Effects of Temperature and Salinity on the LMS (Lysosomal Membrane Stability) Biomarker in Clams *Donax trunculus* and *Chamelea gallina*

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**Abstract:** Population of clams *Donax trunculus* and *Chamelea gallina* have been declining significantly in recent decades, and environmental pollution and accelerated global warming have been proposed as contributing factors to this decline, in addition to overfishing. Lysosomal membrane stability (LMS) is a sensitive indicator of health status of the organisms. In this study, we investigate the LMS in these species after exposure for 21 days to nine combined conditions of water temperature (12, 20, and 27.5 °C) and salinity ranges (27–28, 32–33, and 37–38). LMS was assessed in living hemocytes by using the neutral red retention assay. Mortality and the condition index of the organisms were evaluated as supporting parameters. The results indicated interspecies differences in the LMS under similar environmental conditions. Overall, LMS was found to be more sensitive to temperature than to salinity changes. Although both species can tolerate changes in either salinity or temperature seawater conditions, the tolerance range is narrower for *D. trunculus*, showing a significant cytotoxicity (NRRT < 50 min) at temperatures above 27.5 °C and salinities above 32, and 100% mortality at 27.5 °C and a low salinity range (27–28). This study is the first to assess the combined effect of temperature and salinity on the LMS in *C. gallina* and *D. trunculus*, and provide necessary information before using LMS as contaminant-related biomarker in field studies with these species.

**Keywords:** bivalves; environmental stress; cytotoxicity; biomarker; lysosomal membrane stability; neutral red assay; condition index

1. Introduction

In the western Mediterranean Sea, the wedge clam (*Donax trunculus*) and the striped venus clam (*Chamelea gallina*), among other species, hold significant socio-economic value as fishing resources and consequently face high fishing pressure [1,2]. These filter-feeding bivalve mollusks inhabit fine sandy bottoms [3,4]. *D. trunculus* exclusively resides in the biocenosis of superficial fine sands [5] with strong dependence on the nature and composition of the sediment [6]. On the other hand, the striped venus clam is preferentially distributed in the coastal biocenosis of well-selected fine sand [7], typically found at depths of 3 to 12 m [8], while wedge clams are predominantly distributed between 0 and 2 m deep [9].

Over the recent few decades, the fisheries of these clam species have witnessed a significant decline in the Spanish Mediterranean Region [1,2,10]. This decline has led to the gradual closure of fishing grounds by local authorities or the abandonment of fishing activities in some regions [1]. Management measures, such as legal minimum size requirements...
or spatial-temporal closures to avoid the fishing during the spawning season, have been implemented to address this situation [2,11]. Despite these efforts, the populations of wedge clams and striped venus clams have not recovered [1], resulting in severe consequences, including the loss of employment for hundreds of small-scale fishermen [1].

Marine ecosystems, particularly in the Mediterranean Sea, are experiencing the impacts of global climate change [12], chemical pollution [13], and biological pollution [14], resulting in significant effects on marine biodiversity [15,16]. While the ocean absorbs both CO$_2$ and heat, slowing atmospheric warming, it also undergoes changes such as increased water temperature, reduced pH, decreased dissolved oxygen, salinity fluctuations, heatwaves, rising sea levels, and alterations in ocean currents [17]. Each of these stressors has its own effect, acting both independently [18] and synergistically [19,20]. Warming, acidification, deoxygenation, and salinity changes occur more rapidly in shallow water habitats, influenced by continental factors. Other anthropogenic pressures such as high urban pressure, mass tourism, overfishing, increased maritime traffic, sewage disposal, and extensive use of fertilizers are added to these coastal marine zones [21]. Consequently, the Mediterranean Sea is being pushed into a new and, at least, uncertain state by all these stress sources [14].

Rivetti et al. [22] linked changes in the temperature of surface waters of the Mediterranean Sea to mass mortality events of invertebrates, including mollusks, concluding that the frequency of mass mortality of benthic invertebrates is increasing and correlates with positive thermal anomalies. In the Mediterranean, marine heat waves are not only more frequent but also more intense and of longer duration [23]. Therefore, the two species under study may struggle to maintain good health at high temperatures. Moschino and Marin [24] demonstrated that *C. gallina* at temperatures of 28 °C increased their respiration rate, potentially leading to metabolic imbalances, while Widdows and Staff [25] suggested that wild clams may experience high stress levels at such temperatures potentially causing death [26]. In the case of *D. trunculus*, Ansell et al. [27] observed a high thermal tolerance for this species, reflecting its distribution in shallower waters; however, their determinations were carried out working with short exposure periods of up to 96 h.

