Interaction between a mobile and a sessile species in SW Atlantic mudflats: Lack of negative effects of the mud crab *Cyrtograpsus angulatus* Dana, 1851 on filtration rate and growth of the infaunal clam *Tagelus plebeius* (Lightfoot, 1786)

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**A B S T R A C T**

Biogenic modifications of benthic environment by organism’s activities like feeding, burrowing or excretion have in general negative implications for cohabiting sessile infauna, especially for suspension-feeders. However, the outcome of these interactions depends on several biological and physical characteristics and thus, it is not easily predictable in soft bottom environments. The stout razor clam *Tagelus plebeius*, an estuarine, infaunal and sessile filter feeder species, coexists in Mar Chiquita coastal lagoon (Argentina, 37° 32'S, 57° 19'W) with the mud crab *Cyrtograpsus angulatus*, a very mobile benthic consumer. Due to the intense crab activity and consequent habitat transformation, we predicted negative effects on clams’ filtration rate and their growth. However, an inclusion experiment showed that *T. plebeius* filtration rate was higher in enclosures when crabs were present responding to enhanced turbidity and chlorophyll a. Additionally, another experiment showed that mean condition index as well as mean shell growth increment of clams was not negatively affected by crabs, although maximum values of shell growth increments (mm) were smaller in crab inclusions. Our findings suggest that the clams can exploit the increased food availability (i.e., Chl a) generated by resuspension and are also not affected by the disturbance that crabs can make over sediment surface. In conclusion, contrary to the generalized outcome proposed for this kind of interaction, crabs do not have a negative effect on *T. plebeius*.

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1. Introduction

Benthic species assemblage and interactions are frequently modified by the mobility of the organisms and habitat biogenic transformations (Murray et al., 2002). Through feeding, excretion, burrowing and bioconstruction, macrofauna influences superficial particle transport (Rosenberg, 2001), changes sediment composition and stability (Widdows and Brinsley, 2002), and changes the flux of particles (Boudreau, 1992) and solutes (Michaud et al., 2006) between the sediment and the overlying water (i.e. sediment–water interface). Sessile infauna is especially affected by these changes in the physical environment. For example, changes in physical characteristics of sediment–water interface generated by biotic processes affect filtration activity of benthic filter-feeding species (Pillay et al., 2007) since this interface provide their food supply (Herman et al., 1999). Generally, sediment disruption by animal activities (e.g. feeding, burrowing) is predicted to have negative effects on suspension-feeding infauna (Rhoads and Young, 1970; Wilson, 1991). Nonetheless, it depends on other factors such as sediment properties and/or characteristics and functional traits of interacting species (e.g. body size, frequency of disturbance; Posey, 1987; Snelgrove and Butman, 1994). For example, filtration rate of bivalves may respond positively to increased seston load (Jørgensen, 1996). However, extremely high seston concentrations may lead to clogging of filtration apparatus with negative consequences on growth and survival (Jørgensen, 1996; Ellis et al., 2002). Thus, the outcome of these interactions in soft-bottom sediments is not easily predictable.

