Ecological immunogenetics of life-history traits in a model amphibian

Seth M. Barribeau1,†, Jandouwe Villinger2,† and Bruce Waldman3,4,*

1Experimental Ecology, Institute of Integrative Biology, ETH Zürich, Zürich 8092, Switzerland
2Molecular Biology and Bioinformatics Unit, International Centre of Insect Physiology and Ecology, PO Box 30772-00100, Nairobi, Kenya
3Department of Ecology, PO Box 84, Lincoln University, Canterbury 7647, New Zealand
4Laboratory of Behavioral and Population Ecology, School of Biological Sciences, Seoul National University, I Gwanak-ro, Gwanak-gu, Seoul 151-747, South Korea

†These authors contributed equally to the study.
*Author for correspondence (waldman@snu.ac.kr).

The vertebrate major histocompatibility complex (MHC) encodes cellular recognition of pathogens and influences mating and social behaviour. MHC genotypes of individuals determine immunorecognition of different pathogen repertoires [5]. Host-specific immunological responses shape individuals’ microbiota, both commensal and parasitic [6,7]. These, in turn, produce cues used for social recognition [8]. Mating preferences for MHC-dissimilar partners should facilitate inbreeding avoidance and make offspring less vulnerable to disease by increasing their heterozygosity or by shuffling MHC alleles to confer protection against pathogens that successfully exploit parental genotypes [5]. Social preferences for MHC similarity facilitate cooperation among kin. Examples include communal nesting partner preferences in mice [9] and schooling preferences in tadpoles [10]. Thus, social behaviour both influences and is influenced by the micro-organisms with which individuals live [11].

When associating with MHC-similar conspecifics, individuals may benefit from exposure to low-virulence micro-organisms that are adapted to their immune system [6]. However, new pathogens may pose a significant risk by exploiting shared weaknesses in group members’ immune systems and thus spread rapidly. In contrast, associating with MHC-dissimilar conspecifics may expose individuals to highly virulent micro-organisms. Whether MHC-biased social behaviours are adaptive cannot be fully understood without first determining their immunological costs and benefits, both affecting the individual itself and the conspecifics with which it interacts [12].

To determine the fitness consequences of MHC-based association, we exposed Xenopus laevis tadpoles to water conditioned by adult conspecifics. Tadpoles and adults of this species share the same habitat and microbial environment. MHC loci regulate both immunity to bacteria [13] and association preferences among tadpoles [10]. We assessed three life-history traits—growth, development and survivorship—as a function of the MHC similarity between adults and tadpoles.

2. MATERIAL AND METHODS

We bred X. laevis with known MHC class I and class II sequences (haplotypes f, g and j [14,15]; GenBank accession numbers: class I AF185580, AF185579 and AF185586; class II AF454374, AF454377 and AF454376). The frogs originated from the Basel Institute for Immunology and had been bred for several generations in our laboratory to maintain shared background genetic variation in frogs homozygous for different MHC haplotypes. All frogs were maintained in 60 l holding tanks into which water was fed continuously (3 l h−1). The flow-through system was fed by filtered, aerated water (21 °C) sourced from a deep aquifer.

We mated six adult MHC-homozygous frogs (ff, gg, jj) twice in one night to obtain tadpoles of six MHC genotypes (ff, fg, fj, gg, gj, jj) [13]. We induced oviposition by injecting the females into the dorsal lymph sac with 100 IU of luteinizing hormone-releasing hormone (Argent Chemical Laboratories, Redmond, WA, USA). MHC-identical frogs were paired first, after which amplexed adults were separated and paired with MHC-dissimilar partners. After eggs hatched, we placed 100 tadpoles from each brood into 10 l tanks.

Subsequently, we isolated the maternal frogs, and two additional MHC heterozygous females (fg, gj), in 60 l flow-through tanks and fed them sliced ox liver every 2 days. After 4 days, we stopped the water flow to allow chemical cues and micro-organisms to accumulate, and ceased feeding the frogs to limit fouling. To obtain an estimate of microbial density, we plated 10 µl of the water from each tank in triplicate onto tryptone soya agar (Oxoid, Basingstoke, UK), incubated the plates aerobically at 32 °C, and counted the number of bacterial colonies at 24 and 48 h. We exposed tadpoles

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to water from frogs, standardized by dilution to the microbial concentration of the lowest number of colonies found (gg female: 250 cfu ml$^{-1}$).

Two weeks after hatching, we separated 300 tadpoles, 50 of each genotype, individually into beakers containing 11 of conditioned water. Ten tadpoles of every genotype ($f_1, f_2, b_1, b_2, g_1, g_2, b_1$) were exposed to water from each of $f_2, f_2, g_2, g_2, b_1$ and $b_1$ adults. We randomly assigned the tadpoles into six blocks. We fed tadpoles every 2 days with ground nettle suspension, topped the water up to 1 l every 4 days to replenish evaporation, and moved each beaker one place each day to limit position effects. Three weeks after exposure, we recorded developmental stage [16] and photographed each tadpole from 60 cm above to measure snout–vent length (SVL) using ImageJ v. 1.3 (NIH, Bethesda, MD, USA).