Marine bivalves are osmoconformers capable of maintaining metabolic function despite changes in osmolarity and cytosol composition during changes in salinity [28]. Closing and sealing of the valves are vital responses to stressful salinity [29]. Despite these adaptations, salinity variations can also cause mortality of benthic bivalves [30]. Although *D. trunculus* is an euryhaline species, prolonged exposure to low salinities slows its movements, which can affect its survival in the environment [31]. Matozzo et al. [32] suggest that salinity variations above and below 34 allow for the observation of quantifiable alterations in the immune responses of *C. gallina*.

When investigating the biological effects of environmental factors, the alterations in certain physiological processes or subcellular responses can be used as biomarkers of organism health condition. In this sense, lysosomal membrane stability (LMS) represents the most sensitive biomarker of stress at cellular level. LMS in bivalve hemocytes is a core biomarker of general stress of chemical pollution that has been widely recommended by different organizations, such as the Barcelona Convention, International Council for Exploration of the Sea (ICES), and Oslo and Paris (OSPAR) Conventions, and implemented in mussel monitoring programs of chemical pollution [33–35]. Given the fact that a variety of factors in addition to overfishing, such as environmental pollution, have been proposed to explain the population significant declines of *Donax trunculus* and *Chamelea gallina* mollusks in the Mediterranean Region, interest and opportunities arise to use LMS as a general stress biomarker of chemical environmental pollution in these species. However, the utility of this biomarker in other bivalve species than *Mytilus* sp. has to be supported by knowledge on how they respond to extrinsic abiotic and intrinsic biotic factors [36]. This is due to the fact that LMS is a robust diagnostic indicator of individual health status in bivalves, but it is also affected by non-contaminant factors, such as temperature, prolonged hypoxia, salinity, food deprivation, or the final stage of gametogenesis [37]. Therefore, the
use of LMS to monitor the impacts of environmental chemical pollution on these clam species should involve knowing the response of the LMS biomarker to the temperature and salinity variability to which these species might be subjected in their habitats, as they may act as confounding factors in the assessment of the biomarker response. In this sense, the main objective of this study was to investigate the cellular wellbeing of *Donax trunculus* and *Chamelea gallina* living hemocytes by assessing the lysosomal membrane stability in order to evaluate their health status after short-term exposure to different environmental conditions of temperature and salinity.

## 2. Materials and Methods

### 2.1. Collection of Clams and Maintenance

Specimens of *D. trunculus* and *C. gallina* were collected in the southern part of the Gulf of Valencia (SE Spain), between the beaches of l’Ahuir and Daimús. The organisms were collected with hand dredges at a depth of 0.5 to 2 m and 4 to 7 m, respectively. To avoid keeping individuals for long periods in laboratory conditions, it was decided to sample mollusks when the difference between the natural temperature in the field and the laboratory test temperature was minimal. Consequently, sampling of individuals was carried out during 2023 on three occasions: March (15.4 °C and 36.9 of salinity), May (19.5 °C and 36.8), and June (23.5 °C and 37.1). The size range of *D. trunculus* clams was 16.1 to 28.5 mm, while it ranged from 12.8 to 24.5 mm for *C. gallina* in March, and from 20.8 to 31.4 mm in May and June.

The collected clams (around 450 specimens of each species) were immediately transferred to the laboratory and maintained in aquariums filled with seawater collected from the sampling points. The aquariums were placed in environmental chambers with a controlled temperature and photoperiod (12 h light/12 h dark), constant aeration, and recirculating filtered seawater. Clams were fed daily with microalgae (*Isochrysis galbana*). The organisms were acclimated for 5–7 days to the experimental conditions by progressively decreasing or increasing the temperature or salinity by 1–2 (°C) per day. Organisms sampled in March, May, and June were acclimated to 12 °C (low), 20 °C (medium), and 27.5 °C (high) temperatures, respectively (Figure 1). Experimental temperatures and salinities were chosen considering the characteristic conditions of the Mediterranean Sea, where values close to those tested in this study are recorded. During winter, water temperatures range between 13 and 16 °C, while they are around 22 and 27 °C in the summer [38]. Salinity generally varies between 36 and 38 [38], decreasing in places near river mouths [4].

![Experimental design overview](image-url)

**Figure 1.** Experimental design overview. A visual representation illustrating the arrangement and variables of the experimental design employed in this study.
2.2. Exposure to Experimental Temperatures and Salinities

Nine combined conditions of temperature and salinity were studied in both species. Animals were held for 21 days at three different temperatures (low (12 °C), medium (20 °C), and high (27.5 °C)) and at three salinity ranges (27–28, 32–33, and 37–38) for each temperature condition (Figure 1). Experimental units for each environmental condition consisted of 40 L aquaria that contained 50 individuals. Seawater parameters (temperature, salinity, and dissolved oxygen) and mortality were monitored daily in every aquarium for both species. Temperature, salinity, and pH were measured using a calibrated WTW Model 340i multiprobe (WTW GmbH, Weilheim, Germany), while dissolved oxygen was measured using a proODO YSI (YSI Inc., Yellow Springs, OH, USA) optical probe. Salinity was corrected using distilled water and seawater. Mortality was determined by the number of clams with open shells that did not react to the mechanical stimulus. Organisms were exposed for 21 days as this period reflects the time frame of sustained exposure to different salinity/temperature regimes in the southern sector of the Gulf of Valencia.