The stout razor clam *Tagelus plebeius* is an estuarine, filter feeder species which is distributed along the American Atlantic coast, from Cape Cod (42° N, Mass. USA; Leal, 2002) to the Northern Argentinean Patagonia (41° S, Argentina; Scarabino, 1977), although this distribution is not continuous. This clam is often an important infaunal community component (e.g. Seitz et al., 2006) inhabiting permanent burrows (Gutiérrez and Iribarne, 2004) up to 70 cm depth in estuarine tidal flats with cohesive sandy silt sediments (Holland and Dean, 1977) and salinities ranging from 41 to 2%o (Addino, 2014). The location of
clams is visible by the presence of their siphon holes in the sediment surface (Gutiérrez and Iribarne, 2004) reaching densities of up to 120 ind m⁻² in the low intertidal (Alvarez, 2013). Stable isotope analysis revealed that in the Southwest Atlantic estuaries suspended benthic microalgae and phytoplankton comprise its main food supply (Botto et al., 2005). In its southernmost distribution range T. plebeius coexists with the mud crab Cystograpsus angulatus (41° S, Argentina to 27° S, Brazil; Bosch; 1964) which is an omnivorous benthic consumer (Schwindt et al., 2001; Botto et al., 2005) that inhabits predominantly the lower intertidal and the adjacent subtidal level (Spivak et al., 1994). In estuaries, the adults of this species are found in soft-muddy bottoms characterized by high densities of detritivorous polychaetes mostly of the genus Heteromastus and Laeonoereis (Spivak, 1997). In Mar Chiquita lagoon crabs predate these annelids (Olivier et al., 1972; Schwindt et al., 2001) and also consume pelagic and benthic microalgae, some meiofaunal organisms and plant debris (Schwindt et al., 2001), being also scavengers (Olivier et al., 1972). The adult crabs take shelter remaining superficially buried in the sediment (Spivak, 1997) up to 4 cm depth (Schwindt et al., 2001). In fact, Schwindt et al. (2001) reported sediment reworking by C. angulatus to a depth of 5 cm in some areas of the lagoon. Moreover, they suspend sediment through locomotion (pers. obs.). They also show large spatial displacement mainly related to tidal water movement, usually concentrating many crabs along the water edge (Boschi, 1964; Spivak et al., 1996). Due to this high mobility and great small-scale variability quantitative studies of its population dynamics are very difficult. However, rough estimates in Mar Chiquita coastal lagoon indicate that its density varies from clusters of thousands of crabs near the mouth to single individuals in the inner areas of the lagoon (Spivak et al., 1994). Therefore, due to its feeding habits, superficial burrowing and displacement they could lead to disruption and resuspension of superficial sediments acting as a biodiffuser (see François et al., 2002) and thus modifying sediment–water interface with possible consequences for clam feeding.

Furthermore, it is known that in the SW Atlantic Mar Chiquita coastal lagoon (37°22'S, 57°16'W, Argentina) other crab species (Neohelice (= Chasmagnatus) granulata) negatively affects growth and condition index of T. plebeius due to both, direct contact with consequent shell damage and indirect mechanisms probably related to behavioural changes in clams and their associated energy costs (Lomovasky et al., 2006). These effects are stronger in the lower intertidal and it is suggested that even the displacement of crabs over the sediment surface can lead those behavioural changes affecting energy budget of clams (Lomovasky et al., 2006). In this sense, considering that C. angulatus also wander over the sediment surface, it is probable that its activity over siphon holes (i.e. over the bottom) also disturbs clams. Thus, we expect changes in the clam’s behaviour such as escape responses or lower feeding activity with consequences on the clam’s energy budget and growth. In this context, the aims of this study were to evaluate if 1) the crab C. angulatus modify particle load in the sediment–water interface and thus, filtration activity of the razor clam T. plebeius and 2) if crab presence affects shell and somatic growth of clams.

2. Materials and methods

2.1. Study site

Samplings and field experiments were made in Mar Chiquita coastal lagoon in an area called CELPA located approximately 2.5 km from the lagoon inlet (Fig. 1), at the low intertidal level (0.40 m above mean low tide, following Lomovasky et al., 2006) where sediment mean grain size is 115 μm (SD = 9.55, range = 103.57–131.75) and mean turbidity is 3.06 NTU (0.84, range = 1.55–4.44) (Addino, unpublished data). The lagoon is a brackish water area of 46 km² permanently connected to the sea (Isla, 1997). The entrance channel is approximately 6 km long, 200 m width and its depth is between 1.5 and 2 m (Fasano et al., 1982; Reta et al., 2001). It is characterized by small tidal amplitudes (up to 1 m, Isla, 1997) which are mainly generated by the winds and rain (Isla and Garrido, 2001).

2.2. Displacement of C. angulatus in the tidal flat

To corroborate the pattern of crab displacement with tidal water movement (see Introduction), field observations were made in spring 2009. Fifty eight fixed points were randomly stated (marked with a fine wire stake protruding a few centimetres from the sediment) in the tidal flat. Then the number of crabs that cross through a 10 cm diameter area around each point during 10 min was observed. During the observations, we could see that the use of the marks did not modify the behaviour of crabs (i.e. crabs did not stay around the stakes). Observations were made over a mudflat extent of approximately 500 m in three states of the tidal cycle defined as: stable high tide, flooding and ebbing. Observations were made simultaneously by 3 observers which could see 2 points at the same time, thus the observation periods were conducted for no more than 1 h and 40 min at a time. The null hypothesis of no difference in mean crabs h⁻¹ crossing around a fixed point between tidal moments was tested by ANOVA. Data were square-root transformed to fulfil the analysis assumptions (Zar, 1999).

