Genetic variability in the tolerance of natural populations of *Simocephalus vetulus* (Müller, 1776) to lethal levels of sodium chloride

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Abstract – Using several clonal lineages of *Simocephalus vetulus* (Cladocera, Daphniidae) as a random sample, we investigated the genetic component of the halotolerance of one brackish and two freshwater populations of this littoral filter feeder. We hypothesized that genotypes from the brackish population were more tolerant than freshwater ones, via adaptation to local environmental conditions. Clonal identity was established by a cost-effective molecular fingerprinting technique (microsatellite-primed polymerase chain reaction (MSP-PCR)). Two distinct methodologies were used to assess cladoceran sensitivity to synthetic-grade sodium chloride (NaCl): (i) standard 48-h acute assays and (ii) 12-h survival time (ST) trials. No correlation was found between acute EC₅₀ and ST values. The sensitivity of brackish and freshwater clones was comparable in terms of acute EC₅₀ (varied from 2.28 to 3.83 g.L⁻¹). On the contrary, genetically determined differential tolerance to NaCl among populations was found for ST: all brackish genotypes, except one, were more resilient (ST > 120 min) than freshwater clones (ST < 120 min). Bearing in mind that these results were obtained with isolates from the extant population, it is surprising that the range of acute sensitivity of the freshwater and brackish genotypes was similar, and that the only difference between them was the ability of brackish clones to survive longer under high salinity stress (6 g.L⁻¹, in ST trials). We must conclude that the effect of salinity (original environment context) on the selection of genotypes was weaker than we had expected and than other authors have shown for other stressors.

Key words: Genetic variability / genotype–environment interaction / sodium chloride / halotolerance / *Simocephalus vetulus*

Introduction

Saline intrusion in freshwaters has received considerable attention (e.g., Schallenberg et al., 2003; Sarma et al., 2006; Gonçalves et al., 2007; Nielsen et al., 2008), justified by the predictions of sea level rise and observation of coastal erosion, which threaten coastal lakes (IPCC, 2008). In coastal lakes that face regular (e.g., tidal) or sporadic salinity increases, freshwater organisms must cope with potentially lethal levels of salinity for short-term periods (hours to days). Particularly for large cladocerans (such as *Daphnia* and *Simocephalus*), their ability to survive (or not) these salinity pulses may modulate zooplanktonic taxa succession (Jeppesen et al., 1994, 2007; Schallenberg et al., 2003; Nielsen et al., 2008) and a lake regime shift (see Jeppesen et al., 2007).

Although sensitive to minor salinity increases (Schallenberg et al., 2003; Sarma et al., 2006; Gonçalves et al., 2007), freshwater cladoceran populations also occur in coastal systems with distinct degrees of connectedness to the sea (e.g., Jeppesen et al., 1994; Schallenberg et al., 2003; Marques et al., 2006; Santangelo et al., 2008). Such physiological plasticity is likely due to genetic factors associated with osmoregulation shifts (Arné and Koivistio, 1993; Aladin and Potts, 1995; Martínez-Jerónimo and Martínez-Jerónimo, 2007). The degree of phenotypic plasticity to environmental fluctuations is naturally framed within the populations’ genetic variability, which is greatly influenced by founder effects and locally acting selective forces (De Meester, 1996a, 1996b; Boersma et al., 1999;
De Meester et al., 2002). Many studies have shown that locally acting selective forces (particularly predation) lead to the formation of local races or “ecotypes” (local adaptation, see Weider and Hebert, 1987; De Meester, 1996a, 1996b; Boersma et al., 1999; Cousyn et al., 2001; De Meester et al., 2002). This has also been reported in Daphnia populations from brackish vs. freshwater environments (Weider and Hebert, 1987; Teschner, 1995; Barry et al., 2005).

In this study, Simocephalus vetulus (Müller, 1776) (Cladocera, Daphniidae) populations inhabiting coastal lakes were used as an experimental model. Our interest was focused on a population inhabiting a brackish lagoon in the South of Portugal, which had to cope with salinities close to 2. Our main goal was to determine whether there is a genetic component of salinity resistance in this population. Since these populations are subjected to salinity pulses, due to the heterogeneous and challenging habitat they live in, they must be able to cope with short-term lethal levels of salinity. Therefore, our aim was to compare the acute halotolerance with salinity (using NaCl as a proxy) of this brackish population with that of freshwater populations of S. vetulus. To do so, two distinct methodologies were employed: (i) standard acute assays (OECD, 2004), complemented with (ii) survival time (ST) trials (adapted from Ribeiro et al., 2000; Lopes et al., 2005). We hypothesized that there should be a population differentiation pattern in terms of acute halotolerance, with brackish genotypes being more resilient to NaCl than freshwater ones.