2.3. Lysosomal Membrane Stability (LMS)

After 21 days of exposure, hemolymph samples were obtained from 16 clams from each treatment and hemocyte monolayers were prepared as previously described in Martínez-Gómez et al. [39]. From each clam, 0.1 mL of hemolymph was extracted from the adductor muscle using a 1 mL hypodermic syringe with a 25-gauge needle and pre-loaded with 0.1 mL of filtered aquarium water (0.45 µm) (corresponding with the seawater of the treatment). LMS was measured by using the in vivo cytochemical method named neutral red retention assay (NRRA), following the procedure described by Lowe et al. [40], with modifications as described in Martínez-Gómez et al. [39]. After 15 min of incubation with neutral red (NR) (CAS: 553-24-2; Labkem, Barcelona, Spain), hemocyte samples were observed under a microscope (×400 and ×200 magnification) at 15, 30, 60, 90, 120, and 180 min and examined for both structural abnormalities (lysosomal swelling, enlargement, and rounded-up fragmenting cells) and lysosomal NR leakage. Results of LMS were expressed as the neutral red retention time (NRRT) (minutes), corresponding to the last time period recorded when there was no evidence of dye loss or lysosomal abnormalities in more than 50% of the cells. To enhance the sensitivity of the LMS measurements, results were also expressed as %LMS, by applying the scoring procedure described in Martínez-Gómez et al. [39], which considers not only the neutral red retention time but also lysosomal alterations. In non-stressed cells, lysosomes will accumulate and retain the NR dye for a prolonged period and will not present lysosomal alterations, whereas in stressed cells, NR will leak into the cytosol and hemocytes will show lysosomal abnormalities. The size of the organisms used to analyze lysosomal membrane stability ranged from 13.9 to 23.9 mm (in March) and from 20.8 to 31.44 mm (in May and June) for *C. gallina* and from 16.1 to 28.5 mm for *D. trunculus*.

2.4. Condition Index (CI)

The CI provides information on the nutritional and physiological status of bivalves [24,41] and provides complementary information to interpret LMS results [42].

CI was determined before and after 21 days of exposure of the organisms (n ~10 individuals) to the nine treatments (three temperatures ranges combined with three salinity ranges). Because CI varies according to organism age and physiological status [43], CI was standardized to certain size ranges. For *D. trunculus*, organisms were selected with a size range of 19–23 mm. In the case of *C. gallina*, because the organisms sampled in March were exceptionally small ones, the size range used was 13–16 mm, whereas for the May and June samplings, it was 17–20 mm.

The CI was calculated using the equation applied by Boscolo et al. [44], as follows:

\[
\%CI = \left[\frac{\text{dry soft tissue weight (g)}}{\text{shell length (cm)}}\right] \cdot 100, \tag{1}
\]
The weight of the dried soft tissue was obtained by drying the soft tissue individually at 100 °C until a constant weight was obtained. The shell length was measured with a digital vernier caliper.

2.5. Data Analysis

Values in the text are expressed as mean ± standard error (SE) if not otherwise specified. The 95% confident limits of the mean LMS values (expressed as minutes of NRRT) were assessed against their corresponding background assessment criteria (BAC) and environmental assessment criteria (EAC) developed for the technique [44]. Statistical analyses were carried out using the software packages IBM® SPSS Statistics v.17 and Statgraphics Centurion XVII. All data were preliminarily checked for normality (Shapiro-Wilk test) and homoscedasticity (Levene’s test); when necessary, they were transformed using Arcsine√(p/100) or Box-Cox transformation (lambda was found using the most likely estimate (MLE) approach) to meet the required assumptions [45]. Differences in NRRT, %LMS, and %CI among treatments were identified using multifactorial ANOVA (2-way ANOVA). The factors were temperature with 3 levels (10 °C, 20 °C and 27.5 °C) and salinity with 3 levels (27–28, 32–33, and 37–38). To obtain specific LMS information for all treatments and for each temperature and salinity condition, a pairwise comparison (Tukey’s HSD post hoc, p < 0.05) of 9 treatments was carried out: (i) the 3 ranges of salinity levels for each temperature and (ii) the 3 temperature levels for each salinity. Additionally, a pairwise comparison was performed to determine the differences between the CIs of the reference organisms in the three samplings (March, May, and June—on day 0) with the CI values of each treatment (after 21 days of exposure).

