tested were resistant to Pr_C1 and Pr_P20, while half of the genotypes tested were susceptible to Pr_C19 and Pr_P15. Two genotypes were resistant to all four *Pasteuria* pathogens (clones Belgium_1, BE-OHZ-M5 and Germany, DE-K35-Inh1), while the *Daphnia* genotype from Hungary (HU-H0-2) was susceptible to all four pathogens. The Israel (IL-M1-1) clone died in culture and was not tested for Pr_P15 and Pr_P20.

### Common microbiome among *D. magna* genotypes.

We sequenced the V3 to V5 hypervariable region of the 16S rDNA of 31 samples belonging to eight *D. magna* genotypes including one positive control (mock community) and analyzed 432, 318 sequences after denoising (3893–47570 per sample). Out of 28 genera used in the DNA mock community, 21 were represented at relative abundances similar to the DNA proportions used for sequencing. The use of 97% cut-off in picking Operational Taxonomic Units (OTUs) only gave resolution at the genus or family level identification and not at the strain level (see Supplementary Fig. S1). Strains belonging to *Actinobacteria* were not seen in the DNA mock community sequences but the V3-V5 16S rDNA primers were able to capture the majority of the bacterial groups. *Actinobacteria* sequences were however seen in 4 (out of 31) *D. magna* samples albeit very rare and low in abundances. Qi *et al.*\(^{28}\) have seen *Actinobacteria* sequences in their metagenomics survey of *Daphnia* microbiome and found these to be abundant in *D. pulicaria* but very rare in *D. magna* and *D. pulex*. The DNA mock community sequences were not included in the downstream analysis.

A total of 6548 OTUs (≥97% ID) were represented in the 31 *D. magna* samples but after removing singletons and chloroplast sequences, only 455 OTUs remained to be used for the analysis. Of these OTUs, 13 were present

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**Table 1.** Infection phenotype (i.e., resistotype) of eight *D. magna* genotypes to four *P. ramosa* pathogenic strains tested via attachment testing to the esophagus or hindgut.

| *Daphnia* ID | Pr_C1 | Pr_C19 | Pr_P15 | Pr_P20 | Resistotype | Country Origin | Location | Latitude-Longitude | Life Stage | Collected |
|-------------|-------|--------|--------|--------|-------------|---------------|----------|-------------------|------------|-----------|
| BE-OHZ-M5   | R     | R      | R      | R      | RRRR       | Belgium_1     | Oud Heverlee, Belgium | 50° 50′ 14″, 4° 39′ 48″ | Hatched from ephippium |
| BE-OHZ-M10  | S     | S      | R      | R      | SSRR       | Belgium_2     | Oud Heverlee, Belgium | 50° 50′ 14″, 4° 39′ 48″ | Hatched from ephippium |
| QTL-IXF-1   | R     | S      | R      | R      | RSSR       | Basel, Switzerland | 47° 33′ 37″, 7° 34′ 57″ | Hatched from ephippium |
| FI-Xinh3    | R     | S      | S      | R      | RSSR       | Finland       | Tvärminne, Finland | 59° 50′ 31″, 23° 12′ 6″ | Hatched from ephippium |
| DE-K35-Inh1 | R     | R      | R      | R      | RRRR       | Germany       | Ismaninger Teiche, Germany | 48° 12′ 24″, 11° 42′ 35″ | Hatched from ephippium |
| HU-HO-2     | S     | S      | S      | S      | SSSS       | Hungary       | Kikungsagi-nemzeti Park, Hungary | 46° 50′ 14″, 19° 11′ 41″ | Field-collected individual |
| IR-GG1-1    | R     | S      | R      | R      | RSRR       | Iran          | Lake Guru-gol, Iran | 37° 55′ 3.8″, 46° 42′ 7.8″ | Hatched from ephippium |
| IL-M1-1     | R     | R      | R      | R      | RR         | Israel        | Mamilla Pond, Jerusalem | 31° 42′ 52″, 35° 3′ 3.5″ | Hatched from ephippium |
in 23 (out of 35) of the *D. magna* samples comprising genera and orders at different relative proportions (Fig. 1). The abundant microbiome populations were dominated by bacteria having homologous 16S rDNA sequences to *Limonohabitans* (Beta-proteobacteria, 22–66% of the total counts), *Saprospirales* (Bacteroidetes, 9%–54%), and *Rhodobacter* (Alpha-proteobacteria, 1–22%). Bacterial sequences homologous to *Emticicia*, *Rudanella* and *Rutella* (Bacteroidetes, 0.5–3%); *Rhizobiales* (Alpha-proteobacteria, 0.2–7%); *Ideonella* (Beta-proteobacteria, 0.3–5%); *Halomonas* and *Sinobacteraceae* (Gamma-proteobacteria, ≤0.6%) were the other least abundant but common microbiome among *D. magna*.

