1. Introduction

-Anemonia viridis as food item and restrictions in Andalusia and Spain.

-Reproduction modes in A. viridis, birradial symmetry and anatomy.

- Integrated multitrophic aquaculture. Sustainability. Do not focus on this unless the journal calls for it.

- Oxidative stress as a marker of wellbeing in aquaculture.

-Justificar y/o relacionar los cambios metabólicos asociados 1) al estrés por manipulación (inducción asexual) y 2) a la maduración sexual y como ambos pueden afectar al estado estado oxidativo (para justificar la metodología del trabajo)

2. Materials and methods

2.1. Experimental design and sampling

Wild anemones were obtained from the natural environment off the coast of Granada (Andalusia, Spain) in (month-year) with the pertinent permits from local authorities. Animals were transferred to facilities of Andalmar Biotech S.L., where they were distributed in floating baskets in a concrete 8000 L outdoor re-circulation tank. The tank was equipped with (filtration system details). And used natural seawater, obtained by pumping from a well and filtered before flowing into the aquaculture circuit. (Nº of baskets and anemones, scale of the experiment).

Other organisms with different trophic level were introduced in the tank so as to create an IMTA environment (Coll et al., 2025). These organisms were sea urchins (*Paracentrotus lividus*), sea snails (Monodonta turbinata), sea cucumbers (*Holothuria tubulosa*) and macroalgae (*Ulva rigida*, *Cystoseira mediterranea*). (Include density table?)

Inone month after the beginning of the experimental period, (Nº) anemones from different baskets were selected as experimental group, and they were sectioned in two through their pharyngeal biradial axis. Both halves were then left to heal, so as to generate two different clonal individuals. (More information on healing process).

4 weeks after the sectioning procedure, once the anemones had had time to heal completely a first sampling (T1) took place ,. (Mortality). 9 sectioned anemones were selected (3 per bascket), as well as 9 control anemones (3 per basket) which had not undergone the procedure. The second sampling (T2) took place 16 weeks after, in month year, and again 9 control anemones and 9 sectioned anemones were collected. In both samples, each anemone was snap-frozen in liquid nitrogen and stored at -80 ºC. The column and tentacles of each individual was homogenized separately (Heidolph Instruments) in 100 mM Tris, 0.1 EDTA and 0.1% Triton buffer (pH 7.8) at a 1:4 ratio (w/v). Extracts were then centrifuged at 16 000 rpm for 30 minutes at 4 ºC (Sigma 3 K30), and the supernatant was collected and stored at -80 ºC for posterior analysis.

2.2. Oxidative status assays

Superoxide dismutase (SOD) (EC 1.15.1.1) activity was determined according to McCord & Fridovich (1969) method, consisting on an indirect measurement as the degree of inhibition of cytochrome c reduction. Determination of catalase (CAT) (EC 1.11.1.6) activity was performed using Aebi (1984) method, based on the decrease in absorbance produced by H2O2 consumption by this enzyme.

Glutathione peroxidase (GPx) (EC 1.11.1.9) activity of the samples was determined following Flohé & Günzler (1984), based on an indirect measurement of NADPH oxidation, generated by its coupling with a standard glutathione reductase (GR) (EC 1.8.1.7) reaction. GR activity was measured according to Carlberg & Mannervik (1975) method, consisting on a measure of the absorbance decrement caused by NADPH oxidation.

Glucose 6-phosphate dehydrogenase (G6PDH) (EC 1.1.1.49) activity of the samples was obtained using a modified method of Löhr & Waller (1965), based on recording the change in absorbance due to NADPH production by the enzyme.

Determination of glutathione S-transferase (GST) (EC 2.5.1.18) activity was performed following the method of Frasco & Guilhermino (2002), based on the formation of a conjugate between glutathione and 2,4-dinitrochlorobenzene that increases absorbance. DT-diaphorase (DTD) (EC 1.6.99.2) activity was determined using a modified method of Lemaire et al. (1996), based on measurement of decreased absorbance by reduction of 2,6-dichlorophenol indophenol,

Soluble protein content was quantified following Bradford (1976) method in order to express enzymatic specific activity. One unit of activity was defined as the amount of enzyme required to transform one μmol of substrate per minute under the measurement conditions (Coll et al., 2025). For SOD, units of activity had a different definition, as the amount of enzyme required to generate a 50% inhibition in the reduction of cytochrome c.

Total antioxidant activity was carried on according to Erel (2004) method, based on the change of absorbance due to reduction of 2,2′-azino-bis-(3-ethylbenzothiazoline-6-sulfonic acid)to determine Trolox-equivalent antioxidant capacity (TEAC) of the extracts,.