Finally, Pearson’s correlation coefficient was used to study the correlations between temperature, salinity, LMS, and %CI.

3. Results

Information on the experimental conditions for each treatment is presented in Table 1. The seawater temperature and salinity values of the treatments during the experiment differed slightly from the nominal values. Dissolved oxygen remained within the appropriate range (~100% oxygen saturation) at all experimental conditions. The pH remained fairly constant between 8.0 and 8.2.

Table 1. Seawater parameters (mean ± SE) during the 21 days experiment conducted with Donax trunculus and Chamelea gallina. N: treatment number; T: temperature; S: salinity; DO: dissolved oxygen (mg/L and % oxygen saturation).

<table>
<thead>
<tr>
<th>Test Treatment</th>
<th>Donax trunculus</th>
<th>Chamelea gallina</th>
</tr>
</thead>
<tbody>
<tr>
<td>N</td>
<td>T</td>
<td>S</td>
</tr>
<tr>
<td>1</td>
<td>27-28</td>
<td>12.30 ± 0.05</td>
</tr>
<tr>
<td>2</td>
<td>Low</td>
<td>32-33</td>
</tr>
<tr>
<td>3</td>
<td>37-38</td>
<td>12.17 ± 0.03</td>
</tr>
<tr>
<td>4</td>
<td>Medium</td>
<td>27-28</td>
</tr>
<tr>
<td>5</td>
<td>32-33</td>
<td>19.95 ± 0.04</td>
</tr>
<tr>
<td>6</td>
<td>37-38</td>
<td>20.34 ± 0.05</td>
</tr>
<tr>
<td>7</td>
<td>High</td>
<td>27-28</td>
</tr>
<tr>
<td>8</td>
<td>32-33</td>
<td>26.94 ± 0.09</td>
</tr>
<tr>
<td>9</td>
<td>37-38</td>
<td>26.98 ± 0.11</td>
</tr>
</tbody>
</table>

3.1. Chamelea Gallina

No mortality occurred in any of the treatments after 21 days of exposure, so survival was 100%.

The results of NRRT and %LMS values of C. gallina are presented in Figure 2 and Table S1. There was a significant difference in LMS (expressed as NRRT) as a result of temperature and the interaction of both factors (temperature and salinity) but no effect of salinity (2-way ANOVA; p-value < 0.05) (Table 2). LMS in C. gallina hemocytes (expressed
as mean value of NRRT) was higher in organisms from the 20 °C/27–28, 20 °C/37–38, and 12 °C/32–33 salinity treatments (Figure 2a), where hemocytes showed a longer retention time of neutral red within the lysosomal compartment (Figure 3A). Pairwise comparisons revealed three homogeneous subsets among the nine combined treatments (Tukey-b test; p-value < 0.05) (Table S2). At a low temperature (12 °C), NRRT was significantly higher in animals exposed to salinity of 32–33, compared to those maintained at 27–28 and 37–38 (p-value = 0.003) (Figure 2a and Table S3). However, an opposite pattern was found at a medium temperature (20 °C), with NRRTs being higher at salinities 27–28 (p-value = 0.002).

When LMS was estimated by adding information of lysosomal alterations to the NRRTs (%LMS), no differences were observed due to temperature and salinity, but there was a significant interaction between these factors (2-way ANOVA; p-value ≤ 0.001) (Table 2). No differences were observed in %LMS between the three salinities tested within the three temperatures studied, except for the treatment comprising 12 °C/32–33 of salinity, which was significantly higher (Figure 2b and Tables S2 and S3) (p-value = 0.014). Nevertheless, some organisms showed lysosomal alterations in the cells (Figure 3B).

**Figure 2.** (a) Neutral red retention time (NRRT) and (b) Lysosomal membrane stability percentage (%LMS) in *C. gallina* for 21 days exposure to combined conditions of temperature (low, medium, and high) and salinity ranges (27–28, 32–33, and 37–38). Background assessment criteria (BAC). Environmental assessment criteria (EAC). Data are reported as mean ± SE, n = 16. The lowercase letters above the bars indicate inclusion within homogeneous subsets (2-way ANOVA; Tukey’s post hoc test; p-value < 0.05).
Table 2. Results of the two-way ANOVA, showing the effects of temperature (T), salinity (S), and their interaction in *C. gallina* and *D. trunculus*. Both factors were considered fixed and three dependent variables were examined: lysosomal membrane stability (expressed as NRRT and %LMS) and condition index (%CI).