Bacterial OTUs that were >1% in relative abundance but were only found in 15 out of 31 *D. magna* samples had homologous 16S rDNA sequences to the Flavobacteriales of the genera *Chryseobacterium* and *Flavobacterium* and the Sphingomonadales of the genera *Sphingomonas* and *Sphingobium*. Some bacteria such as *Hydrogenophaga*, *Caulobacter*, *Myxococcales*, *Pseudomonas*, *Pedobacter* and *Bacillus* were abundant in some *Daphnia* genotypes (Supplementary Fig. S2).

**Microbiome similarity by genotype and resistotype.** We plotted the principal coordinates of the Weighted and Unweighted UniFrac distances (Fig. 2a,b) to assess the similarity of the microbiome composition between eight *Daphnia* genotypes. A number of distinct clusters between genotypes were seen, suggesting variation in the microbiome composition between host genotypes. ANOSIM analysis supports statistically significant differences in bacterial community composition between host genotypes (Unweighted UniFrac, ANOSIM R = 0.50, p = 0.001). Similar results were obtained using ADONIS both from Unweighted (R² = 0.39, p = 0.001) and Weighted UniFrac Distances (ANOSIM R = 0.22, p = 0.016; ADONIS: R² = 0.42, p = 0.007) and Bray-Curtis Distance (ANOSIM R = 0.22, p = 0.004; ADONIS: R² = 0.39, p = 0.006). Dissimilarity in the microbiome composition corresponded with the geographic origin of *D. magna* genotypes. For instance, the Iran clone showed a more distinct microbiome composition than the Finland clone (Unweighted UniFrac distance = 0.67, p < 0.01), even though both were hatched from resting eggs in the lab. A similar observation was seen between the Germany and the Hungary clones (Unweighted UniFrac distance = 0.58, p = 0.052, black and red polygons, respectively, in Fig. 2a,b). Notably, these two *Daphnia* clones have opposite resistotypes to four *P. ramosa* strains. The F1 hybrid clone (F1_lab breed) had a microbial composition similar to its parents, the Finland and Germany clones (Unweighted UniFrac distance = 0.50 and 0.53 respectively, p > 0.05), although this sexual offspring has been hatched from ephippia.

To further determine if the resistotype of the *D. magna* clones against four *P. ramosa* strains can explain the variation in the microbiome composition besides the genotypic effect, we assessed the variation of the microbiome composition between *Daphnia* using their resistotype to individual *Pasteuria* strains as the constraining variable employing three distance matrices. Distance-based Redundancy Analyses (db-RDA) based on Unweighted UniFrac distance matrix showed significant clustering of *Daphnia* microbiome by resistotypes to Pr_C1, Pr_P15 and Pr_P20 (p values < 0.05, Fig. 3a,c and d) but not with Pr_C19 (p > 0.05, Fig. 3b). However, db-RDA performed on Weighted UniFrac distance matrix did not show significant clustering in the microbiome composition between *Daphnia* based on resistotypes to Pr_C1, Pr_C19, Pr_P15 and Pr_P20 (Weighted UniFrac: p = 0.07, 0.63, 0.11 and 0.07, respectively; Supplementary Fig. S3A–D). Similarly, db-RDA analysis performed using Bray-Curtis distance matrix also did not show significant clustering in the microbiome composition between *Daphnia* resistotypes to Pr_C1, Pr_C19 and Pr_P15 but showed significant microbiome clustering based on Pr_20 (Bray-Curtis:

