Locally adapted gut microbiomes mediate host stress tolerance

Shira Houwenhuyse1 · Robby Stoks2 · Shinjini Mukherjee3 · Ellen Decaestecker1

Received: 10 July 2020 / Revised: 29 January 2021 / Accepted: 11 February 2021 / Published online: 3 March 2021
© The Author(s) 2021. This article is published with open access

Abstract
While evidence for the role of the microbiome in shaping host stress tolerance is becoming well-established, to what extent this depends on the interaction between the host and its local microbiome is less clear. Therefore, we investigated whether locally adapted gut microbiomes affect host stress tolerance. In the water flea Daphnia magna, we studied if the host performs better when receiving a microbiome from their source region than from another region when facing a stressful condition, more in particular exposure to the toxic cyanobacteria Microcystis aeruginosa. Therefore, a reciprocal transplant experiment was performed in which recipient, germ-free D. magna, isolated from different ponds, received a donor microbiome from sympatric or allopatric D. magna that were pre-exposed to toxic cyanobacteria or not. We tested for effects on host life history traits and gut microbiome composition. Our data indicate that Daphnia interact with particular microbial strains mediating local adaptation in host stress tolerance. Most recipient D. magna individuals performed better when inoculated with sympatric than with allopatric microbiomes. This effect was most pronounced when the donors were pre-exposed to the toxic cyanobacteria, but this effect was also pond and genotype dependent. We discuss how this host fitness benefit is associated with microbiome diversity patterns.

Introduction
Many organisms face different environmental conditions across their species’ range and thereby evolved local adaptation with organisms being relatively fit in their local conditions and relatively unfit in others [1, 2]. Local adaptation is often studied in macro-organisms across large-scale environmental gradients [3–7]. Nevertheless, it can also occur at much smaller geographical scales and in micro-organisms, as shown for parasites and microbiomes [8–11]. While there is ample evidence for local adaptation among natural populations, it is not always detected [8–10]. One reason may be that local adaptation is widely considered to be the result of fitness trade-offs across environments [11, 12], which may be more likely to be detected under stressful conditions that limit energy stores [13–16]. Local adaptation may, however, also truly be absent because of out-of-phase dynamics in coevolutionary processes, a weak response to selection, and gene flow between host populations [9, 17–19].

Insights in the genomic targets and the molecular mechanisms underpinning local adaptation are starting to accumulate thanks to the increased affordability of high-throughput sequencing [20–22]. However, several challenges, such as the identification of the true targets of local adaptation still remain [23]. Recently, the gut microbiome of different organisms has emerged as a key determinant of many aspects of organismal biology, capable of shaping developmental, physiological and reproductive phenotypes [24–29]. The composition of gut microbiomes differs spatially in function of the regional environmental conditions [30–34], and there is some evidence that different host genotypes differ in their gut microbiome [35, 36]. It is therefore likely that when hosts show local genetic adaptation, this may be misleading as their gut microbiome may, at least partly, be contributing to this pattern [37]. There is recent evidence that the gut microbiome plays a role in local
adaptation to environments differing in, for example, pollution [38], aridity [39] and salinity [40, 41]. These studies, however, mainly focused on how the microbiome differed between locations, but did not explicitly investigate the host fitness effects caused by carrying a local microbiome.

In this study, we tested through reciprocal transplants, whether having a sympatric vs. an allopatric gut microbiome gives the host an advantage in stressful environments. The underlying hypothesis is that hosts profit from having a locally adapted microbiome, which increases their fitness under stressful conditions. This was hypothesized to be associated with two different scenarios in terms of gut microbiome diversity and establishment. Based on Macke et al. [42, 43], *Daphnia magna* selects beneficial bacteria from the environment. We expect stronger host-mediated selection for beneficial bacterial strains in sympatric than in allopatric microbiomes. If so, the gut will be occupied by locally selected bacteria leaving less opportunity for other bacteria to establish, which may result in more convergent microbiomes over the host genotypes (see Fig. 1) [44]. Alternatively, hosts with a sympatric microbiome can be expected to have a larger bacterial spectrum with beneficial gene functions [45], hence showing a higher bacterial diversity than with an allopatric microbiome.

To unravel whether a local microbiome could play a role in increased tolerance, *D. magna* genotypes from different ponds out of two separate regions were exposed to toxic cyanobacteria in a reciprocal transplant experiment in which germ-free *Daphnia* were inoculated with a local or foreign gut microbiome. The zooplankter *D. magna* is an ecological and evolutionary model organism for which local adaptation has been shown in response to stressors, among which cyanobacteria [46–48]. A commonly occurring cyanobacterium genus is *Microcystis*, which is poor food, interferes with *Daphnia* filtering activity, and produces a wide range of deleterious cyanotoxins [49–51]. Because *Microcystis* exposure is stressful for *Daphnia*, exposing *Daphnia* to it makes it likely to detect trade-offs and local adaptation in the host. Because gut microbiomes pre-exposed to *Microcystis* have been shown to provide a higher tolerance to *Microcystis* for the *D. magna* host [42, 43], we tested whether local adaptation in the recipients was dependent on pre-exposure of the donors to toxic *Microcystis aeruginosa*. We addressed following questions: (1) Is fitness higher in
D. magna receiving a sympatric than an allopatric microbiome and does this depend on the donor diet? (2) Do D. magna have a lower or higher microbial diversity when they receive a sympatric vs. an allopatric microbiome? and (3) how is this related to their fitness and genotype?