Studies concerning genetic differentiation in quantitative traits of cladocerans have been conducted in daphniids (Cladocera, Daphniidae) living in contaminated vs. uncontaminated habitats (e.g., Lopes et al., 2005; Muyssen et al., 2005) and fish vs. fishless lakes (e.g., De Meester, 1996a; Boersma et al., 1999). Such studies rely on the parthenogenetic mode of reproduction, short life cycle and ease in laboratory culturing of cladocerans (Muyssen et al., 2005), which assure continuous rearing of progeny bearing the genetic information of the founding mothers (De Meester, 1996a; Pfrender and Lynch, 2000). By maintaining clonal lineages under similar laboratory conditions for several generations, environmental and maternal effects can thus be eliminated.

Material and methods

Cultures and test organisms

S. vetulus (Müller, 1776) occupies a different ecological niche from most daphniids, due to its habitat preferences and swimming behaviour (Alonso, 1996). It is a macrophyte-associated filter feeder and an important regulator of primary production and nutrient cycling in littoral habitats (Hann, 1995; Alonso, 1996). Clonal lineages of S. vetulus were established from three extant populations, which were sampled with oblique trawls of plankton net. Two of these populations came from coastal freshwater ecosystems in central Portugal, namely a small shallow lake (Lagoa de Mira – LM; lat. 40°26′29″N, long. 8°45′09″W) and a nearby semi-artificial ditch or canal (Vala da Fervença – VF; lat. 40°25′36″N, long. 8°45′19″W). Although there is hydrological connectivity between these systems and the Ria de Aveiro estuary (i.e., indirect connection with the marine environment via a semi-artificial system of small ditches and channels), salinity does not reach upstream due to a system of small dams and dikes. Thus, there is low dissolved solids concentration and conductivity is typically <500 μS.cm⁻¹. The third population originated from inhabitants of a brackish lagoon (Lagoa de Melides – M; lat. 38°08′01″N, long. 8°46′52″W). This coastal lake has an intermittent connection with the sea (usually once or twice a year, either by human or natural action). Samples were taken at the farthest point from the sea entrance, where salinity was 2.3.

Field-collected samples were immediately brought to the laboratory, where a random subsample of ovigerous mothers was isolated from each lake and transferred to 50 mL glass beakers with lake water. As a starting point, we assumed each mother to represent a putatively distinct genotype, until genetic fingerprinting tools clarified clonal identity (see below). Asexually produced newborns (F₀ generation) from these mothers were used as starting inocula of monoclonal cultures. The genotype collections of S. vetulus were reared in moderately hard reconstituted water (pH 7.4–7.8, hardness 80–100 mg.L⁻¹ CaCO₃, alkalinity 57–64 mg.L⁻¹ CaCO₃), comprised of: 123 mg.L⁻¹ MgSO₄·7H₂O, 96 mg.L⁻¹ NaHCO₃, 60 mg.L⁻¹ CaSO₄·2H₂O, and 4 mg.L⁻¹ KCl, sensu ASTM (1980) and USEPA (2002). This synthetic medium was supplemented with 4 mL.L⁻¹ of a standard organic additive (algal extract as described by Baird et al., 1989) and vitamins (as described for M4 and M7 medium – Elendt and Bias, 1990). Cyclically, neonates from the third or fourth brood were isolated and their mothers discarded, thus providing continuous culture renewal (F₁, F₂, F₃, ...) for at least 10 generations. This provided a required refractory period, ensuring normalization of environmental and maternal influences; both freshwater and brackish genotypes were cultured in the described freshwater reconstituted medium. Cultures were reared under a temperature of 20 ± 2 °C and a 16 h:8 h photoperiod, and organisms were fed three times a week (Monday, Wednesday and Friday) with a Pseudo-kirchneriella subcapitata ration of 1.5 × 10⁵ cells.mL⁻¹ (for more information on algal culture and ration, see Antunes et al., 2004). Vessels with acclimated animals were used as stock cultures for all subsequent experiments.