We analysed how tadpole mortality varied with the number of shared MHC haplotypes ($0/1/2; n = 110/40/50$) and block by binomial generalized linear models (GLM) using R v. 2.10 (www.r-project.org). We examined how tadpole development varied in response to the percentage of shared amino acids at MHC class I and II loci peptide-binding region (PBR) domains [14,15] by quadratic regressions with tadpole SVL as a covariate. Developmental stage was analysed with Gaussian GLMs including SVL as a covariate. We also analysed the effects of relatedness of frogs (whether water was conditioned by the maternal parent), and the ratio by which conditioned water was diluted, on tadpole size, developmental stage and mortality in additional GLMs.

3. RESULTS

Tadpole mortality decreased with the number of MHC haplotypes shared with frogs that had conditioned the water (figure 1; $\chi^2 = 4.23, p = 0.04$). Three times as many tadpoles died when exposed to water conditioned by frogs with which they shared no haplotypes compared with when they shared one haplotype. No tadpoles died when reared in water conditioned by MHC-identical donor frogs (figure 1).

Tadpole development slowed in proportion to the number of shared MHC haplotypes ($F_{1.291} = 4.31, p = 0.039$) and PBR amino acid sequence similarity (figure 2a), class I: $t_{289} = 3.09, p = 0.0022$, overall $F_{5.289} = 4.15, p = 0.0067$; class II: $t_{289} = 4.92, p < 0.0001$, overall $F_{5.289} = 10.80, p < 0.0001$). With increasing PBR sequence similarity, development at first slowed to a minimum (95% similarity, MHC class II) and then increased (figure 2a,b).

Tadpole size varied independently of developmental stage ($F_{1.290} = 0.07, p = 0.80$) and the number of shared MHC haplotypes ($F_{1.291} = 0.62, p = 0.29$). Nor did tadpole size explain stage differences in our regression analyses (class I: $t_{289} = 0.33, p = 0.74$; class II: $t_{289} = 0.46, p = 0.65$). Kinship cues and dilution factor did not significantly influence development (relatedness: $F_{1.291} = 0.85, p = 0.36$; dilution: $F_{1.291} = 0.11, p = 0.74$), growth (relatedness: $F_{1.291} = 0.18, p = 0.67$; dilution: $F_{1.291} = 1.75, p = 0.19$) or survival (relatedness: $\chi^2 = 0.11, p = 0.74$; dilution: $\chi^2 = 0.46, p = 0.50$).

4. DISCUSSION

More tadpoles died in water conditioned by MHC-dissimilar than MHC-similar frogs, but surviving tadpoles developed faster. Tadpoles exposed to cues from conspecifics dissimilar to themselves in MHC type suffered reduced fitness. These results demonstrate that the MHC identity of conspecifics with which individuals interact not only influences the social behaviour but also life-history traits of group members.

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causing tadpoles to approach metamorphosis at smaller sizes typically occurs in unfavourable conditions. In *Xenopus*, more rapid metamorphosis also can be induced by exposure to predators, resulting in smaller juveniles [18]. In other anurans, such individuals suffer a reduced likelihood of survival and reproduction [19]. Adults may be more resistant to infection than tadpoles, which might favour rapid metamorphosis in some conditions. However, immunity is suppressed during metamorphosis, hence vulnerability to disease is then high [20].

Selection for a social environment with a compatible microbiota may explain the preferential association of tadpoles with others sharing the same MHC alleles [10]. MHC-similar individuals are more likely to transmit locally adapted pathogens that express low virulence. Individuals thereby might benefit from avoiding novel pathogens associated with immunologically dissimilar conspecifics. The advantages of preferential association with immunogenetically similar conspecifics might extend beyond pathogens to confer protection against other types of parasites as well [21]. However, because costs and benefits are environmentally modulated, we do not expect MHC-biased social assortment to be invariably selected nor expressed.

In sub-Saharan Africa, adult and larval *X. laevis* often live in turbid water. Little is known about their social behaviour in the wild [22], although adults cannibalize tadpoles in the laboratory. Thus, we cannot be certain of the functional significance of our results. Moreover, we determined neither the microbial composition nor the proportion of resident and transient microbiota [7] of water conditioned by adults. Resident microbiota would be more likely to be adapted to host frogs’ immune systems than transient, possibly pathogenic, organisms.

Social groups are dynamic; individuals are exposed to a changing array of micro-organisms that vary with group composition and shifting environmental factors. Our results suggest that immunogenetic identity is important in assessing the costs and benefits of sociality. How microbiota associated with specific MHC genotypes, spanning the full spectrum from mutualistic to pathogenic organisms, affect the fitness of hosts and their conspecifics remains fertile ground for future work.

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