Materials and methods

Sampling of the Daphnia magna genotypes and pond water

Five D. magna genotypes from each of two regions in Belgium separated by circa 100 km (Kortrijk and Leuven) were collected (Table S11). In Kortrijk, we obtained D. magna genotypes from the Blauwe Hoeve (K_BH), Kennedypark (K_KP) and Morinnewstraat (K_MS), by sampling one individual from the active population in each pond. In addition, two genotypes were obtained from a recent (top layer) resting eggs in Zwevegem (K_ZWE1 and K_ZWE2) as at the time of sampling no active population of D. magna was present. In Leuven, all genotypes were obtained from recent (T = top layer sediments) resting eggs of two ponds: Heverlee (L_OM2) and Oud-Heverlee (L_T2, L_T3, L_T7 and L_T8; coding is based on Cousyn et al. [52]). After being kept as stock cultures, they were raised as three independent clonal (maternal) lines (i.e., genetically identical Daphnia individuals raised independently from each other to control for maternal effects). The ponds from Kortrijk out of which the genotypes were sampled did not have a Microcystis bloom at sampling (S. Houwenhuysse, personal observation), but in some ponds of Leuven (i.e., L_OM2) Microcystis blooms have been observed (E. Decaestecker, personal observation). It could be possible for the Leuven genotypes to be locally adapted to cyanobacteria. However, Daphnia adaptation to Microcystis strains is assumed to be locally structured [53], and we here test the role of a local vs. a foreign microbiome towards a 'neutral' toxic cyanobacterial strain. Hence, irrespective of the absence/presence of cyanobacteria in the ponds, no adaptation to the here tested toxic cyanobacterial strain was expected in Daphnia. Maternal lines of the D. magna genotypes were cultured in filtered tap water at a temperature of 19±1 °C and under a 16:8 h light:dark cycle in 2 L glass jars (at a density of 30 individuals/L). They were fed three times per week with saturating amounts of the green algae Chlorella vulgaris. The medium of these cultures was refreshed once per week. Three months before the start of the experiment 10 L pond water was collected from each pond and stored at 4 °C to maintain the bacterioplankton community. At the time of pond water sampling, there was no visible presence of cyanobacteria in any pond. After sieving the pond water over a 100 µm sieve, two mixtures (=filtered local pond water) were made, one from the Kortrijk ponds, and one mixture from the Leuven ponds to be inoculated in the experimental jars during the transplant experiment.

Cultivation of green algae and cyanobacteria

D. magna individuals were fed with the unicellular green algae Chlorella vulgaris (which is good-quality food for D. magna [54]) or a combination of C. vulgaris and M. aeruginosa (which is a toxic cyanobacterial strain for D. magna [42]). We used the M. aeruginosa strain PCC 7806, isolated from the Braakman reservoir in the Netherlands (51°19’22” N, 3°44’16” E) and part of the Culture Collections at Institute Pasteur (Paris, France). C. vulgaris and M. aeruginosa were grown in Wright’s Cryptophyte (WC) medium [55] and modified WC medium (without Tris), respectively. The algae were cultured under sterile conditions in a climate chamber at 22±1 °C with a light:dark cycle of 16:8 h in 2 L glass bottles, with constant stirring and aeration. Filters (0.22 µm) were placed at the input and output of the aeration system to avoid any bacterial contamination. The axenity of the algal cultures was checked on Lysogeny Broth medium [56] agar plates. Ash-free dry weight of the cultures was determined following Mohieman et al. [57].