Clonal identification

Microsatellite-primed polymerase chain reaction (MSP-PCR) or inter-simple sequence repeats (ISSR) is a fast and simple genotyping technique that uses repetitive
and highly variable regions of the genome (microsatellites) to design universal primers, which produce highly variable fingerprinting profiles (Zietkiewicz et al., 1994; Karp and Edwards, 1997; Abbot, 2001). This technique has shown to be useful in studies on systematics and in the differentiation of strains and breeds within the same species (e.g., Alves et al., 2007) and within cell lineages from the same individual (e.g., Grasela and McIntosh, 2003).

DNA was extracted individually from preserved specimens (70–80% ethanol) of each population using a standard protocol (Schwenk et al., 1998), which has undergone minor adjustments over time (see Brede et al., 2006). To maximize extraction, only adult females were used and a preliminary step consisting of specimen maceration was included in the extraction procedure. All monoclonal cultures successfully established in the laboratory were analysed, in a total of 54 putative lineages (18 per population).

Each MSP-PCR reaction was performed in a total volume of 24 μL, containing 2.0 mM MgCl₂, 0.2 mM dNTP, 1x PCR buffer, 0.3 μM primer, and 1 unit of DNA polymerase (Go Taq Flexi DNA polymerase, Promega). Three primers [(GTG)₅, (CAG)₆, and (ACC)₆] were chosen to discriminate genotypes, after preliminary testing on PCR conditions and reproducibility of a wider battery of primers (see Abbot, 2001; Grasela and McIntosh, 2003; Dusinsky et al., 2006; Alves et al., 2007). Cycling conditions were as follows: (i) a denaturation step consisting of one cycle of 120 s at 95 °C; (ii) 35 cycles of 45 s at 94 °C, 45 s at 50° or 57 °C (depending on the primer used) and 90 s at 72 °C; (iii) a final elongation step of 10 min at 72 °C. Primer (GTG)₅ required an annealing temperature of 50 °C, while (CAG)₆ and (ACC)₆ were amplified at 57 °C. PCR products were separated on 1.5% agarose gels (run at 90 V for 4 h) and visualized under UV light, after immersion of gels in an ethidium bromide solution for 10 min. A Promega loading buffer (Go Taq Flexi) was used and a 100 bp molecular marker (100–1500 bp) was loaded in the gel as a reference.

The analysis of genetic variability among individuals was performed by comparison of the banding patterns obtained. The patterns were converted to a bidimensional binary matrix (1 = presence of band, 0 = no band). Samples were always run twice (amplification and fragment separation) and bands were only scored “1” if present in both amplifications. From this initial matrix, we derived similarity matrices using the Jaccard’s coefficient (Culley, 2005), which allowed us to build dendrograms reflecting the genetic architecture of populations sampled. The dendrograms were constructed using unweighted pair group method with arithmetic mean (UPGMA). All analyses (gel and fingerprint data) were performed with GelCompar II (Applied Maths, USA), using its default optimization and tolerance levels. We used a conservative cut-off value (> 30% dissimilarity) to differentiate clonal lineages, based on previous optimization of PCR consistency.

<table>
<thead>
<tr>
<th>Assays</th>
<th>[NaCl] (g·L⁻¹)</th>
<th>Salinity</th>
<th>Conductivity (μS·cm⁻¹)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Acute assays</td>
<td>0.0</td>
<td>0.0 ± 0.00</td>
<td>216.4 ± 44.73</td>
</tr>
<tr>
<td>2.0</td>
<td>2.1 ± 0.06</td>
<td>3650 ± 100.0</td>
<td></td>
</tr>
<tr>
<td>2.4</td>
<td>2.5 ± 0.00</td>
<td>4323 ± 40.4</td>
<td></td>
</tr>
<tr>
<td>2.9</td>
<td>3.0 ± 0.06</td>
<td>5127 ± 41.6</td>
<td></td>
</tr>
<tr>
<td>3.5</td>
<td>3.7 ± 0.06</td>
<td>6047 ± 35.1</td>
<td></td>
</tr>
<tr>
<td>4.2</td>
<td>4.4 ± 0.10</td>
<td>7157 ± 127.4</td>
<td></td>
</tr>
<tr>
<td>5.0</td>
<td>5.2 ± 0.17</td>
<td>8333 ± 203.1</td>
<td></td>
</tr>
<tr>
<td>Survival-time</td>
<td>6.0</td>
<td>6.3 ± 0.06</td>
<td>9977 ± 98.2</td>
</tr>
</tbody>
</table>

