Infection dynamics of the monogenean parasite *Gyrodactylus gasterostei* on sympatric and allopatric populations of the three-spined stickleback *Gasterosteus aculeatus*

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Abstract: Parasites with high host specificity maximally depend on their hosts, which should increase the likelihood of coevolution. However, coevolution requires reciprocal selection exerted by the host and the parasite, and thus a considerable level of parasite virulence. In species of the monogenean ectoparasite genus *Gyrodactylus* consecutive generations are confronted with a single host, which may constrain the evolution of virulence. Transmission, which is often important in the ecology of *Gyrodactylus* species, may have the opposite effect, but may also lead to the avoidance of coevolutionary arms races. We investigated the potential outcome of coevolution between *Gyrodactylus gasterostei* Gläser, 1974 and its host, the three-spined stickleback (*Gasterosteus aculeatus* L.) by determining the strength of genotype by genotype (*G*×*G*) interactions on two levels: within and between sympatric and allopatric host populations. To do so, we compared the parasite’s infection dynamics on laboratory-reared sympatric (Belgian) and allopatric (German) hosts. We found that a parasite line successfully infected a range of sympatric host genotypes (represented by families), while it failed to establish on allopatric hosts. Phylogeographic studies suggest that neutral genetic divergence between the host populations cannot explain this dramatic difference. Provided that this result can be generalised towards other parasite lines, we conclude that coevolution in this host-parasite system is more likely to lead to local adaptation on the population level than to *G*×*G* interactions within populations.

Keywords: Monogenea, *Gyrodactylus gasterostei*, ecological immunity, fish parasite, *Gasterosteus aculeatus*, host specificity, local adaptation, parasite-host coevolution

Resistance against parasites is shaped by the antagonistic interaction among hosts and parasites. This interaction may result in coevolution between host and parasite, producing patterns like local adaptation, genotype by genotype (*G*×*G*) interactions, and reciprocal selection (Woolhouse et al. 2002, Webster et al. 2004). All of these facets of antagonistic coevolution have been described in a number of host-parasite systems. For example, negative frequency-dependent selection causing Red Queen dynamics were driving local and temporal adaptation in host-parasite systems with direct transmission such as *Daphnia magna* and its microparasites (Ebert 1994, Decaestecker et al. 2007). Local adaptation was also observed in parasites with more complex life cycles (Ballabeni and Ward 1993, Lively and Dybdahl 2000). Similarly, reciprocal selection has been demonstrated in feather lice completing their life cycle directly on pigeon hosts (Clayton et al. 1999), but also in parasites with intermediate hosts such as *Schistosoma* species (Webster et al. 2004).

These examples demonstrate that coevolution is possible, irrespective of the parasite’s life cycle. The role of host specificity for local adaptation, on the other hand, has rarely been investigated. In this study, we were interested in coevolution of parasites characterised by high host specificity and a direct life cycle. Such parasites are confronted with few selective forces other than the ones imposed by their hosts, and this close interaction should maximise the likelihood of coevolution. Coevolution is most likely observed when a specialist pathogen exerts a strong selection pressure on its host and *vice versa* (Woolhouse et al. 2002). This can lead to strong *G*×*G* interactions and maintain host and parasite genotypic diversity by negative frequency-dependent selection. However, when consecutive generations of parasites are confronted...
with genetically similar hosts (as is often the case in host-specific parasites with a direct life cycle), coevolution may arise without the need for high virulence (Agrawal 2006). Relaxation of parasite selection for host counteradaptations might then lead to generalist strategies in parasites leading to comparable infection success on larger groups of related genotypes up to the whole population.

A group of parasites combining high host specificity with a direct life cycle is the flatworm genus *Gyrodactylus* (Monogenea, Platyhelminthes). *Gyrodactyli*ds represent common ectoparasites of fish species, living mainly on fins and gills. Embryos of *Gyrodactylus* species develop within each other’s uterus like Russian dolls, allowing relatively short generation times and exponential population growth (Cable and Harris 2002). Such features parallel those of directly transmitted microparasites, for which substantial theories on the likelihood and consequences of coevolution have been developed and tested (Woolhouse et al. 2002). In contrast, data on the interaction of *Gyrodactylus* species and their hosts at the micro-evolutionary level are scarce, and the consequences for coevolution are unknown. Boeger et al. (2005) suggested that transmission has the potential to minimize coevolutionary arms races in *Gyrodactylus* species and their hosts. Transmission, for instance by direct host contact, indeed plays a central role in the biology of *Gyrodactylus* species. Other studies have focussed on the susceptibility of different species and populations of fishes, as infection with *Gyrodactylus* species may have severe pathological consequences (Soleng and Bakke 1998, Bakke et al. 1999, 2002). Van Oosterhout et al. (2003) found marked variation in resistance for *Gyrodactylus* species between guppy populations facing high and low predation risk. Intra-specific variation in *Gyrodactylus* species is largely neglected in experiments. Given the economical damage of *Gyrodactylus* species in aquaculture (Nielsen and Buchmann 2001), empirical data on the evolution of virulence and coevolution are highly desirable.