Gut microbiome transplant experiment

To determine if D. magna shows a better performance upon exposure to toxic cyanobacteria in the presence of a sympatric vs. an allopatric gut microbiome, a reciprocal gut microbiome transplant experiment was conducted (Fig. 2). We first exposed donor individuals to M. aeruginosa to test whether pre-exposure of the microbiome results in a higher stress tolerance in the recipients. Donor genotypes were pooled per region (five genotypes per region), per algal treatment (pre-exposed to a mixture of 60% toxic M. aeruginosa and 40% C. vulgaris, or to 100% C. vulgaris) and per replicate and were kept at a density of 50 (i.e., ten per genotype) individuals in 2 L experimental jars with filtered local pond water. Donor diets had a final carbon concentration of 1 mg/L and were given to the D. magna three times per week. The pooled donor D. magna genotypes stayed in these conditions for 10 days to adjust the gut microbiome when exposed to M. aeruginosa [42]. Afterwards, 25 D. magna individuals per donor region × diet combination were randomly isolated and kept in sterile-filtered tap water for 24 h to remove food particles from the gut, as well as environmental bacteria on the carapace and filter apparatus. Then, guts were extracted with dissection needles under a stereo microscope and placed in an Eppendorf tube containing 1 mL sterile milliQ water. These
dissected guts were then stored at 4 °C for 7 days. Each donor condition was set up in triplicate, which resulted in 12 pooled donor inocula (two donor regions × two donor diets × three replicates). After 7 days, guts from these donor populations were dissected and used as in inoculum. In the recipient phase, the five (and one, one of the dissected guts were then stored at 4 °C for 7 days. Each donor condition was set up in triplicate, which resulted in 12 pooled donor inocula (two donor regions × two donor diets × three replicates). After 7 days, the stored donor inocula were crushed and given to sterile D. magna juveniles. In the recipient phase, each genotype received a pooled and crushed gut microbiome of one of the four donor treatment combinations. This resulted in a fully crossed design: donor region (two levels) × donor diet (two levels) × D. magna genotype (nested within two recipient regions, with four genotypes for Kortrijk as one genotype did not hatch after sterilization and five genotypes for Leuven) with three replicates per genotype (one per independent maternal line). In sympatric combinations D. magna received a local donor inoculum, in allopatric combinations they received a foreign donor inoculum. Sterile D. magna juveniles were obtained following the protocols of Callens et al. [58, 59]. Therefore, per maternal line of each genotype, 30 parthenogenetic eggs with external membranes were collected in a six-well plate with 15 eggs in 5 mL filtered tap water per well. Afterwards, they were placed in a laminar flow hood to disinfect the eggs by submerging them in 5 mL of a 0.1% glutaraldehyde solution and gently agitating them for 10 min. Then, the eggs were transferred to another well, containing 5 mL sterile filtered tap water to remove glutaraldehyde residues. The eggs stayed in this rinsing step for 10 min after which the rinsing step was repeated. Afterwards, the eggs were transferred in groups of 15–5 mL sterile filtered tap water. The six-well plate was sealed with parafilm and placed in an incubator. Eggs were allowed to hatch for 72 h at a temperature of 20 ± 0.5 °C and a 16:8 h light:dark cycle. The genotype K_ZWE1 did not hatch after sterilization and therefore excluded from the recipient phase of the experiment, resulting in a total of nine recipient genotypes. Per maternal female line, eight germ-free juveniles were placed per two in 50 mL falcon tubes filled with 45 mL sterile filtered tap water. Each of those pairs received one of the four different pooled donor inocula. The recipient D. magna were incubated with the pooled donor inocula for 48 h, after which D. magna were individually transferred to new 50 mL falcon tubes filled with filtered local pond water. In these 48 h, the recipient D. magna will have taken up a substantial part of the bacteria from the pooled donor inoculum. After the transfer to (unsterile) filtered local pond water (filtered over a 100 µm sieve), all recipient D. magna were subjected to a cyanobacterial diet (i.e., 60% toxic M. aeruginosa and 40% C. vulgaris), which was administered every 48 h to test if there was an effect from the donor gut microbiome on Daphnia tolerance to toxic cyanobacteria in terms of host fitness, survival and reproduction, monitored every 48 h during 21 days. Body size was measured at days 3, 6, 10 and 20. After 21 days, the guts from the surviving D. magna were dissected for amplicon sequencing. The D. magna
were placed in sterile-filtered tap water for 24 h to remove food particles from the gut, as well as environmental bacteria on the carapace and filter apparatus. After these 24 h, they were dissected under a stereo microscope using dissection needles and collected in 10 µL sterile milliQ water, after which they were stored in −20 °C.

**Library preparation and sequencing**

To characterize the gut microbial communities from the donor sets and from the recipient *D. magna* at the end of the 21-day monitoring period, DNA was extracted using the PowerSoil DNA isolation kit (Qiagen). DNA was dissolved in 20 µL milliQ water. The total DNA yield was determined using a Qubit dsDNA HS assay (Invitrogen) on 1 µL of sample. A nested PCR was applied to increase specificity and amplicon yield. The full-length 16S rRNA gene was first amplified with EUB8F and 1492R primers on 10 ng of template using a high-fidelity SuperFi polymerase (Life Technologies) for 30 cycles: 98 °C—10 s; 50 °C—45 s; 72 °C—30 s. PCR products were subsequently purified using the QIAquick PCR purification kit. To obtain dual-index amplicons of the V4 region, a second amplification was performed on 5 µL (=20–50 ng) of PCR product using 515F and 806R primers for 30 cycles: 98 °C—10 s; 50 °C—5 s; 72 °C—30 s. Both primers contained an Illumina adapter and an 8-nucleotide (nt) barcode at the 5′-end. For each sample, PCRs were performed in triplicate, pooled and gel-purified using the QIAquick gel extraction kit. An equimolar library was prepared by normalizing amplicon concentrations with a SequalPrep Normalization Plate (Applied Biosystems) and subsequent pooling. Amplicons were sequenced using a v2 PE500 kit with custom primers on the Illumina Miseq, producing two ×250-nt paired-end reads. This way, 96 recipient samples were generated representing eight genotypes (one of the nine recipient genotypes, L_T2, did not have any surviving individuals after 21 days) × two donor regions × two donor diets × three replicates, and the 12 donor samples. The number of sequenced *D. magna* individuals per treatment is represented in the Table SI2.

DNA sequences were processed following Callahan et al. [60]. Sequences were trimmed (the first ten nucleotides and from position 190 onwards) and filtered (maximum of two expected errors per read) on paired ends jointly. Sequence variants were inferred using the high-resolution DADA2 [61] method, which relies on a parameterized model of substitution errors to distinguish sequencing errors from real biological variation. Chimeras were subsequently removed from the dataset. After filtering, a total of 907,846 reads were obtained with on average 24,536.38 reads per sample (minimum = 23 reads and maximum = 53,480 reads), with most samples having more than 10,000 reads (only four exceptions with 23, 9080, 8478 and 3840 reads). Taxonomy was assigned with a naïve Bayesian classifier using the Silva v132 training set. OTUs with no taxonomic assignment at the phylum level or which were assigned as “chloroplast” or “cyanobacteria” were removed from the dataset. OTUs for which the mean relative abundance was below 10⁻³ were removed from the analysis. To visualize the bacterial families that differed between the treatments, OTUs were grouped at the family level, and families representing <1% of the reads were discarded (this was not done for the analyses). To test for differences in α- and β-diversity, all samples were rarefied to a depth of 1000 reads. One gut sample (KP, with a sympatric donor microbiome pre-exposed to *M. aeruginosa*) had a lower number of reads, and was removed from the analysis.