*The contribution of the test medium in terms of Na⁺ and Cl⁻ was assumed to be unimportant, given the small amount of these ions in its composition (see text).*

Halotolerance assays

Standard acute assays were conducted by exposing distinct *S. vetulus* clonal lineages (M63, M48, M66, M13, M7, M50, M20, M36, LM2, LM38, LM5, LM48, LM42, LM41, LM64, LM25, VF7, VF35, VF18, VF70, VF29, VF48 and VF69) to NaCl aqueous solutions, following standard protocols (ISO, 1996; USEPA, 2002; OECD, 2004). A stock solution (50 g·L⁻¹) was prepared by dissolving analytical grade NaCl in moderately hard reconstituted water. Test solutions (Tab. 1) were then obtained by diluting the appropriate volumes of stock solutions in the synthetic medium. Salinity and conductivity measurements were performed with a WTW conductivity meter. A fixed volume (10 mL of test solution) was added to the corresponding test vessels. Neonates (<24 h old and born between the 3rd and the 5th brood) were exposed to a geometric sequence of NaCl concentrations (2.0–5.0 g·L⁻¹; Tab. 1), under the same conditions of rearing cultures. The range of concentrations used was selected based on preliminary tests and toxic levels reported in the literature (e.g., Gonçalves et al., 2007). Our goal was to find a range of concentrations which could be applied to all clones. Four replicates per treatment were used and five neonates were randomly assigned to each 10 mL replicate (i.e., a total of 20 organisms per test concentration were used). In all tests, a negative control was used, where no NaCl was added (nominal concentration 0.0 g·L⁻¹; Tab. 1). Vessels were screened for immobilized daphnids after a 48 h exposure period. During these acute exposures, neither organic extract nor food was added.

ST trials were conducted with the same clonal lineages as the standard acute assays; experimental procedures were adapted from Ribeiro et al. (2000) and Lopes et al. (2005). Neonates (less than 24 h old and born between the third and the fifth brood) were exposed to a fixed concentration of NaCl (6.0 g·L⁻¹), which is normally lethal within 12 h. Salinity and conductivity measurements were also performed (Tab. 1). Assays were performed in 24-well
cell culture plates, where six organisms of each clone were individually exposed to 2 mL of test medium (moderately hard reconstituted water spiked with NaCl). Four genotypes could thus be tested in each plate. For each clone, trials were performed in triplicate, with each replicate consisting of neonates produced in different days or from different culture flasks of the same clone. Organisms were observed every 15 min from the beginning of the trial until 180 min, every 30 min from 180 to 360 min, every 1 h from 360 to 720 min, and, if necessary, every 2 h after 720 min (12 h). Similarly to the standard acute assays (see above), we used immobilization as the trial endpoint. Organisms were scored as immobile under the stereoscope (8–10× magnification) if they were not able to swim after gentle and repeated prodding. Trials ended when all organisms were considered as immobilized in two consecutive observations. In parallel to all ST trials, controls (neonates exposed to unspiked culture medium) were performed, following the same experimental design; trials were valid only if no immobilized individuals were recorded in the control at the end of the test.

**Statistical analysis**

Acute toxicity was expressed as 48 h immobilization EC₅₀ values (and respective 95% confidence intervals), as estimated with Probit analysis using Minitab® software. Average ST for each replicate and corresponding acute EC₅₀ values was integrated in the computation of a halotolerance index (HI):

\[
HI = 100 \times \text{relST} \times \text{relEC}_{50},
\]

where relST and relEC₅₀ represent the relative ST or EC₅₀, i.e., the value of the clone under consideration divided by the maximum of all clones. This normalization step was introduced in the calculation to give similar weight to both variables, while allowing HI to vary between 100 (extreme tolerance) and close to 0 (extreme sensitivity). This weighted metric is thus a simple way to order genotypes in terms of their sensitivity.