**Figure 1.** Common microbiome found in 75% of *D. magna* samples (n = 31) originated from different populations. These thirteen genera/families also represent the abundant OTUs in the *D. magna* microbiome consortium. Bubbles represent OTU counts shown in relative abundance (%).
Cytophagaceae and mice, host genetics is reported to be a factor in shaping the composition and diversity of the microbiome17,29. Comamonadaceae a significantly lower proportion of bacteria from their infection phenotype regardless of their microbiota or even in the absence of microbiota. Daphnia of all Germany genotype replicates in all treatments. This suggests that the respective bacteria-free, if it contains its native microbiota or if the Pasteuria showed that Pr_C1 attached to the esophagus wall of all Hungarian genotype replicates, regardless if the animal is ota swapping will change the original resistotype of the respective Daphnia and German (RRRR) genotypes for the reverse microbiota transplant experiment to determine if microbiota swapping will change the original resistotype of the respective Daphnia genotype from Germany is significantly less species rich than the Hungary genotype based on Chao1 and number of observed OTUs metrics (Students T-test; p = 0.03 and 0.03, respectively). In terms of bacterial abundance, the highly-resistant German genotype contains a much higher proportion of bacteria from the family Saprospirales (p = 0.05), Sphingomonadaceae, Sphingobacteriaceae (p = 0.05) and Rhizobiaceae while it has a significantly lower proportion of bacteria from Comamonadaceae (p = 0.01), Rhodobacteraceae (p = 0.008) and Cytophagaceae (p = 0.06) (Fig. 4) than the highly-susceptible Hungarian genotype.

Microbiota swamping between D. magna of opposing resistotypes. We used Hungarian (SSSS) and German (RRRR) genotypes for the reverse microbiota transplant experiment to determine if microbiota swamping will change the original resistotype of the respective Daphnia clone. Pasteuria attachment tests showed that Pr_C1 attached to the esophagus wall of all Hungarian genotype replicates, regardless if the animal is bacteria-free, if it contains its native microbiota or if the Pasteuria-susceptible Daphnia contains the microbiota of a Pasteuria-resistant German genotype (Table 2). Similarly, Pasteuria Pr_C1 failed to attach on the esophagus wall of all Germany genotype replicates in all treatments. This suggests that the respective Daphnia genotypes retained their infection phenotype regardless of their microbiota or even in the absence of microbiota.

Discussion
Genetics and microbiota can have important functions in health and performance of animals. Here we report that genotypes of the model system D. magna from various geographic backgrounds still differed in their microbiome composition even after years of culturing the animals under the same dietary and environmental growth conditions. This indicates that the effects of the host genotype and of the host microbiota on the expression of phenotypic traits can be confounded. Both effects can be very important in the ecology and evolution of the system, but in different ways. For example, genes are only transmitted vertically, while microbiota can be transmitted vertically and horizontally. During migration, microbiota may be lost and in new habitats, new members may be acquired. To understand the expression and evolution of phenotypic traits, it is therefore necessary to disentangle host genetics from microbiota effects. Here we tested whether the first line of defense (i.e., the attachment of parasite in the esophagus or foregut entry) characterized by resistotypes of the water flea D. magna, are influenced by host genetic differences, or by the host’s microbiota as has been suggested in other systems.

We found that genotypic variation and parasite susceptibility and resistance of D. magna to specific P. ramosa strains, can correlate with differences in the microbial composition of the daphnid hosts (see Figs 2–4). In humans and mice, host genetics is reported to be a factor in shaping the composition and diversity of the microbiome.
and the abundance of some members of microbiota are associated with immune-related genes. However, despite significant variation in the microbiome composition and structure between daphnid hosts (especially between two highly-opposite resistotypes), we have not seen a microbiota influence on the resistance trait of *D. magna* against the parasite *P. ramosa* (at least with Pr_C1 strain) after performing a reciprocal microbiome

**Figure 3.** Biplot of db-RDA analysis using Unweighted UniFrac Distance, showing the clustering of microbiome composition between *D. magna* samples, with the analysis constrained by their resistotype to each *P. ramosa* strain (a) Pr_C1, (b) Pr_C19, (c) Pr_P15, and (d) Pr_P20. R (resistant) and S (susceptible) act as centroids to the clusters. Red colored samples belong to the susceptible *D. magna* resistotype and black colored samples belong to the resistant *D. magna* resistotype for the respective *P. ramosa* strains. Pseudo-F and p values < 0.05 denotes statistically significant variation between the microbiome of the *Daphnia* resistotypes.