<table>
<thead>
<tr>
<th>Factor</th>
<th>NRRT</th>
<th>%LMS</th>
<th>%CI</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>df</td>
<td>F</td>
<td>p-Value</td>
</tr>
<tr>
<td><em>Chamelea gallina</em></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>T</td>
<td>2</td>
<td>4.28</td>
<td>0.016</td>
</tr>
<tr>
<td>S</td>
<td>2</td>
<td>0.65</td>
<td>ns</td>
</tr>
<tr>
<td>T × S</td>
<td>4</td>
<td>6.75</td>
<td>0.000</td>
</tr>
<tr>
<td><em>Donax trunculus</em></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>T</td>
<td>2</td>
<td>35.92</td>
<td>0.000</td>
</tr>
<tr>
<td>S</td>
<td>2</td>
<td>4.57</td>
<td>0.012</td>
</tr>
<tr>
<td>T × S</td>
<td>4</td>
<td>4.49</td>
<td>0.005</td>
</tr>
</tbody>
</table>

NRRT: neutral red retention time. %LMS: percentage of lysosomal membrane stability. Statistically significant differences (p-value < 0.05). Ns: not significant (p-value > 0.05).

Figure 3. Hemocytes (small granulocytes) from *Chamelea gallina* showing the retention of neutral red within the lysosomal compartment (A), and stressed cells showing loss of neutral red from pathologically enlarged lysosomes into the cytosol, enlargement, and leakage but colorless lysosomes (B).

There was a significant difference in the condition index for *C. gallina* due to temperature but there was no effect of salinity and no interaction between these factors (2-way ANOVA; p-value < 0.05) (Table 2). The CI of the organisms did not change significantly after 21 days of treatment, except in organisms maintained at the highest temperature (p-value = 0.007), where CI after 21 days was lower than at the beginning of the experiment (Table 3 and Table S4). Although the CIs obtained in the low-temperature treatments were similar to those obtained at a high temperature, they were not comparable due to the difference in size between the organisms studied (13–16 mm vs. 17–20 mm).

Table 3. Size and condition index (CI) of *C. gallina* and *D. trunculus* at the beginning and after 21 days exposure to nine combined treatments of temperature and salinity. Data are reported as mean ± SE.

<table>
<thead>
<tr>
<th>Reference (Day 0)</th>
<th>Salinity (Day 21)</th>
</tr>
</thead>
<tbody>
<tr>
<td>27–28</td>
<td>32–33</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>T (°C)</th>
<th>S (%)</th>
<th>CI (%)</th>
<th>T (°C)</th>
<th>S (%)</th>
<th>CI (%)</th>
<th>T (°C)</th>
<th>S (%)</th>
<th>CI (%)</th>
<th>T (°C)</th>
<th>S (%)</th>
<th>CI (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Low</td>
<td></td>
<td></td>
<td>Medium</td>
<td></td>
<td></td>
<td>High</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>15.4/36.9</td>
<td>–</td>
<td>–</td>
<td>17.93</td>
<td>2.78ab</td>
<td>± 0.23</td>
<td>18.91</td>
<td>2.52a</td>
<td>± 0.12</td>
<td>19.03</td>
<td>2.89ab</td>
<td>± 0.35</td>
</tr>
<tr>
<td>Medium</td>
<td>19.5/36.8</td>
<td>2.61a</td>
<td>18.33</td>
<td>2.02b</td>
<td>± 0.18</td>
<td>18.01</td>
<td>2.07b</td>
<td>19.83</td>
<td>2.18b</td>
<td>0.007</td>
<td></td>
</tr>
<tr>
<td>High</td>
<td>23.5/37.1</td>
<td>± 0.08</td>
<td>± 0.26</td>
<td>22.65</td>
<td>± 0.09</td>
<td>± 0.22</td>
<td>± 0.09</td>
<td>± 0.15</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Lowercase letter superscripts indicate homogeneous subsets for each species and temperature treatment (Tukey’s post hoc). Ns: not significant (p-value > 0.05). “—” represents no data, and “X” indicates that there were no organisms due to mortality.
After 21 days of exposure to nine combined treatments, the CI in *C. gallina* (*n* = 9) did not correlate with the temperature or salinity conditions tested, nor with the LMS values measured in this species (Figure 4 and Table S5).

### 3.2. *Donax Trunculus*

In contrast from what was observed with *C. gallina*, there was acute mortality of individuals of *D. trunculus* during the experiments (Figure 5). Overall, the most notable decrease in percentage of survival was associated with the treatments conducted at the highest temperature, with a 100% mortality of individuals after 14 days at 27.5 °C/27–28 salinity (Figure 5c). The survival rate of the organism in the other treatments (low and medium temperatures and at all three salinity levels) ranged between 70 and 90% (Figure 5a,b).