ST was analysed for differences between populations, following a nested ANOVA design (mixed-effects model), with population as the fixed factor and clone (nested within population) and replicate trial (nested within clone) as the random factors. Individual ST measurements constituted the error term of the ANOVA. No replicates per clone were available for EC₅₀ values, so a one-way ANOVA was used to assess differences among Simocephalus populations, using clone as the error term. All analyses were computed using Minitab® software, which also allowed extracting variance components for each source of variation.

**Results**

MSP-PCR band profiles allowed the differentiation of the three populations (Fig. 1). The only exception was a
subset of six LM genotypes, which was associated with the M population cluster. For this study, 23 clonal lineages (eight clones for M, eight clones for LM and seven clones for VF) were used in the experiments (see list in Table 2), after being scored as genetically distinct genotypes (Fig. 1; the most similar genotypes used were M20 and M36). Genetic identity was confirmed by additional dendrograms for each population (data not shown), using a predetermined cutoff level of 70% similarity. This allowed a security margin in the discrimination of clones.

The acute immobilization EC50 values (Table 2 and Fig. 2) ranged from 2.28 to 3.83 g·L⁻¹, thus demonstrating a large range of halotolerance in the tested genotypes. Comparing clone sensitivity within each population, the EC50 values obtained ranged from 2.48 to 3.83 g·L⁻¹ in population M (brackish lagoon), 2.81 to 3.59 g·L⁻¹ for population LM (freshwater lake), and 2.28 to 3.60 g·L⁻¹ in population VF (freshwater ditch). These ranges show an overlap of the values of halotolerance between populations (see distribution of clones in the x-axis of Fig. 2), confirming no differentiation between populations (one-way ANOVA: d.f. = 2, 20; \( P = 0.48; \ P = 0.63 \)). Comparing all clones, the lowest (2.28 g·L⁻¹) EC50 was observed in VF (a freshwater population), while the highest values (3.83 and 3.65 g·L⁻¹) were recorded in clones from the brackish population (Table 2 and Fig. 2).

The EC50 values were not consistent with the halotolerance of clones in ST trials. In fact, the correlation between these variables (Fig. 2) was weak and not significant (\( r = -0.083; \ P = 0.71 \)). A good example of this lack of correspondence is clone M63 (Fig. 2 and Table 2), which obtained an extreme ST value (241 min) but whose EC50 was in the middle region of the plot (2.9 g·L⁻¹). Comparing clone sensitivity within each population, the ST values obtained ranged from 112.5 to 241 min in population M (brackish lagoon), 76.7 to 115.8 min for population LM (freshwater lake), and 62.5 to 116.7 min in population VF (freshwater ditch). These intervals showed that brackish clones were more halotolerant than the freshwater clones. The distribution of clones along the y-axis of Figure 2 shows that ST of freshwater genotypes was always below 120 min, while all brackish clones – except M7 (ST = 112.5 min) – displayed STs above this threshold. On average, clones from the brackish population (M) resisted 1 h more than the two freshwater populations (Fig. 3). This produced statistically significant differences among populations (Tab. 3), despite intra-populational (clonal) variability. Indeed, inter-populational variability was higher than the within population variance component (17% vs. 10%; Tab. 3).

HI values ranged from 29.9 to 64.0 for population M (brackish lagoon), 21.4 to 34.3 for population LM (freshwater lake), and 18.27 to 35.45 in population VF (freshwater ditch). Consistently with ST values, six out of eight genotypes from the brackish population scored the highest HI values (M20 and M36 were the exception) of all tested genotypes.

**Discussion**

Our study reports the differential tolerance to NaCl between one brackish and two freshwater populations of *S. vetulus*. All except one brackish genotype were able to
cope with high salinity stress (6 g L$^{-1}$) for a longer period than freshwater genotypes, in ST trials. Since most maternal and environmental influences were previously discarded (by culturing clones for several generations under standardized conditions), these differences are assumed to be genetically determined. Therefore, this population differentiation pattern is shaped by locally-acting selective forces, in this case salinity. The increased resilience of brackish clones to high salinity levels, expressed as increased survivorship, is in agreement to what was found by Weider and Hebert (1987) and Teschner (1995) for Daphnia. Local selective forces play an important role in the genetic structure and diversity of invertebrate populations, especially those with reduced dispersal capabilities (De Meester, 1996b; De Meester et al., 2002, 2006). Thus, even minor salinity increments in freshwaters (of a progressive, long-term nature) may cause subtle impacts in animal populations, such as the appearance of locally-selected races or “ecotypes” (sensu Weider and Hebert, 1987).