Lastly, oxidative damage to lipids was assayed according to thiobarbituric acid reactive substances (TBARS) contentfollowing a modified method from Buege & Aust (1978) and using malondialdehyde (MDA) as a standard.

2.3. Non-specific immune parameters

Acid (EC 3.1.3.2) and alkaline (EC 3.1.3.1) phosphatases (AP and AlP) activity were measured following the method of Easy & Ross (2010) and Huang et al. (2011) respectively?, based on the change of absorbance at 405 nm produced by the activity of the enzyme on p-nitrophenyl at different pH. Lysozyme activity (EC 3.2.1.17) was determined according to Swain et al. (2007), by using a M*icrococcus lysodeikticus* suspension and an egg lysozyme standard. Determination of myeloperoxidase (MPx) activity (EC 1.11.1.7) was carried out following a modified method of Mohanty & Sahoo (2010), based on the increase of absorbance due to oxidation of 3,3',5,5'-tetramethylbenzidine (TMB) by the products of MPx activity.

Both oxidative status and immune parameters measurements were carried out with a PowerWave microplate spectrophotometer (Bio-Tek Instrument, Inc.) at a stable temperature of 25 °C for enzymatic determinations.

2.4. Statistical analysis

All statistical analysis and data processing was carried out using R. 4.4.3 and Rstudio 2024.12.1. Results were expressed as mean ± standard error of the mean (SEM). A confidence level of 95% (α = 0.05) was established for all statistical tests. For each variable, a two-way ANOVA was conducted to examine the effect of the sectioning procedure at short and long term on the oxidative status of *A. viridis*. When interaction between both variables was significant, a t-test was performed at each level of the variable time (T1 and T2) to test for differences between sectioned and control animals. Normality of residuals was assessed via Shapiro-Wilk’s normality test, while homoscedasticity was tested using Levene’s test. All residuals were normally distributed (p > 0.05) and all variables had homogeneity of variances (p > 0.05). Obtained p-values were adjusted using Benjamini-Hochberg correction for multiple testing.

3. Results

3.1. Oxidative status parameters

SOD and CAT

According to SOD activity in handled and non handled groups in tentacle and columnar tissue of *A .viridis* at different seasonal time…(Figure 1) interaction between the two variables was not significant, and there was no effect of the sectioning procedure or the time variable on SOD activity, neither on columnar nor tentacular samples. A light tendency was found on tentacular SOD activity, where sectioned individuals yielded a slightly smaller activity, but this pattern was not found to be statistically significant. CAT activity (Figure 2) showed different responses in tentacular and columnar samples. Columnar activity was increased by the sectioning procedure, while there was no effect of time or interaction between both variables. Tentacular CAT activity, however, featured a significant interaction effect. At T1, sectioned anemones displayed a significantly higher CAT activity than control anemones (p = 0.000274, p < 0.001). However, at T2, this relationship became inverted and the sectioned individuals exhibited a lower activity (p = 0.00599, p < 0.01).

GR activity is displayed in Figure 3. Time had a significant effect on both columnar and tentacular GR, as samples from T2 exhibited lower activity than T0 measurements. Furthermore, columnar GR activity was significantly increased for all sectioned individuals.

Figure 4 shows GST activity on both body regions. While there were no significant differences in columnar GST activity, tentacular samples featured significant interaction between the two variables. At T1, sectioned individuals had their GST activity significantly increased (p = 0.000656, p < 0.001). These differences were not reflected T2, where control and sectioned individuals were found to be similar (p = 0.345). ´

Columnar DTD activity (Figure 5. A) increased significantly on sectioned anemones. Tentacular DTD activity featured a significant interaction effect (Figure 5.B). Anemones at T1 exhibited this same pattern of higher activity for sectioned individuals (p = 0.000103, p < 0.001) detected in columnar activity. At T2, however, no differences were found between control and sectioned samples (p = 0.393).

Total Antioxidant Capacity, measured as TEAC, is displayed on Figure 6. Neither columnar or tentacular TEAC showed any differences associated with sectioning or time.

Lipid peroxidation, measured as MDA concentration (Figure 7) did not vary significantly in columnar samples, but tentacular MDA was found to be lower on sectioned individuals compared to control ones.

3.2. immune status

Neither acid phosphatase (AP, Figure 8) nor alkaline phosphatase (AlP, Figure 9) showed any significant effect of handling or interaction between…….

MPx activity showed significant interaction in columnar samples. At T1 columnar MPx activity was lower for sectioned individuals (p = 0.000895, p < 0.001), while there were no significant differences at T2 (p = 0.674). The effect of sectioning and time on tentacular activity also resulted non-significant.

Columnar lysozyme activity showed no significant differences associated with the studied variables.

4. Discussion

5. Conclussions

6. References

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