There was a significant difference in LMS (expressed as NRRT and %LMS) as a result of both temperature and salinity, and there was a significant interaction between these factors (2-way ANOVA; *p*-value < 0.05) (Table 2). LMS in *D. trunculus* hemocytes (expressed as mean value of %LMS) was significantly lower in survival clams from the treatments conducted at 27.5 °C than in the other treatments (Figure 6). The highest mean values of LMS were observed in clams from the 20 °C/32–33 of salinity (NRRT = 165.9 ± 10.7 min; %LMS = 82.7 ± 5.1) and 20 °C/37–38 of salinity (NRRT = 136.9 ± 13.1 min; %LMS = 72.8 ± 6.7) treatments (Table S1).

Pairwise comparisons revealed the existence of LMS response subsets (6 subsets for NRRT and 4 subsets for %LMS) (Tukey’s post hoc test; *p*-value < 0.05) (Figure 6a,b and Tables S6 and S7). The treatments carried out at the same temperature but at different salinities had similar LMS results in %LMS (Figure 6b and Tables S6 and S7) (Tukey’s post hoc test; *p*-values > 0.05).

There was a significant difference in condition index due to temperature, but there was no effect of salinity or the interaction of both factors (Table 2). Unlike *C. gallina*, the CI of *D. trunculus* decreased with increasing temperature (Table 3). Organisms exposed to a low temperature exhibited the highest CI values, showing no significant differences due to salinity. At a medium temperature (20 °C), a lower CI was obtained compared to those exposed at a low temperature (12 °C). The reference CI (of 19.5 °C and 36.8 salinity) differed from the CI obtained at 27–28 and 32–33 (*p*-value = 0.007, Table 3). Finally, organisms exposed to a high temperature (27.5 °C) had the lowest CI values. At this temperature, significant differences were observed between the reference CI (23.5 °C and 37.1 salinity) and those exposed to salinities of 32–33 and 37–38 (*p*-value = 0.000, Table 3). It should be noted that, during exposure to a high temperature, spawning occurred in *D. trunculus*, coinciding with the low LMS and CI recorded, pointing to a temperature-induced spawning event in this species.
Correlation analysis also revealed positive correlations between the two “endpoints” (Table S8). In *D. trunculus* (*n* = 8), a negative correlation was observed between CI and treatment temperature (Pearson’s correlation Coeff. = −0.976; *p*-value = 0.000) and a moderate positive correlation between CI and %LMS (Pearson’s correlation Coeff. = 0.737; *p*-value = 0.037) (Figure 4).

![Graph](image)

**Figure 5.** Percentage of *Donax trunculus’* survival as a function of salinity at each treatment temperature during 21 days of exposure at a (a) low water temperature (12 °C), (b) medium water temperature (20 °C), and (c) high water temperature (27.5 °C) at salinities of 27–28, 32–33, and 37–38, respectively.
There was a significant difference in condition index due to temperature, but there was no effect of salinity or the interaction of both factors (Table 2). Unlike C. gallina, the CI of D. trunculus decreased with increasing temperature (Table 3). Organisms exposed to a low temperature exhibited the highest CI values, showing no significant differences due to salinity. At a medium temperature (20 °C), a lower CI was obtained compared to those exposed at a low temperature (12 °C). The reference CI (of 19.5 °C and 36.8 salinity) differed from the CI obtained at 27–28 and 32–33 (p-value = 0.007, Table 3). Finally, organisms exposed to a high temperature (27.5 °C) had the lowest CI values. At this temperature, significant differences were observed between the reference CI (23.5 °C and 37.1 salinity) and those exposed to salinities of 32–33 and 37–38 (p-value = 0.000, Table 3). It should be noted that, during exposure to a high temperature, spawning occurred in D. trunculus, coinciding with the low LMS and CI recorded, pointing to a temperature-induced spawning event in this species.

Correlation analysis also revealed positive correlations between the two “endpoints” (Table S8). In D. trunculus (n = 8), a negative correlation was observed between CI and treatment temperature (Pearson’s correlation Coeff. = −0.976; p-value = 0.000) and a
In this study, we found that certain combined conditions of seawater temperature and salinity maintained for at least 21 days are capable, by themselves, of causing significant cellular stress by affecting lysosomal membrane permeability in the clam species *D. trunculus* and *C. gallina*. We found a differential response to temperature, dependent on acclimation salinity, which has also been observed in other bivalves [53]. In *C. gallina*, at salinities of 32–33, the neutral red was retained within the lysosomal compartment for the shortest duration at 20 °C, increasing as the acclimation salinity was raised or lowered (Figure 2). However, an opposite pattern was observed in the treatments conducted at 12 °C, with longest retention times at 32–33 salinities. At the highest temperature (27.5 °C), no significant changes in lysosomal membrane stability were found at different salinities. Furthermore, our findings show that this differential response varied between the clam species under study. The lysosomal permeability to intermediate salinities (32–33) at 12 °C and 20 °C in *C. gallina* has opposite patterns in comparison to *D. trunculus*. Different patterns of variation in total hemocytes counts has been described between *C. gallina* and *Mytilus galloprovincialis* submitted to the same environmental conditions [56], which partially supports the differential interspecies responses found on LMS in our study. In both species, the LMS at 27.5 °C showed little response to changes in salinity. Consequently, our findings show that the adaptive capacity of the cellular processes in these species was reduced at the highest temperature tested. For *D. trunculus*, at the highest tested temperature (27.5 °C), the NR remained within the lysosomes for only a short time (Figure 6), signaling that the lipid membranes within the cell were less stable.