Unlike ST trials, no population differentiation pattern was found with standard acute assays. These contradictory results are portrayed by the lack of correspondence between acute (48 h) EC$_{50}$ and ST. In a work with D. longispina, Lopes et al. (2005) observed positive correlations between LT$_{50}$ (lethal time) and LC$_{50}$ (lethal concentration) only with copper, but not for zinc, cadmium, or hydrogen ions. Their original population was historically exposed to acid mine drainage contaminants, including copper and low pH. The lack of association between moderately lethal and extremely lethal levels of stressors, such as metals or pH in the study by Lopes et al. (2005) and NaCl (in our study), has an unknown biological meaning. High stressor levels trigger fast-response genes, such as hormones or protective enzymes, which modulate behavioural and physiological responses to increase resistance to environmental stress (see Hoffmann and Parsons, 1993). In the case of metals and hydrogen ions – and probably NaCl – animals die out of a series of related complications besides ionoregulatory failure (Wilson and Taylor, 1993). Hence, the regulation mechanisms to moderately or extremely lethal levels of toxicants should be the same; thus, other explanations must be at the basis of the contradictory results observed for ST trials and standard acute assays. One hypothesis is that the lack of association between both endpoints is related to their temporal resolution (see below).

Differences in ST between freshwater and brackish populations were observed in a limited time scale, varying from minutes to a few hours (between 35 and 178 min). Differences in sensitivity at such a time scale are not perceptible in standard acute exposures. In ST trials, a lethal concentration of NaCl is used and frequent measurements are taken, while in standard acute assays the organisms are checked at larger time intervals (24–48 h) after exposure to a range of increasing concentrations. There is considerable loss of information when daphniids are screened at such large time intervals (≥ 24 h). Thus, the finer temporal resolution of ST trials seems theoretically more advantageous and sensitive than standard acute exposures for exploring differentiation in quantitative genetic traits. Also, ST allows one to analyse the data using an ANOVA approach to partition-associated

### Table 3. Nested ANOVA summary tables for ST, including variance components (%Var). ANOVA model includes a fixed factor (Pop – population) and two (Clone and Rep – replicate) nested random factors.

<table>
<thead>
<tr>
<th>Model</th>
<th>d.f.</th>
<th>Adj. MS</th>
<th>F</th>
<th>P</th>
<th>%Var</th>
</tr>
</thead>
<tbody>
<tr>
<td>ST</td>
<td>Pop</td>
<td>2</td>
<td>130,938</td>
<td>9.88</td>
<td>0.001</td>
</tr>
<tr>
<td></td>
<td>Clone (Pop)</td>
<td>20</td>
<td>13</td>
<td>254</td>
<td>3.48</td>
</tr>
<tr>
<td></td>
<td>Rep (Clone)</td>
<td>46</td>
<td>3807</td>
<td>1.04</td>
<td>0.412</td>
</tr>
<tr>
<td></td>
<td>Error</td>
<td>345</td>
<td>3670</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Fig. 2. Relative halotolerance of S. vetulus genotypes from three populations (M, LM, and VF): acute EC$_{50}$ values vs. ST. Each circle, square, and triangle represents a genotype.

Fig. 3. Mean ST for each of the three populations studied (M, LM, and VF). Error bars represent standard error.
sources of variation (as done here), which is powerful and useful.

In this study, field clones were isolated from extant populations. Alternatively, many authors (e.g., Teschner, 1995; De Meester, 1996a; Boersma et al., 1999; Barry et al., 2005) use resting eggs (which result from sexual reproduction) as source of isolates. Both the active and dormant population have been subjected to different selective forces in each habitat throughout generations (De Meester, 1996b; De Meester et al., 2002), with emphasis on salinity in this case. The active population is less diverse, as it is constituted by individuals that have been subjected to local forces also during their ontogeny (De Meester et al., 2006), passing through various bottlenecks (hatching success, ability to feed and grow, asexual reproductive success). Therefore, the genotypes used in this study are the most adapted to the specific habitat of origin (freshwater or brackish). Thus, it is surprising that the range of acute sensitivity of the isolated freshwater and brackish genotypes is similar, with the only difference between them being the ability of brackish clones to survive longer under high salinity stress (6 g.L\(^{-1}\)). This is contrary to the findings of Lopes et al. (2004, 2005), who found remarkable differences between the acute sensitivities to copper (using standard acute assays) of historically-stressed and reference populations. We must conclude that the effect of salinity (original environment context) on the selection of genotypes was weaker than we had expected and than other authors have shown for other stressors.