**Figure 4.** Differences in microbiome composition between highly-resistant (RRRR) *D. magna* genotype from Germany (DE-K35-Inb1) and highly-susceptible (SSSS) *D. magna* genotype from Hungary (HU-HO-2) shown as average relative abundance at the family level. Statistical comparison was performed using Student’s T-test with levels of significance denoted by *for p < 0.05 and ** for p < 0.01.
shown that gut microbiome plays a protective role against trypanosomatid parasites5, with the host genotype playing a secondary role in their immunity3, and in wild Anopheles gambiae where a member of the midgut microbiome renders the insect resistant to the malaria-causing Plasmodium32. In a marine crustacean Palaemon macrodactylus, a bacterium constantly isolated from the eggs can protect the shrimp embryo from a fungal pathogen by releasing an anti-fungal chemical33. In human disease-related pathogenesis, a fecal microbiota probiotic could combat infection of the pathogen Clostridium difficile to alleviate suffering from chronic intestinal bowel disorder34,35. Furthermore, the human gut microbiota member, Bacteroides fragilis, has been proposed as a potential probiotic to improve autistic behavioral symptoms36. The disparity between our study and the emerging paradigm that the microbiome plays a major role in the resistance against parasite infection or disease pathogenesis emphasizes the idea that the microbiome’s role in immunity could be organism-dependent (or at least parasite-dependent) and should not be generalized across all taxa. Notably, the Daphnia-Pasteuria system exhibits strong host-parasite species interaction, with host infection rate dependent on the presence of diverse parasite strains that would specifically infect matching host genotypes37. The co-evolutionary arms race between Daphnia and Pasteuria may be a stronger driving factor on the evolution of innate immunity in D. magna against the pathogen and could mask the influence of microbiota (if there is any). Moreover, the difference of the Daphnia-Pasteuria system compared to other host-parasite systems may lie with how the Pasteuria infection process takes place, the decisive step being the attachment of Pasteuria to the esophagus. P. ramosa can only infect Daphnia if it manages to attach to the esophagus of the daphnid host25. It is evident that this inhibition of attachment is accomplished by the epithelial immunity or barrier defenses present in the esophagus (the esophagus being the extension of the foregut) of resistant Daphnia, and appears to be entirely independent from the microbiome influence. Interestingly, the absence of microbiome in resistant Daphnia did not alter the barrier defenses. Bacteria-free organisms are known to have abnormal and immature gut development38–40 which impairs mucosal immunity in the gut leading to disease susceptibility. In tse-tse flies, microbiota-free organisms are susceptible to trypanosome infection because of a compromised peritoneal matrix that act as a physical barrier to the pathogen, as demonstrated by attachment test of the parasite in the host’s esophagus. More importantly, despite the known contribution of the microbiome to host immunity, a Pasteuria-resistant Daphnia that was exchange. Remarkably, the resistotypes of the two Daphnia genotypes to the parasite were also not altered even when the animals were in a bacteria-free state, which strongly supports our conclusion, that the microbiome may not play a role in the innate immunity (first line of defense) of D. magna against a P. ramosa pathogen. These results are in direct contrast from what is reported in other taxa such as in bumblebees where it has been shown that gut microbiome plays a protective role against trypanosomatid parasites1, with the host genotype playing a secondary role in their immunity3. This is the first study to show that despite considerable differences in the microbiome composition between Daphnia genotypes, particularly that of P. ramosa-susceptible and P. ramosa-resistant Daphnia, reciprocal swapping of their microbiota did not alter the susceptibility or resistance of the respective daphnid hosts against a Pasteuria pathogen, as demonstrated by attachment test of the parasite in the host’s esophagus. More importantly, the genetic basis of the differing tolerance of natural populations of Daphnia to high copper concentrations44,45 and temperature46 has been investigated previously. The interaction of host genetics and the microbiome in this context could be an interesting area for further investigation.