In this study we explore the coevolutionary dynamics between *Gyrodactylus gasterostei* Gläser, 1974 and its principal host, the three-spined stickleback *Gasterosteus aculeatus* L. The *Gyrodactylus* community of sticklebacks in Europe includes several species, among which *G. gasterostei* and *G. arcuatus* are most commonly observed (Harris 1985, 1998). The epidemiology of *G. gasterostei* in Western Europe suggests narrow host specificity for the three-spined stickleback (Harris 1985, Raeymaekers et al. 2008a). Therefore, *G. gasterostei* can be considered a specialist and thus represents a suitable candidate to investigate the impact of host specialisation on the potential outcome of coevolution. Our experiment was carried out to compare the fitness (i.e., infection success and population growth) of an isogenic (i.e., derived from a single worm) *G. gasterostei* line on three laboratory-reared (i.e., immunologically naive) sibships from its sympatric Bel-

**MATERIALS AND METHODS**

**Source material.** The experimental infections were performed with a *Gyrodactylus gasterostei* line collected from a single three-spined stickleback stemming from Westerkerke (Belgium; 3°00’E, 51°10’N). This site represents a small eutrophic polder creek near the coast, with a very slow freshwater current and a high density of macrophytes. The stickleback population here, i.e., the sympatric host population, belongs to the lowland ecotype (Raeymaekers et al. 2005, 2007, Van Dongen et al. 2009; Fig. 1). *Gyrodactylus gasterostei* is the most common *Gyrodactylus* species on this host population (Raeymaekers et al. 2008a). A German stickleback population from Vierer See (Plön, Germany; 10°25’E, 54°09’N; Fig. 1), a lake draining into the Baltic Sea, was selected as the allopatric host population because of the common occurrence of *G. gasterostei* (Kalbe M., M.P.I.-Plön, Germany; pers. comm.).

The experiment was performed with the F₁ offspring derived from crosses from sympatric Belgian (B) or allopatric German (G) fish, caught in March 2003. These fish were disinfected with a 1:8000 formalin solution and crossbred in the laboratory in July 2003. This resulted in three sympatric F₁ sibships, coded B1, B2 and B3, and three allopatric F₁ sibships, coded G1, G2 and G3. After spawning, clutches were collected and kept in aerated glass jars until hatching, before fry were transferred to 20-l flow-through aquaria. These fish reached maturity in spring 2004 and were then used for our experiment (see below). Prior to the experiment, they were never exposed to the parasite and can therefore be regarded as immunologically naive.

**Experimental design.** The experimental design is shown in Fig. 2. All fish were kept individually (see below). The infection experiment was preceded by a worm breeding stage, in order to obtain sufficiently large numbers of worms. In spring 2004, the Westerkerke site was revisited to catch infected three-spined sticklebacks. These donor fish were transported alive to the laboratory, and their worms were transmitted within 1–2 days to naive fish of family B₁, i.e., one of the sympatric families. The required number of worms was reached by passaging one isogenic line (i.e., starting from a single worm) on four subsequent sets of randomly selected B1 hosts. Set sizes during the
four rounds were \( n = 1, 10, 32 \) and 39 sticklebacks, with infections lasting 20, 30, 24 and 30 days, respectively (Fig. 2). The isogenic line was initially selected out of 10 lines as the one with the best growth on the first host and was later confirmed to be \( G. \) gasterostei, based on the ITS rDNA region encompassing the Internal Transcribed Spacers (ITS1 and ITS2) and the small ribosomal subunit (5.8S rRNA) gene (Zietara et al. 2002).