**Statistical analyses**

Statistical analyses of the life history traits were performed in R 4.0.0 (R studio version 1.1.463). We used the Akaike information criterion (AIC) to select the best subset of variables to represent the best model (Table SI3). We evaluated general linear models (GLM with Gaussian or normal distribution, as the data was normally distributed) and linear mixed-effects models (LMER). We first compared models with the same fixed effects, but different random effect (i.e., (1|Region:Pond:Genotype) and (1|Region:Pond)). Secondly, we tested the significance of the fixed factors in the model with the best random effects factor. Type II ANOVA tables for fixed-effect terms with Satterwhaite and Kenward–Roger methods for dominant degrees of freedom for F-tests and p values were created (Anova function of the car package [62]). In the final model, we included donor diet (toxic cyanobacteria absent or present), donor microbiome type (allopatric or sympatric microbiome), and *D. magna* genotype (=clone) as fixed factors, with genotype nested within pond (see Table SI1) and pond nested within region as random factor. We also included all possible interactions. Survival was analyzed in two different ways, first using a log-rank or Mantel-Haenszel test (from the OIsurv, survival and survminer packages in R [63]) on a random selection of one genotype per pond (fully randomized over the genotypes, and thus correcting for dependency between genotypes within the L2 pond) and on a total of nine recipient genotypes. The survival times of individuals that were still alive at the end of the 21-day experiment were coded as right-censored. The second way the survival was analyzed was on the percentage *D. magna* individuals that survived until the end of the experiment. This was done with a general linear mixed-effects model (from the lme4 package in R [64]), controlling for unbalanced design with a restricted maximum likelihood estimation. Total fecundity and body size were
analyzed on a total of eight recipient genotypes. Total fecundity was analyzed for these *D. magna* individuals that survived until day 21 with a linear mixed-effects model, controlling for unbalanced design with a restricted maximum likelihood estimation. Body size was analyzed with a linear mixed-effects model with the four subsequent measurements (days 3, 6, 10 and 20) as repeated measures and also controlled for unbalanced design with a restricted maximum likelihood estimation. To specifically analyze relationships in the response patterns to *M. aeruginosa* pre-exposure between traits, we calculated delta values as the clonal average trait value when pre-exposed to *M. aeruginosa*—the clonal average trait value when not pre-exposed. We did so for survival, total fecundity and body size (days 3, 6, 10 and 20). We then tested with a Pearson correlation test, with bonferroni correction for multiple testing (using the ggpubr package in R [65]), for pairwise relationships between delta values of different traits (e.g., between survival and body size).

Measures for α-diversity of the gut microbial communities within the different treatments, OTU richness and the Shannon entropy (taking into account both OTU richness and the relative abundance of OTUs) were calculated using the vegan package in R [66]. Shannon entropy was calculated as the exponential function of the Shannon entropy, this represents the true diversity of the bacterial community in the sample [67, 68]. We used the AIC criterion to select the best subset of variables to represent the best model (Table SI3). We evaluated GLM (with Gaussian or normal distribution, as the data was normally distributed) and LMER. We first compared models with the same fixed effects, but different random effect (i.e., (1|Region:Pond:Genotype) and (1|Region:Pond)). Secondly, we tested the significance of the fixed factors in the model with the best random effects factor. Type II ANOVA tables for fixed-effect terms with Satterhwaite and Kenward-Roger methods for dominator degrees of freedom for F-tests and p values were created (Anova function of the car package [62]). In the final model, the effects of donor diet, microbiome type, genotype, and all possible interactions, were assessed through a linear mixed-effects model, controlling for unbalanced design with a restricted maximum likelihood estimation with as random factor genotype nested in pond and pond nested within recipient region on a total of eight recipient genotypes. To investigate differences in community composition (β-diversity) between the different microbial communities, Bray–Curtis dissimilarity indices were calculated and plotted using principal coordinates analysis with the phyloseq package in R [69]. The effects of donor diet, microbiome type, genotype, and all possible interactions, with random factor genotype nested in pond and pond nested within recipient region on β-diversity were assessed through a permutation MANOVA, using the Adonis2 function in the vegan package in R [70] on the total dataset (i.e., donors and recipients), and on the donors and the recipients separately. To identify bacterial families that differed between donors and recipients, between diets and between microbiome types, OTUs were grouped at the family level, and families representing <1% of the reads were discarded. Differential abundance analyses were then performed with the Bioconductor package DESeq2 [71]. A Unionplot was created using the wilcox/unionplot function from GitHub, to show the OTUs that are unique and shared between donor, allopatric and sympatric treatments. In addition, the Bray Curtis dissimilarity matrix was extracted to perform an ANOVA on this matrix, to determine the difference in distance between donor and recipients across the treatments. Based on AIC the best model was a general linear model with diet, microbiome type and genotype as fixed factors.

### Results

Given the complexity of the results, we only focus on significant effects in the text below. For full results, see Tables SI 1–12.