**Conclusion**

This is the first study to show that despite considerable differences in the microbiome composition between Daphnia genotypes, particularly that of P. ramosa-susceptible and P. ramosa-resistant Daphnia, reciprocal swapping of their microbiota did not alter the susceptibility or resistance of the respective daphnid hosts against a Pasteuria pathogen, as demonstrated by attachment test of the parasite in the host’s esophagus. More importantly, despite the known contribution of the microbiome to host immunity, a Pasteuria-resistant Daphnia that was

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**Table 2.** Results of the infection phenotyping of bacteria-free animals and reciprocally transplanted microbiota between highly resistant Daphnia from Germany and highly-susceptible Daphnia from Hungary.

| Treatment | Positive attachment |
|-----------|---------------------|
| A. Pr_C1 Susceptible Daphnia (SSSS, Hungary clone) |  |
| BacFree (n = 20) | 100% |
| Susceptible Daphnia + own microbiota (n = 17) | 100% |
| Susceptible Daphnia + microbiota of Resistant Daphnia (n = 18) | 100% |
| B. Pr_C1 Resistant Daphnia (RRRR, Germany clone) |  |
| BacFree (n = 7) | 0% |
| Resistant Daphnia + own microbiota (n = 8) | 0% |
| Resistant Daphnia + microbiota of Susceptible Daphnia (n = 14) | 0% |
Methods

Animals. Eight D. magna genotypes were collected from different geographic regions, either as planktonic individuals or as resting eggs which were subsequently hatched and kept as clonal lines for several years (Table 1). Two clones included are the parents of a standing QTL panel27,28, one of which (Fl-Xinb3) was selfed three times, while the other (DE-K35-liinb1) was selfed once. The F1 lab_breed (QTL-IXF-1) genotype is the sexual offspring of these two parent clones and was hatched from a resting egg in the lab27. Animals were maintained in artificial Daphnia medium (ADaM48, for at least 20 generations under standardized environmental conditions; 20 °C, 16:8 light dark photoperiod, the green algae Scenedesmus sp. as only food and animals cultured.

Only female adults were sampled for the 16S rDNA profiling. Five jars with 80 ml ADaM containing two Daphnia of each genotype were grown for 4 weeks under standard laboratory conditions fed with 10 million cells algae per jar every other day. ADaM was replaced every week and only 2 adult individuals were kept in the jar every ADaM change. Animals were washed twice in sterile ADaM to remove unassociated bacteria before storage in −20 °C prior to DNA extraction.

Resistotype determination via esophagus-attachment test. Different D. magna genotypes were attachment-tested against four Pasteuria clones/isolates as described by Duneau et al.25 Daphnia show a binary infection outcome to Pasteuria pathogens; susceptible animals allow P. ramosa attachment to the esophagus or to the hindgut while resistant animals display no sign of attachment. P. ramosa Pr_C1 and Pr_C19 clones were isolated from infected D. magna from Moscow (Russia) and Gaarzerveid (Germany), respectively, and previously characterized in Luijckx et al.26. Pr_C1 and Pr_C19 mode of attachment is through the esophagus. P. ramosa Pr_P15 and Pr_P20 are isolates that have been propagated in the same D. magna host clones several times, but where not cloned. Pr_P20 was isolated from an infected D. magna female that was collected in 2011 from Aegelsee, Switzerland. Pr_P20 attaches to the host in the esophagus. Pr_P15 is derived from an infected D. magna collected in Belgium and is unique in that it attaches usually to the hindgut of the host.

P. ramosa spores were fluorescently labeled and attachment tests performed as described by Duneau et al.25. Briefly, spores were labeled with fluorescein-5(6)-isothiocyanate (FITC) dye at a concentration of 1 mg in 0.1 M NaCO3 (pH 9.1). The solution was kept in the dark for 2 hours and washed 3–5 times via centrifugation. Daphnia were exposed to resuspended spores (5000 spores per individual) for 1 hour in the dark. Animals were then examined for spore attachment with an inverted epifluorescent microscope (Leica DM 2500) at 400 × magnification. Resistotypes were defined as 4 letter codes (e.g. RSSR) composed of R (for resistance, no attachment) and S (susceptibility, attachment) for P. ramosa strains Pr_C1, Pr_C19, Pr_P15, Pr_P20, respectively.