After the last passage, an experiment was initiated to compare the success of the parasite line on sibship B1 with its success on the other sympatric sibships (B2 and B3), and on the allopatric \( F_1 \) sibships (G1, G2 and G3). On day zero of the experiment, 16 fish of each sibship were infected. The infections were organised such that the worms from each final worm breeding host were distributed equally over the six sibships (B1–B3, G1–G3). The experiment was evaluated by anaesthetising all fish weekly to count the number of worms.

**Infection and fish maintenance.** Fish used for worm breeding and experimental fish were briefly anaesthetized with 50 mg l\(^{-1}\) MS222 and infected by putting a \( G. \) gasterostei individual on the right pectoral fin, following standard methods of Cable et al. (2000). Subsequently, all fish were placed individually in 2-l aquaria, which were positioned randomly in a controlled cooling system at 12 °C and a 12 h light:dark photoperiod. Fish were fed three times a week a mix of brine shrimp \( Artemia \) salina nauplii and bloodworms (chironomid larvae). Water was changed weekly using dechlorinated tap water.

**Feeding activity.** Feeding activity was registered at the beginning of the infection experiment and 5 weeks later as an indicator of the health of the sticklebacks. It was measured as the number of feeding lunges made at \( Artemia \) salina nauplii during 1 min averaged over two trials on consecutive days. This test was performed on all experimental fish by placing them in their individual aquarium in a light-tight observation box, illuminated from above. After 5 min, a standardised number of nauplii (>1000 ind. l\(^{-1}\)) were introduced and feeding lunges were counted by a single observer (J.A.M. Raeymaekers). Feeding activity is considered to be a good health indicator in guppies (Van Oosterhout et al. 2003). We found this measure to be correlated between the two consecutive days (Pearson correlation = 0.51; \( P < 0.0001 \)). At the beginning of the experiment, there were no significant differences in feeding activity among populations (fixed effect; \( F_{1,90} = 3.29, P = 0.20 \)) or among sibships nested in population (random effect; \( F_{4,90} = 1.41, P = 0.24 \)). The average feeding activity here was 10.71 ± 6.64 lunges per minute.

**Data analysis.** Stickleback feeding activity after 5 weeks was evaluated as above, i.e., we investigated with a general linear model whether the infection experiment had induced differential feeding activity among populations (considered as fixed effect) or among sibships nested in population (considered as random effect).

Total worm load was calculated as the log-transformed sum of weekly infection intensities. This sum always included the initial worm, making the logarithmic transformation always defined. A general linear model was performed on total worm load to investigate the effect of population (considered as fixed) and the effect of sibship nested in population (considered as random). To assess epidemiological differences over time, weekly infection intensity was modelled with a generalised linear mixed model (GLMM; Molenberghs and Verbeke 2005), including time as a fixed factor. Infection intensity was assumed to be in-

![Fig. 1. Unrooted neighbour joining tree of Cavalli-Sforza and Edwards (1967) genetic distances (based on six microsatellite loci) among two experimental and seven neighbouring threespined stickleback \( Gasterosteus aculeatus \) populations, recalculated from Raeymaekers et al. (2007, 2008b) and Reusch et al. (2001). Experimental populations are Westkerke (observed heterozygosity \( H_o = 0.81 \)) and Vierer See (\( H_o = 0.64 \)). Belgian populations belong to the upper Scheldt drainage (S), the upper Meuse drainage (M), or the coastal lowlands (L). Vierer See belongs to the prevailing lake clade found in northern Germany (Reusch et al. 2001).](image1)

![Fig. 2. Graphical presentation of worm breeding and experimental design. Codes indicate \( Gasterosteus aculeatus \) population/genotype combinations. Populations are Westkerke (WK) and Vierer See (VS), and genotype codes refer to sibship B1, B2, B3, G1, G2 and G3, or wild individuals (W). Numbers in bold along arrows represent days of infection before passaging. Numbers in parentheses represent sample size of fish, which were kept individually in 2-l aquaria.](image2)
RESULTS

The experimental infections did not induce host mortality. After 5 weeks, fish from different sibships within populations displayed differential feeding activity (sibships nested in population: $F_{1,88} = 3.61, P = 0.0090$). In particular, sibship B1 showed a lower activity ($5.53 \pm 4.39$ lungen min.$^{-1}$) than sibship B2 ($10.94 \pm 7.26$ lungen min.$^{-1}$) and sibship B3 ($11.78 \pm 6.69$ lungen min.$^{-1}$). As will be seen below, these differences cannot be attributed to the experimental infection. Furthermore, fish from different populations did not differ in feeding activity ($F_{1,88} = 1.53, P = 0.28$).