#### D. magna life-history traits

For survival, the three-way interaction between donor diet (pre-exposed to *M. aeruginosa* or not), donor microbiome type (sympatric vs. allopatric) and recipient pond, was highly significant ($X^2=57.9$, df = 23, $p<0.0001$; Table SI4). When tested separately per donor diet and using one genotype per pond, there was a significant recipient genotype × donor microbiome type only when the host received a microbiome pre-exposed to the toxic *M. aeruginosa* strain ($X^2=23.7$, df = 11, $p=0.01$), but not when the microbiome was not pre-exposed to *M. aeruginosa* ($X^2=16.2$, df = 11, $p=0.1$). When the microbiome was pre-exposed to the toxic *M. aeruginosa* strain, seven out of the nine genotypes (K_MS, K_ZWE2, L_OM2, L_T2, L_T3, L_T7 and L_T8) had a higher survival percentage when they received a sympatric than an allopatric microbiome, for the other two genotypes (K_BH, K_KP) the survival percentage was higher when they received an allopatric than a sympatric microbiome (Fig. 3, Table SI5). This result is confirmed when looking at the interaction plots (Fig. 4A, B), most of the genotypes had a higher survival when they received a sympatric microbiome pre-exposed to *M. aeruginosa*.

For *D. magna* fecundity, there was a significant three-way interaction between donor diet × donor microbiome type × genotype ($F=2.6$; df = 7, 63; $p=0.02$; Table SI6 and Fig. SI1). Separate analyses per donor diet showed no significant interaction between microbiome type and genotype for each diet. There was, however, a marginal significant diet × microbiome type ($F=3.8$; df = 1, 63;...
reflecting that when *D. magna* individuals received a microbiome that was pre-exposed to the toxic *M. aeruginosa* strain more juveniles were produced when they received a sympatric than an allopatric microbiome (Fig. 4C, D and Fig. SI2). This was not the case when the donor inoculum was not pre-exposed to *M. aeruginosa*, there less juveniles were reproduced when they received a sympatric than an allopatric microbiome. Separate analysis, with a model excluding the fixed factor genotype, showed that the diet x microbiome type interaction was dependent on the genotype, as the interaction between diet and microbiome type was no longer significant in this model ($F = 2.7; df = 1, 80; p = 0.1$).

For *D. magna* body size, there was a three-way interaction between donor diet, microbiome type, and genotype (Table SI7). Separate analysis per donor diet showed that when *D. magna* received a microbiome pre-exposed to *M. aeruginosa* there was no significant interaction between microbiome type and genotype ($F = 1.6; df = 7, 241; p = 0.13$). However, when the *D. magna* received a microbiome not pre-exposed to *M. aeruginosa* there was a marginally significant interaction between donor microbiome and genotype ($F = 1.9; df = 7, 219; p = 0.066$). In addition, there was a significant interaction between donor diet and microbiome type across the repeated size measurements ($F = 6.5; df = 1, 248; p = 0.011$).

*D. magna* receiving a donor microbiome pre-exposed to *M. aeruginosa* had a smaller body size than when not pre-exposed to *M. aeruginosa*, and this effect was most pronounced when they received a sympatric microbiome (Figs. 4E, F and Fig. SI3). This diet x microbiome type interaction was not dependent on the genotype. A separate analysis, with a model excluding the fixed factor genotype, showed that the diet x microbiome type interaction was still significant ($F = 6.5; df = 1, 352; p = 0.01$). There was also a marginally significant interaction between microbiome type and genotype ($F = 1.9; df = 7, 617; p = 0.057$, Fig. SI4), and time and genotype ($F = 2.04; df = 21, 248; p = 0.005$) for *D. magna* body size, and a significant main effect of genotype ($F = 5.9; df = 7, 648; p < 0.0001$) and time (day 3, 6, 10 and 20; $F = 1114; df = 3, 248; p < 0.0001$).

Analysis of the differences between control and cyanobacterial donor diet revealed a marginally significant negative correlation between the *D. magna* survival and body size (at day 10) of the recipients (Pearson correlation using the average per genotype: $r = -0.48, t = -2.08, df = 14, p = 0.055$; Table SI8) reflecting that genotypes that showed a more positive response in survival tended to show a more negative response in body size, when they received a microbiome that has been pre-exposed to toxic *M. aeruginosa* (Fig. 5).
Gut microbiome composition

The donor microbiomes consisted mainly of Gammaproteobacteria and Bacteroida (Flavobacteria). The recipient gut microbiomes also harbored Gammaproteobacteria and Bacteroida, but were more diverse than the donor microbiomes (Shannon entropy: $F = 201; df = 1, 30; p < 0.0001$; Fig. SI5) and contained also bacteria from following classes: Actinobacteria, Alphaproteobacteria, Bacilli, Planctomycetacia and Verrucomicrobiae (Fig. 6 and Table SI9).