When genotypes are isolated from the extant population, such as the case, the use of molecular fingerprinting is essential to assure clonal identity, since active populations are a mix of parthenogenetic and sexual clones (De Meester et al., 2006). In this study, we used a simple and cost-effective technique (MSP-PCR), which allowed satisfactory discrimination of genotypes (clones), analogously to RAPD (e.g., Weber and Declerck, 1997), microsatellite (e.g., Chopelet et al., 2008), or allozyme (e.g., Weider and Hebert, 1987) analyses. However, a conservative approach was needed, both in terms of band profile analysis and dissimilarity cut-off level, in order to avoid false positives (i.e., scoring identical genotypes as distinct). A reproducibility analysis of the technique showed that caution is needed because of variability among amplifications (probably due to misprints during PCR). Nonetheless, a few markers allowed the successful differentiation of the three studied populations (except for a small group of LM genotypes), even among populations with some connectivity degree (LM and VF). Although this was not the purpose of the study, we foresee a potential application of this technique in the decomposition of the genetic architecture of daphnid natural populations, provided a more comprehensive set of markers is used and that the technique’s reproducibility and amplification specificity are improved.

Acute EC\(_{50}\) for \(S. \text{v}etulus\) genotypes ranged between 2.3 and 3.8 g.L\(^{-1}\) of NaCl in this study. This is in agreement with the microcosm experiments of Sarma et al. (2006), whose \(S. \text{v}etulus\) experimental populations barely survived at 4 g.L\(^{-1}\), but were able to reproduce up to a concentration of 2 g.L\(^{-1}\). Halotolerance values for other cladocerans have also been reported using laboratory assays. Martínez-Jerónimo and Martínez-Jerónimo (2007) reported a 48 h LC\(_{50}\) of 5.48 g.L\(^{-1}\) for \(D. \text{magna}\) while demonstrating that acclimated organisms could survive and reproduce in NaCl concentrations up to 6 g.L\(^{-1}\). Gonçalves et al. (2007) reported an acute 48 h EC\(_{50}\) of 2.9 and 5.9 g.L\(^{-1}\) for \(D. \text{longispina}\) and \(D. \text{magna}\), respectively. These data suggest that \(S. \text{v}etulus\) halotolerance, regardless of their habitat of origin (brackish vs. freshwater), is comparable with that of \(D. \text{longispina}\), and both are more sensitive than \(D. \text{magna}\). Indeed, the latter species is documented to be halotolerant and is a common inhabitant of brackish lakes and rock pools (Arnérr and Koivisto, 1993; Teschner, 1996b; De Meester, 1996; Alonso, 1996). Ceriodaphnia lacustris, \(D. \text{thomsonii}\), \(D. \text{pulex}\) and \(D. \text{pulcarias}\) have also been found in habitats with moderate (5–13) salinities (Weider and Hebert, 1987; Barry et al., 2005; Northcote and Hall, 2010). However, most cladocerans are restricted to freshwater (Alonso, 1996) and are only able to cope with low salinities (usually <2; see Sarma et al., 2006) (for field data, see Jeppesen et al., 1994; Schallenberg et al., 2003; Marques et al., 2006; Santangelo et al., 2008). In fact, Schallenberg et al. (2003) predicts that severe alterations of zooplankton community structure and abundance may occur at salinities >1.2. These authors found *Simocephalus* sp. to tolerate salinities up to 1.2, while in our study \(S. \text{v}etulus\) occurred in Melides (population M) at a salinity of 2.3. However, we only found partial evidence that this population possesses genetically-determined tolerance mechanisms to endure saline stress, as portrayed by the lack of correspondence between standard acute assays (tolerance overlap) and ST (population differentiation).

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