It is known that the interaction between two or more environmental variables may cause changes in other variables [57], such as a decrease in dissolved oxygen (DO) in the water, which may cause stress in organisms [58,59]. In our study, the DO concentration was kept under control (>7 mg/L and 100% oxygen saturation), and the observed effects cannot be attributed to significant changes in this variable (Table 1).

The neutral red retention assay has been and is widely used to determine lysosomal membrane stability in invertebrate hemocytes, with special reference to bivalves [60,61]. Threshold values used as assessment criteria for lysosomal membrane stability using the NRRA have been determined from *Mytilus* sp. on the bases of numerous studies performed around the UK and other European waters, including Mediterranean waters [39,62]. There is a limited number of studies investigating LMS in *C. gallina* hemocytes [52,54,63]. To our knowledge, there are no LMS data in the literature obtained by using the NRRA in the hemocytes of *D. trunculus*. In the absence of specific evaluation criteria for these species, absolute values for LMS measurements in *Mytilus* sp. could be assumed to be comparable. In this sense, NRRT shorter than the EAC level suggested the organism were severely stressed and probably exhibiting pathology. An NRRT shorter than the BAC level but longer than the EAC level was considered to represent stressed but compensating organism. Concerning *C. gallina*, neutral red retention times within the lysosomal compartment of the small granulocytes were in general longer (mean time higher than 50 min for all treatments) than the retention times reported on an earlier field study performed in Adriatic waters with the same species (NRRT ≤ 30 min in spring (20 °C salinity of 35) and autumn (16 °C salinity of 36)) [52]. According to the assessment criteria established for this biomarker, our results point out that the treatments performed with *C. gallina* caused less cellular stress compared to *D. trunculus*, since the organisms showed retention times below the environmental assessment criteria established for this biomarker (EAC < 50 min) in none of the treatments [62]. In general, for both species, the environmental conditions that led to greater LMS were those at a medium temperature (20 °C) at the three studied salinity ranges. In other bivalves, temperatures close to 20 °C have been found to be optimal for maintaining hemocyte lysosomal stability, with membrane stability being decreased both when increasing and decreasing the temperature [61,64]. In addition, in the case of *C. gallina*, organisms exposed to the lowest temperature (12 °C) and salinity 32–33 also presented high stability. For *D. trunculus* at 20 °C and salinity 32–33 and 37–38, the NRRTs were higher.
than the background assessment criteria value, while at the temperature 27.5 °C, at these same salinities, the NRRTs obtained were lower than the EAC value.

Moore et al. [65] observed that mussels exposed to aquatic hyperthermia conditions (28 °C) for 48 h were able to recover LMS values. In our current investigation, prolonged exposure to elevated temperatures (27.5 °C) combined with a salinity of 27–28 resulted in 100% mortality in D. trunculus after 14 days. These results suggest that 27.5 °C could be the threshold temperature level for this species at low salinities. Moreover, in the surviving treatments at that temperature (salinity 32–33 and 37–38), the lowest LMS and the lowest CI values of all treatments were observed. The cellular responses analyzed revealed extremely severe stress in the organisms of those treatments 27.5 °C/32–33 and 27.5 °C/37–38 (NRRT of 28.1 ± 12.7 min and 42.2 ± 9.9 min, respectively) and the lowest mean %LMS values (20.8 ± 7.0 and 36.9 ± 5.4, respectively). Overall, the results of this study indicated that LMS responses were more affected by temperature than salinity, as NRRTs greater than 50 min at 27.5 °C were never detected, regardless of salinity. In the case of C. gallina, although Moschino and Marin [24] reported alterations of physiological measurements, such as growth and condition index, as well as induction and alterations of sub-cellular biomarkers at temperatures higher than 25 °C, our findings indicate that the values at 27.5 °C were similar to others obtained at 12 and 20 °C.