Fig. 4 Interaction plots representing the three-way interaction for survival, reproduction, body size and Shannon entropy. Interaction plots representing: percentage survived *D. magna* at day 20 (**A, B**), total fecundity (**C, D**), body size at day 20 (**E, F**) and exponential value of the Shannon entropy (**G, H**) of the *D. magna* genotypes. Left column shows variable in function of microbiome type (allopatric vs. sympatric) with donor microbiomes pre-exposed to *M. aeruginosa* in full lines and donor microbiomes not pre-exposed to *M. aeruginosa* in dotted lines. Right column shows variable in function of diet (pre-exposed or not to *M. aeruginosa*) with sympatric microbiomes in full lines and allopatric microbiomes in dotted lines.
There was no correlation between the number of guts per sample and Shannon entropy per sample (Pearson correlation test between paired samples: $r = 0.24, t = 1.38, df = 30, p = 0.17$). There was a significant distinction between the gut microbiome of the four donor sets and the recipient $D. magna$ measured by $\alpha$- and $\beta$-diversity ($\alpha$-diversity: $F = 38.2, df = 1.18, p < 0.0001$; $\beta$-diversity: $F = 11.04, R^2 = 0.26, df = 1, p = 0.001$; Figs. S15 and S16). In the donor populations there was a significant interaction between region and diet ($F = 5.1 \times 10^5, df = 1, 1; p = 0.0008$). In the population from Kortrijk the Shannon entropy was higher when they were fed with $M. aeruginosa$ than when they were not fed with $M. aeruginosa$. In Leuven the opposite was true, there
the Shannon entropy was higher when they were not fed with *M. aeruginosa* than when they were fed with *M. aeruginosa* (Fig. S17). In the recipient *D. magna*, there was a significant three-way interaction between donor microbiome type, donor diet and genotype for the Shannon entropy ($F = 131; \text{df} = 54; p = 0.0001$, Table SI10 shows the results of the linear fixed-effects model with diet, microbiome type and genotype as fixed factors and genotype nested within pond and pond nested within region as random factor). Per donor diet, the interaction between microbiome type and recipient *D. magna* genotype was significant (pre-exposed to *M. aeruginosa*: $F = 129; \text{df} = 5, 2; p = 0.007$, not pre-exposed to *M. aeruginosa*: $F = 174; \text{df} = 6, 2; p = 0.005$). We observed inconsistent patterns on the genotype level. When the recipient *D. magna* received a donor microbiome in which the *D. magna* were pre-exposed to *M. aeruginosa*, three out of the eight genotypes (K_MS, L_OM2, and L_T8) had a lower Shannon entropy when they received a sympatric than an allopatric microbiome. For two genotypes (K_BH and K_KP) the sequencing partly failed making it impossible to compare the sympatric and allopatric treatments, 24% of the OTUs were shared between the donor and the sympatric treatments and 22.7% of the OTUs were shared between donor, allopatric and sympatric treatments. In addition, 10.4% of the OTUs in the recipients were also present in the donor samples. This was very similar in the allopatric (10.8%) and in the sympatric (11.1%) conditions. Analysis of the Bray Curtis dissimilarity matrix, to investigate the difference in distance between donor and recipients across the treatments, showed a significant three-way interaction between microbiome type, diet and genotype ($F = 7.3, \text{df} = 2, p = 0.001$; Table SI12). Investigation per microbiome type shows that in the allopatric treatments there was a main genotype effect ($F = 5.05, \text{df} = 2, p = 0.01$), reflecting strong variation in the microbial community between the recipient genotypes (Fig. 7B). In the sympatric treatments this genotype effect depended on the donor diet (diet × genotype: $F = 6.5, \text{df} = 2, p = 0.003$). In the sympatric treatments, there was a main genotype effect in *D. magna* that received a donor diet not pre-exposed to *M. aeruginosa* ($F = 9.1, \text{df} = 2, p = 0.001$). However, when they received a donor diet pre-exposed to *M. aeruginosa*, this
genotype was no longer present ($F = 0.8$, df = 2, $p = 0.45$; Fig. 7B).

**Discussion**

We here performed a reciprocal gut transplant experiment, in which germ-free *D. magna* were inoculated with a sympatric or an allopatric gut microbiome that was pre-exposed to toxic *M. aeruginosa* or not. All inoculated recipients were exposed to *M. aeruginosa* and the tested *Microcystis* strain was a neutral lab strain that has not been interacting with the *D. magna* genotypes in nature. We found strong *D. magna* donor diet (pre-exposed to *M. aeruginosa* or not) × recipient genotype type (sympatric vs. allopatric) × recipient genotype interactions for the life history traits survival and fecundity. Independent of the tested region, more than half of the genotypes showed a higher survival (seven out of nine) or fecundity (five out of eight) when they received a sympatric vs. an allopatric microbiome, especially if the donor microbiomes were pre-exposed to *M. aeruginosa*. This pattern where most *D. magna* genotypes had a higher fitness, in terms of survival and fecundity, when they received a local (sympatric) than a foreign (allopatric) microbiome, is in agreement with what we expected. It has earlier been shown that in stressful environments, hosts may benefit from locally adapted microbes, as hosts may use specific microbial communities with large phenotypic effects to specialize and persist in novel niches [37–41]. However, because of the reciprocal transplant procedure, our study is unique by providing a causal link between fitness effects and local microbiomes. Note, however, that we studied local adaptation sensu lato, given that the adaptation is not at the level of a particular population, but rather on the level of the region, reflecting regional adaptation. In future work it would be very interesting to investigate these effects further on a microscale within a region.

For body size there was also a significant donor diet × donor microbiome type × recipient genotype interaction, and the interactions between donor diet and donor microbiome type, and between donor microbiome type and recipient genotype were significant. Body size was smaller in *D. magna* with a donor microbiome that had been pre-exposed to *M. aeruginosa* than to the control diet and this was especially so for the sympatric microbiomes. We found a marginally significant negative correlation between the delta values of the donor microbiomes (*M. aeruginosa—control treatment*) for survival and body size in the recipient *D. magna* genotypes. This indicates that *D. magna* genotypes that had a higher survival with a microbiome pre-exposed to *M. aeruginosa* were smaller. This suggests that carrying a microbiome that is associated with a higher tolerance upon *M. aeruginosa* exposure comes with a cost in terms of body size. A trade-off between growth and survival is shown in many organisms [72–74]. Here we show that this trade-off in *Daphnia* may be mediated through the microbiome, as also suggested by Callens et al. [58]. Hosts with particular microbiomes apparently divide energy towards growth or longevity, but not to the two traits simultaneously: they (1) grow slower with higher tolerance to stressful conditions, or (2) they grow faster without a higher tolerance [74].