DNA extraction. Frozen animal tissues were bead-beat with two pieces of acid-washed and autoclaved 1.0 mm beads and a 100 µl of 0.1-mm silica zirconian beads in PCR water in a FastPrep F120 cell disruptor (Qbiogene, USA) for 30 s at 4 m·s−1. DNA extraction was continued following the steps described in the DNeasy Blood and Tissue Kit (Qiagen, Netherlands). PCR water was used as a control for the extraction procedure and went through the same extraction and PCR amplification steps as the Daphnia samples.

454 Barcode Tagging. Two PCR reactions steps were carried out in amplifying the V3-V5 hypervariable region of the 16S rDNA with Roche 454 barcodes. The first PCR amplification step involved using the primer pairs F (5′-CACCCGGYXCAACTCTCTAC-3′) and R (5′-TTCGWCCTGAAATTCWCAC-3′) that targets the 327–969 bp of the 16S rDNA using Phusion High Fidelity Taq (New England Biolab, USA) with this PCR program: 98 °C for 1 min, 15 cycles of 98 °C for 10 s, 50 °C for 20 s and 72 °C for 30 s with a final extension time of 72 °C for 5 min. PCR reactions were treated with ExoSap (Fermentas, USA), to remove unincorporated nucleotides and primers. A second PCR reaction step was prepared using the first PCR purified product as DNA template and re-amplified with F/R primer pairs containing the Libl fusion sequence and barcodes (PFx: 5′-CCATCTCATCCCTGGCGTGCTCTCCGACTCAGbarcodeFF primer-3′.

PRs: 5′-CCATCCCTGGCTGTGACCTTGGATACCGGCTAC primer-3′) using the same PCR program and cycles. Amplified products were purified using Agencourt Ampure XP beads (Beckman and Coulter, USA) and eluted in sterile TE buffer. DNA concentrations were measured using Qubit 2.0 Fluorometer (Invitrogen, USA) and a pool of equimolar amounts (30 ng/µl) of barcoded products were sequenced on 454 GS-FLX instrument with Libl Titanium chemistry (Roche, USA) at Microsynth (Balgach, Switzerland). A DNA mock community consisting of 76 bacterial strains belonging to 28 genera and families of various phyla was also tagged with a barcode and was sequenced as a positive control (see Supplementary Table S1). The mock community consisted of bacterial isolates previously isolated from Daphnia animals either freshly collected from the field or maintained in the lab, the algal food cultures (Scenedesmus sp.) or ADaM (Daphnia culture medium). Isolates were chosen to represent different bacterial phylogenies and based on the availability of bacterial cultures.

Sequence Data Processing and Analysis. Demultiplexed samples were denoised and pre-clustered at 99% level similarity using the method described by Reeder and Knight49. Samples were checked for chimeras, low quality sequence and short sequence reads (<150 bp) prior to post-analysis with the Quantitative Insights Into Microbial Ecology (QIIME 1.9.1)46. OTUs were clustered at 97% sequence similarity with UCLust51. Representative sequences were picked for taxonomic identification based on Blast 2.2.2752. OTU representative sequences were aligned with Pynast53, while gaps in the alignment were filtered using the default 16S rDNA
Greengenes alignment lane mask before making the phylogenetic tree using FastTree 2.1.34. Singletons and chlo-roplast sequences were filtered out of the OTU table before downstream analyses.

**Microbial community composition analysis.** To control for sequencing effort, samples were rarified at a minimum of 10 sequences and a maximum depth of 3893 sequences in steps of 50, and performing 20 iterations for each step. The microbial diversity for each *D. magna* sample was assessed using 4 alpha diversity indices (Observed_OTUs, Phylogenetic Diversity, Chao1 and Shannon index) at 3893 sequence depth and statistically compared using Student’s T-tests in JMP 12.0 (Cary, NC, USA). Since our main objective is to compare the microbial community between genotypes with contrasting resistotypes for the reciprocal transplant experiment, only alpha diversity of RRRR (Germany, n = 5) and SSSS (Hungary, n = 3) host genotypes were reported in this study.

For microbial composition assessment of *Daphnia* genotypes and resistotypes, the OTU table was rarified at an equal depth of 3893 sequences to generate pairwise distance matrices. Weighted and Unweighted UniFrac and Bray-Curtis distance matrices derived from this rarified beta diversities were used to statistically test for differences in microbial composition using ANOSIM and ADONIS methods with 999 permutations. Distance-based redundancy analysis (db-RDA) was also applied to determine the clustering of *D. magna* microbiome composition when constrained by their resistotype to each *P. ramosa* strain.