Overall, the number of infected individuals gradually decreased from 77% after the first week towards 20% after 6 weeks. From the start of the experiment allopatric sibships showed a remarkably lower susceptibility compared to sympatric sibships (31% and 6% initial resistance, respectively). Infection intensity peaked after 5 weeks for sympatric sibships (average worm load $\pm$ S.E.: $17.81 \pm 3.45$ worms; max. 90) (Fig. 3), whereas maximal infection was already reached on allopatric sibships after the first week (average worm load $\pm$ S.E.: $1.46 \pm 0.21$ worms; max. 5).

Total worm load was significantly lower on the allopatric host population than on the sympatric host population, but did not differ between sibships within populations (Table 1; Fig. 4). A repeated Poisson regression analysis on weekly infection intensities generated a significant sibship by time interaction, pointing to a faster decline on sibship B1 at the end of the experiment ($F_{10,223} = 5.92, P < 0.0001$, Fig. 3). Differences in total worm load between the sympatric and the allopatric host population, quantified as $Q_{S\text{st}}$, dramatically exceeded neutral genetic divergence ($Q_{S\text{st}} = 0.96, 95\% \text{CI} = [0.68–0.99]$; $F_{\text{st}} = 0.14, 95\% \text{CI} = [0.08–0.21]$, Fig. 5). This indicates that selection on resistance of German hosts or strong local adaptation of the parasite in Belgium or a combination of both shaped this trait, rather than neutral divergence of the host populations. The observed differences in feeding activity, on the other hand, were rather caused by neutral divergence than selection, because the most probable value of genetic differentiation of this trait did not exceed neutral expectations (Fig. 5).

DISCUSSION

This study documents the results of an infection experiment comparing infectivity and infection intensity of the monogenean ectoparasite *Gyrodactylus gasterosteii* between its sympatric and an allopatric host population. The isogenic parasite line used here tended to show weak genotype by genotype (G×G) interactions on sympatric immunogenetically naive hosts, with hosts from the family used for worm rearing (B1) displaying lower infection intensities over time (Fig. 3). The main variation, however, was observed between the sympatric and the allopatric host population (Figs. 3, 4). The difference in total worm load between both populations was most likely caused by selection, as genetic differentiation for this trait ($Q_{S\text{st}}$) by far exceeded neutral genetic divergence (Fig. 5). Differences in feeding activity, on the other hand, showed no stronger differentiation than expected under neutral divergence, indicating that little adaptive variation is present for that trait. Wide confidence envelopes of $Q_{S\text{st}}$ estimates are not surprising, as only a very small number of families was used to assess genetic variation. The point estimate of highest probability for the difference in susceptibility nevertheless reflects a clear pattern of selection when reflected against the close clustering of stickleback populations from Belgium and northern Germany (including Vierer See) based on microsatellite (Fig. 1) and

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Table 1. ANOVA table of log-transformed total worm load on a sympatric and an allopatric three-spined stickleback *Gasterosteus aculeatus* population after 6 weeks of infection with an isogenic *Gyrodactylus gasterosteii* line. Host sibship (nested in population) was included as a random effect. Significant P-values are in bold.
**Fig. 3.** Temporal dynamics of *Gyrodactylus gasterostei* infection intensities on three sympatric Belgian (B1, B2, B3 – left panel) and three allopatric German (G1, G2, G3 – right panel) sibships of *Gasterosteus aculeatus*. Worms were initially reared on fish from sibship B1 (solid line, left panel). Error bars indicate standard errors.

**Fig. 4.** Average log-transformed total *Gyrodactylus gasterostei* load in three sympatric Belgian (B1, B2, B3) and three allopatric German (G1, G2, G3) sibships of *Gasterosteus aculeatus*. Worms were initially reared on fish from sibship B1 (shaded bar). Error bars indicate standard errors.

**Fig. 5.** Population differentiation between sympatric Belgian and allopatric German populations measured as $Q_{st}$ for susceptibility (total worm load) and life history (feeding activity) traits. Points show the highest posterior density of 1000 random samples from the posterior distribution and lines show 95% confidence intervals of the posterior distribution. Phenotypic differentiation is compared to neutral genetic divergence measured as $F_{st}$ (dashed line) and its 95% C.I. (shaded area).
mitochondrial (Mäkinen et al. 2006, Mäkinen and Merila 2008) phylogeographic studies. In summary, our results suggest that individual parasite lines are rather adapted to local host populations than to specific host genotypes.