There was a significant difference in α- and β-diversity in the donor vs. the recipient microbiomes. The donor microbiomes consisted mainly of Gammaproteobacteria and Bacteroidia (Flavobacteria). In the study of Macke et al. [42], *D. magna* genotypes that were tolerant to *Microcystis*, contained more Flavobacteria. Gammaproteobacteria and Flavobacteria have been described to be part of the mainly aerobic *D. magna* gut bacterial community [75]. Gamma-proteobacteria typically break down and ferment complex sugars and provide particularly important digestive roles [40, 76]. Members of the Flavobacteria group cause the lysis of *Microcystis* cells and degrade dissolved organic matter derived from intracellular products of *Microcystis* [77, 78]. The presence of Flavobacteria in the gut of *Daphnia* may therefore provide the host individual access to otherwise inaccessible nutrients in addition to detoxification of cyanotoxins. The recipient gut microbiomes were more diverse than the donor gut microbiomes. Next to Gammaproteobacteria and Bacteroida, the recipient gut microbiomes also contained bacteria from the following classes: Actinobacteria, Alphaproteobacteria, Bacilli, Planctomycetacia and Verrucomicrobiae. The recipient microbiomes were inoculated with the donor microbiomes for 48 h in sterile-filtered tap water in our transplant experiment. After these 48 h, the recipient *D. magna* were placed in a mixture of pond water from the region of origin, which may explain the relatively higher diversity and the presence of environmental bacteria in the gut microbiome of the recipient [79] *D. magna*. According to the Unionplot, 71.2% of the OTUs were shared between donors and recipients, showing that in the 48 h the recipients took up a substantial part of the bacteria from the pooled donor inoculum. The analysis of the Bray Curtis dissimilarity matrix, showed that *D. magna* that received a sympatric microbiome that was pre-exposed to *M. aeruginosa* had a more convergent microbiome across the genotypes. This was demonstrated by the distance between donors and recipients varying less across the genotypes in the sympatric treatments that were pre-exposed to *M. aeruginosa*, than across the genotypes in allopatric and sympatric treatments not pre-exposed to *M. aeruginosa* (Fig. 7B).

The host *D. magna* may follow two patterns with respect to the selective uptake of microbial strains to obtain an
increased tolerance upon toxic cyanobacterial exposure: (1) selection of a few adapted strains, or (2) selection for a high diversity of strains with complementary gene functions (Fig. 1). In this study, we found support for both patterns. Five of the eight genotypes (K_BH, K_MS, L_OM2, L_T7 and L_T8) had higher fitness traits (survival and/or fecundity). In three of these recipient genotypes (K_MS, L_OM2, L_T8) there was a lower Shannon entropy in the sympatric vs. allopatric donor microbiome when the donors were pre-exposed to *M. aeruginosa*. This suggests a selective uptake of a microbiome with beneficial functions, especially in stressful environments. This resembles to patterns observed in plants that recruit their microbiome in a host species-specific way and whereby the plant genotype affects the accumulation of micro-organisms that help the plant to defend itself, e.g., against pathogen attacks [80–82]. Selection of specific OTUs by these *D. magna* genotypes may lead to a lower chance of establishing other OTUs. This can be enhanced by priority effects of the first establishing bacterial strains leaving less space for other bacteria from the environment to enter and settle. These bacteria by interacting strongly with the host, will be strong competitors, and outcompete other incoming bacteria. In turn, this may then affect the host phenotype. However, two out of the eight genotypes (K_ZWE and L_T3) followed the second pattern, where both *D. magna* fitness (survival and/or fecundity) and bacterial diversity were higher in the sympatric than in the allopatric treatment. Hosts can also benefit from a more diverse microbiome. For example, it has been shown that mice with a natural, more diverse, microbiome had a higher fitness and a lower inflammation response to two diseases than mice with a laboratory, less diverse, microbiome [45]. It is possible that a more diverse community contains a wider array of metabolic capabilities, allowing for example a better growth [83]. Alternatively, the second pattern can also be explained through *Daphnia* genotypes being non-selective and picking up randomly a high diversity of (also non-adaptive) strains in stressful environments.

We conclude that most *D. magna* genotypes had a higher fitness in terms of survival and fecundity, when they received a sympatric than an allopatric microbiome. These fitness benefits from a local microbiome were, however, genotype dependent, which conforms the hypothesis that different *Daphnia* genotypes differ in their tolerance to *M. aeruginosa* [42] and different genotypes select different microbiomes [36]. As such the microbiome can help in structuring the wider freshwater bacterioplankton community through microbiome mediated eco-evolutionary dynamics [43, 84]. In some *D. magna* genotypes fitness was higher while bacterial diversity was lower for sympatric than for allopatric microbiomes, suggesting selection of certain microbiome strains. While in others both the fitness and bacterial diversity were higher in sympatric than allopatric treatments, suggesting selection of a high-bacterial diversity. Future research should confirm if particular adaptive strains interact with the host genotype to structure the microbial community in the *Daphnia* gut in stressful cyanobacterial environments.

**Data availability**

The datasets generated for this study can be found in the NCBI, under accession number PRJNA690081.

**Acknowledgements** Funding was provided by the KU Leuven research project C16/17/002 and the FWO project G092619N. We are grateful for the assistance of Anne Desreveaux, Maxime Tavernier and Marlies Van de Maele during the experimental work. We thank Isabel Vanoverberge for her guidance and assistance during the molecular work of this experiment and Wim Meert of the KU Leuven—UZ Leuven Genomics core for his help and feedback on the sequencing. We thank Lore Bulteel, Manon Coone, Martijn Callens, Emilie Macke, Luc De Meester and two anonymous reviewers for their stimulating discussions.