**Reciprocal transplant of microbiota between susceptible and resistant *Daphnia.*** To determine if microbiota influences the resistotype of *D. magna* to *Pasteuria* infection, we removed the bacteria, or swapped the microbiota of two *Daphnia* genotypes that were previously tested to be highly-susceptible (SSSS, Hungary clone) and highly-resistant (RRRR, Germany clone) to four *P. ramosa* clones and isolates (see Table 1). The detailed methods for synchronizing the animals, the production of bacteria-free *Daphnia* from asexual eggs, the preparation of bacterial suspension, preparation of axenic *Scenedesmus obliquus* algae and subsequent PCR screening for bacteria-free *Daphnia* and algae were previously described in Sison-Mangus et al.24 Briefly, eggs from 100 individuals of synchronized Hungarian and German *Daphnia* mothers were extracted from the mother’s brood pouch and washed with sterile ADaM three times. Eggs were treated with freshly made Ampicillin (1 mg/mL) and Kanamycin (100 μg/mL) for 48 hours until development of hatchlings. Sterile ADaM and antibiotics were replaced each day. The bacteria-free state of the hatchlings after antibiotics treatment was verified by 16S rDNA PCR amplification.

Two independent reciprocal transplant experiments were performed using the Hungarian (SSSS) and the German (RRRR) clones as respective experimental animals. After antibiotics removal, bacteria-free *Daphnia* hatchlings were divided into groups representing 3 treatments: 1) Bacteria-free *Daphnia*, 2) *Daphnia* inoculated with microbiota from the same clone, and 3) *Daphnia* inoculated with microbiota coming from clone with opposite resistotype (e.g., German hatchlings received the microbiota from Hungarian clone while the Hungarian hatchlings received the microbiota from German clone). Bacterial suspensions were harvested from the respective *Daphnia* clone sources following the method of Sison-Mangus et al.24 with 1 ml bacterial suspension given to respective hatchlings from each group. The optical density (OD$_{600}$) of the bacterial suspension ranged from 0.72–0.73. Hatchlings were exposed to bacteria for 24 hours and subsequently washed once before placing in individual experimental jars with 25 million axenic algal food cells.

Hatchlings were grown under the same environmental conditions as above and maintained daily as in Sison-Mangus et al.24 *P. ramosa* Pr_C1 stick testing was done at Day 6 post-hatching using the method described above, but with antibiotic treated spores. For this, the FITC-dyed *P. ramosa* spores were treated with triple antibiotic solution (Ampicillin (2 mg/mL), Kanamycin (1 mg/mL) and Tetracyclin (100 μg/mL) for 36 hours in the dark prior to stick testing. This is to prevent unwanted bacteria that maybe present from *Pasteuria* spore solution from contaminating the experimental treatments during the attachment test. Preliminary test showed that *P. ramosa* C1 spores remain viable (able to attach to the esophagus of *Daphnia*) after being treated with antibiotics.

**Data availability.** The raw sequencing reads have been deposited at the NCBI Short Read Archive under BioProject ID PRJNA407858 with the following BioSample accession numbers: SAMN07665251 to SAMN07665254 for IR-GG1–1 clone, SAMN07665255 to SAMN07665258 for IL-M1–1 clone, SAMN07665259 to SAMN07665261 for HU-HO-2 clone, SAMN07665262 to SAMN07665264 for BE-OHZ-M5 clone, SAMN07665265 to SAMN07665267 for BE-OHZ-M10 clone, SAMN07665268 to SAMN07665272 for FI-Xin3 clone, SAMN07665273 to SAMN07665277 for DE–K35-Inb1 clone, SAMN07665278 to SAMN07665281 for QTL-IXF-1 clone and SAMN07695326 for Positive Control.

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**Author Contributions**

M.P.S.M. and D.E. conceived the study. M.P.S.M., D.E. and C.M. designed the study. C.M. conducted the esophagus-attachment test. M.P.S.M. conducted the *Daphnia* experiments and performed the microbiome data analysis. All authors contributed to the final version of the manuscript.

**Additional Information**

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