In addition to variations in environmental parameters, the lysosomes in the vascular system respond to physiological changes associated with gonadal maturation during the spawning period [66,67]. Spawning is also an important physiological stressor in bivalves, which significantly increases lysosomal membrane permeability [61,68,69]. There is a wide discussion among different authors on the impact of water temperature in relation to the spawning period of bivalves [67,70]. Concerning mussels, some researchers argue that lower NRRT values are obtained in the pre-spawning period due to the maintenance of their metabolic functions [36,71], while Harding et al. [72] suggest that these values are reached just after spawning. On the other hand, some indicate that NRRT is not affected by hormonal regulation during reproductive activity in mussels [34]. In our study, for D. trunculus, organisms collected in June (when this species would be in the spawning stage in our study area [73]) and exposed to high temperatures spawned, adding an additional stress factor to temperature and salinity. Thus, the lowest LMS and low CI values obtained in D. trunculus at 27.5 °C could be at least partially due to the spawning observed in the aquaria.

Hemocytes represent the primary line of internal defense for bivalves (endocytosis, encapsulation, inflammation, wound repair, enzyme secretion, elimination by diapedesis), and changes in ambient temperature and salinity may enhance or decrease the hemocytes activity [74]. The LMS of hemocytes in D. trunculus was significantly affected at 27.5 °C. In any case, temperature was a dominant factor in the physiological and immunological responses of D. trunculus, showing lower survival, lower stability of lysosomal membranes, and lower CI value when reaching a high temperature (27.5 °C), thus indicating a higher level of stress. This fact is relevant considering the current situation of the Mediterranean Sea, where the effects of climate change are appreciated: marine heat waves are more frequent and intense with a longer duration so species such as Donax and Chamelea are subjected to extreme temperatures and for longer periods in the environment. After conducting this study, certain limitations were identified that could be addressed in future research. Kruft Welton et al. [20] states that infaunal species, such as ours, may not be negatively affected by climate change stressors due to their ability to burrow as a strategy to avoid high temperatures. However, Monari et al. [63] found that the C. gallina kept at 20 and 25 °C burrowed completely, whereas at 30 °C, the clams progressively emerged from the sediment and then remained on the surface. Nonetheless, including a sand bed in the aquaria during the experimental period would provide results ecologically more relevant concerning the effects of water temperature on the LMS in these species with burying behavior.
5. Conclusions

This study contributes to clarifying the cellular stress caused by the action of combined temperature and salinity conditions in the marine clams *C. gallina* and *D. trunculus*. This investigation demonstrates interspecies differences in the background response of the lysosomal membrane stability biomarker under similar environmental conditions. Given the importance of considering environmental variables when evaluating the toxicity of pollutants in aquatic organisms, our results have a relevant interest in the potential use of LMS as a contaminant-related biomarker in *C. gallina* and *D. trunculus* in field studies.

Although both species can tolerate changes in either seawater salinity or temperature conditions, the tolerance range is narrower for *D. trunculus*, showing a significant cytotoxicity (NRRT < 50 min) at temperatures above 27.5 °C and salinities above 32, and causing 100% mortality at 27.5 °C and low salinities (27–28). This research contributes to consolidating the findings already obtained from other bivalve species that climate changes can affect hemocyte functionality, which is directly related to bivalve immune defense capabilities.

**Supplementary Materials:** The following supporting information can be downloaded at: https://www.mdpi.com/article/10.3390/app14072712/s1, Table S1: Descriptive variables of the organisms used in LMS of each species: size, neutral red retention time (NRRT), and percentage lysosomal membrane stability (%LMS). T: temperature; S: salinity; N: individuals. Data are expressed as mean ± SE., Table S2: Means for groups in homogeneous subsets of Tukey’s HSD (p < 0.05) test of neutral red retention time (NRRT) and percentage of lysosomal membrane stability (%LMS) among the 9 treatments for *C. gallina* (n = 16), Table S3: Pairwise comparisons of LMS (expressed as NRRT and %LMS) at three salinity (S) levels (27–28, 32–33, and 37–38) for each temperature (T) and at three temperature levels (low, medium, and high) for each salinity, for both species, Table S4: Pairwise comparisons of the condition index (CI) of the four salinities (T) levels (reference, 27–28, 32–33, and 37–38) for each temperature (S) (reference: low, medium, and high), Table S5: Correlation analysis between the different variables for *C. gallina*: salinity (S), temperature (T), condition index (CI), neutral red retention time (NRRT), and percentage of lysosomal membrane stability (%LMS) (n = 9), Table S6: Means for groups in homogeneous subsets of Tukey’s HSD (p < 0.05) test of neutral red retention time (NRRT) among the 9 treatments for *D. trunculus* (n = 16), Table S7: Means for groups in homogeneous subsets of Tukey’s HSD (p < 0.05) test of percentage of lysosomal membrane stability (%LMS) among the 9 treatments for *D. trunculus* (n = 16), Table S8: Correlation analysis between the different variables for *D. trunculus*: salinity, temperature, condition index (CI), neutral red retention time (NRRT), and percentage of lysosomal membrane stability (%LMS) (n = 8).


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