**Author contributions** SH and ED designed the experiment. SH performed the experiment and analyzed the data in concert with ED and RS. SH wrote the first draft of the manuscript. SH, RS, SM and ED participated in discussions and editing of the manuscript.

**Compliance with ethical standards**

**Conflict of interest** The authors declare no competing interests.

**Publisher’s note** Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.

**Open Access** This article is licensed under a Creative Commons Attribution 4.0 International License, which permits use, sharing, adaptation, distribution and reproduction in any medium or format, as long as you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons license, and indicate if changes were made. The images or other third party material in this article are included in the article’s Creative Commons license, unless indicated otherwise in a credit line to the material. If material is not included in the article’s Creative Commons license and your intended use is not permitted by statutory regulation or exceeds the permitted use, you will need to obtain permission directly from the copyright holder. To view a copy of this license, visit http://creativecommons.org/licenses/by/4.0/.

**References**

1. Kawecki TJ, Ebert D. Conceptual issues in local adaptation. Ecol Lett. 2004;7:1225–41.
2. Fox JW, Harder LD. Using a “time machine” to test for local adaptation of aquatic microbes to temporal and spatial environmental variation. Evolution. 2014;69:136–45.
3. Halbritter AH, Billeter R, Edwards PJ, Alexander JM. Local adaptation at range edges: comparing elevation and latitudinal gradients. J Evol Biol. 2015;28:1849–60.
4. Zhang M, Suren H, Holliday JA. Phenotypic and genomic local adaptation across latitude and altitude in *Populus trichocarpa*. Evol. 2019;11:2256–72.
Located adapted gut microbiomes mediate host stress tolerance
49. von Elert E, Martin-Creuzburg D, Le Coz JR. Absence of sterols constrains carbon transfer between cyanobacteria and a freshwater herbivore (Daphnia galeata). Proc R Soc B. 2003;270:1209–14.

50. Chen W, Song L, Ou D, Gan N. Chronic toxicity and responses of several important enzymes in Daphnia magna on exposure to sublethal microcystin-LR. Environ Toxicol. 2005;20:323–30.

51. Schwarzenberger A, Zitt A, Kroth P, Mueller S, von Elert E. Gene expression and activity of digestive proteases in Daphnia: effects of cyanobacterial protease inhibitors. BMC Physiol. 2010;10:6–20.

52. Cousyn C, De Meester L, Colbourne JK, Brendonck L, Verschuren D, Volckaert F. Rapid, local adaptation of zooplankton behavior changes in predation pressure in the absence of neutral genetic changes. PNAS. 2001;98:6256–60.

53. Lemaire V, Brusciotti S, van Gremberghe I, Veyver P, Vanoverbeke J, De Meester L, Genotype x genotype interactions between the toxic cyanobacterium Microcystis and its grazer, the waterflea Daphnia. Evol Appl. 2012;5:168–82.

54. Muniruzz S, Uthajakumar V, Arunkumar P, Ramasubramanian V. The effect of different feeds such as Chlorella vulgaris, Azolla pinnata and yeast on the population growth of Daphnia magna commonly found in freshwater systems. Int J Fish Aquac. 2016;4:05–10.

55. Guillard RRL, Lorenzen CJ. Yellow-green algae with chlorophyllide c. J Phycol. 1972;8:10–14.

56. Sezonov G, Joseleau-Petit D, D’Ari R. Escherichia coli physiology in Luria-Bertani broth. J Bacteriol. 2007;189:8746–9.

57. Moheimani NR, Borowitzka MA, Isdepak A, Singh HS, Standard Methods for Measuring Growth of Algae and Their Composition. In: Moheimani NR, Borowitzka MA, Isdepak A, Singh HS, editors. 5th ed. Netherlands: Springer; 2013. p. 265.

58. Callens M, Macke E, Muylaert K, Bossier P, Lievens B, Wauw M, et al. Food availability affects the strength of mutualistic host-microbiota interactions in Daphnia magna ISME J. 2016;10:911–20.

59. Callens M, Macke E, Muylaert K, Vanoverbeke I, Decaestecker E. Optimization of experimental methods for investigating host-microbiota interactions in Daphnia magna. Ch. 1. In: Environmental dependent effects of host-microbiota interactions in Daphnia magna. Callens M, editor. Belgium: PhD thesis Published Martijn Callens at KULeuven; 2017.

60. Callahan BJ, Sankaran K, Fukuyama JA, McMurdie PJ, Holmes SP. Bacterial and archaeal microbiome data analysis: from raw reads to community analysis. F1000Research. 2016;5:1492.

61. Callahan BJ, McMurdie PJ, Rosen MJ, Han AW, Johnson AJA, Holmes SP. DADA2: high resolution sample inference from illumine amplicon data. Nat Methods. 2016;13:581–3.

62. Langsrud O. ANOVA for unbalanced data: using type II instead of type III sums of squares. Stat Comput. 2003;13:163–7.

63. Fox J, Wiesberg S. Cox proportional-Hazards regression for survival data in R. An appendix to an R companion to applied regression. 1st ed. New York: SAGE Publications, Inc; 2002.

64. Bates D, Mächler M, Bolker B, Walker S. Fitting linear mixed-effects models using lme4. J Stat Softw. 2015. https://doi.org/10.18637/jss.v067.i01.

65. Curtin F, Schulz P. Multiple correlations and bonferroni’s correction. Biol Psychiatry. 1998;44:775–7.