Generalisation of these conclusions is difficult mainly for two reasons. Firstly, although this is the first study testing for differences in susceptibility for a Gyrodactylus species between sympatric and allopatric host populations, we only tested a single parasite line. It is possible that parasite lines are highly variable with respect to their specificity for host genotypes and host populations. Secondly, the experiment did not allow for a formal test of local adaptation, as we did not include the reciprocal sympatric and allopatric combinations (Kaltz and Shykoff 1998, Kawecki and Ebert 2004). Therefore, the strong genetic differentiation in resistance can be explained by higher immune competence of German hosts (Scharsack et al. 2007), local adaptation of the worm isolate to Baltic hosts, or a combination of both. It is remarkable here that the allopatric host appeared highly unsuitable, as most worms died in the first week, after giving birth only once. This observation requires further investigation.

Despite of these shortcomings, the experiment nevertheless suggested that the virulence of G. gasterostei is comparatively low. Worms induced no host mortality, and host feeding activity was not related to worm load. Low pathogenicity seems to be characteristic of G. gasterostei, even when infecting naive three-spined stickleback populations (de Roij et al. 2011). However, virulence may also be highly dependent on environmental conditions (Wagner et al. 2008), and will not necessarily be expressed under benevolent laboratory conditions. In the case of Gyrodactylus species, population growth rate is particularly sensitive to environmental factors (Bakke et al. 2002), the hormonal status of the host (Harris et al. 2000) and food availability (Kolluru et al. 2006). In the case of G. gasterostei, it has been shown that populations of worms grow larger on weak or stressed hosts than on healthy fish (de Roij et al. 2011). While some congeners can be highly virulent (e.g. G. salaris; Bakke et al. 1999), virulence of G. gasterostei in good conditions seems not sufficient to drive the evolution of host defences.

Next to virulence, the presence of genetic variation in host susceptibility and parasite infectivity is another prerequisite for coevolution. In general, fishes have been shown to exhibit heritable variation or apparent differences in susceptibility to Gyrodactylus species (Madhavi and Anderson 1985, Bakke et al. 1999, Van Oosterhout et al. 2003). We now know that variation in resistance to G. gasterostei is present among host populations (de Roij et al. 2011; this study). Furthermore, a potential genetic basis for variation in resistance at the individual level has been identified by showing an association between the prevalence of G. gasterostei and a single class IIb allele of the major histocompatibility complex in a wild, riverine stickleback population (Eizaguirre et al. 2009). In our experiment, the breeding of the parasites was performed in a way that maximised the interaction with a single sympatric host sibship. Interestingly, the isolate tended to show lower infection intensities on this host sibship than on other sympatric host sibships. However, the differences were rather small, and since there were no directional changes in infection intensity during the four rounds of worm breeding (data not shown), these results suggest that G×G interactions within populations were rather weak when infected with this worm isolate. Such generalist strategy of the parasite facilitates parasite transmission, as unsuitable hosts will be rare. If these results also apply to other parasite isolates, then coevolution fuelled by G×G interactions might be of lesser importance in this host-parasite system.

Conclusion

In contrast to macroparasites with complex life cycles, consecutive generations of Gyrodactylus species are confronted with a single host genotype. Hence, Gyrodactylus species are expected to reach a balance between its need to reproduce and the cost of harming its host – similar to vertically transmitted parasites (Ebert and Herre 1996). We demonstrated that the fitness of laboratory-reared three-spined sticklebacks was largely unaffected following infection with an isogenic line of the monogenean parasite G. gasterostei. With such a low virulence, reciprocal selection pressures driving coevolutionary processes may be rather weak. Furthermore, the infections revealed small differences in susceptibility within host populations, but strong differences between host populations. Provided that this result can be generalised towards other parasite lines, we conclude that coevolution in this host-parasite system is more likely to lead to local adaptation on the population level than to G×G interactions within populations. Such a specificity level is in agreement with the central role of transmission in the ecology of Gyrodactylus species (Boeger et al. 2005). In general, Gyrodactylus species differ considerably in the level of host specificity, transmission capacity, and reproductive strategies (Harris 1993). These characteristics may influence the evolution of virulence, and may vary among closely related Gyrodactylus species on a single host species. Further exploration of this group of parasites, especially on small vertebrate hosts with reasonably short generation times like sticklebacks, guppies and gobies, may reveal how these characteristics affect the chances for coevolution to occur.

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