2.3. Effect of C. angulatus on suspended material and filtration activity of T. plebeius

To evaluate 1) if C. angulatus changes the particle load and clam’s food availability in the sediment–water interface and 2) the effect on clam’s filtration rate (FR), an inclusion/exclusion field experiment was carried out in spring 2009. Thirty two plastic cylinders (20 cm height, 18 cm diameter) were established in the mudflat where they were flooded every high tide. Cylinders were buried 10 cm into the sediment to ensure no exchange of water during sampling procedure in ebbing tide. The volume into the containers (delimited by cylinder walls and sediment surface) was 2.5 L. Each of 16 cylinders contained one clam which was already buried there since their siphon holes were visible in the sediment surface. The remaining 16 containers were established above sediment without clams. After 7 days of acclimation we were able to use 14 with clam and 11 without, the other containers were lost by storms or discarded because of the intrusion of crabs. Water samples from the sediment–water interface were taken in the following way: just after the tide ebbed when the containers were uncovered but full of water (time 0) the first sample was taken by carefully submerging a 3 ml plastic tube. This procedure was made the more closely possible to the sediment surface but carefully enough to avoid sediment removal. Also, we ensure that the procedure was exactly the same in all treatments; hence any difference found is due to the treatment itself. Immediately after, 1 crab (∼3 cm carapace) was added into each of 14 containers, 7 with clam and 7 without clam. Thus, we completed 4 treatments: only clam (Clam, n = 7), clam and crab (Clam–Crab, n = 7), only crab (Crab, n = 7) and control (without clam and crab, Control, n = 4). Thereafter, a water sample (3 ml) was taken from each container every 10 min in a total period of 40 min (five samples in total). Immediately after water samples were taken turbidity (NTU) and Chl a (μg l⁻¹) were measured in the field with an Aquaflo handheld Fluorometer (Turner Design, Model 8000-01, detection range: Turbidity = 0.5–150 NTU, Chl a = 0.05–300 μg l⁻¹) following Bruschetti et al. (2008). Turbidity was used as representative of particle load in the sediment–water interface (Ellis et al., 2002) and Chl a as food availability. To evaluate the effect of crab on sediment–water interface we compared water turbidity and Chl a between treatments Crab and Control at the beginning (time 0) and at the end of the experimental time (at 40 min) by t-test (corrected t-test, Welch approximation, Zar, 1999). As well as evaluating if crabs affect FR of clams, we compared FR (see calculation details below) between treatments Clam and Clam–Crab in each time interval of 10 min by t-test. Before this final test (at the end of the experiment) clams were extracted and before their length (L,
anterior–posterior axis; precision ±0.01 mm) was measured to compare mean L between treatments and to analyse the relationship between L and FR by simple regressions (Zar, 1999) to dismiss the effect of clam’s size on FR (see Section 3.2).

The FR \((1\ h^{-1})\) of clams was calculated with the indirect method (following Cohen et al., 1984) which measures the rate of exponential decay of particle concentration in a time interval:

\[
\text{FR} = \frac{V_i \times \left( \ln \left( \frac{T_{r_b}}{T_{r_a}} \right) - \ln \left( \frac{T'_{r_b}}{T'_{r_a}} \right) \right)}{t}
\]

where \(V_i\) is the water volume \((l)\) into the containers, \(T_{r_a}\) and \(T_{r_b}\) are turbidity values at the beginning and at the end of a time interval respectively, and \(t\) is the time elapsed in a time interval \((i.e. = 10\ min)\). \(T'_{r_a}\) and \(T'_{r_b}\) are also the turbidity values at the beginning and at the end of a time interval but in containers without filter-feeder \((i.e.\ without\ clam)\). These latest values are included in Eq. (1) to measure and subtract the natural settling rate of particles from the actual FR (see Cohen et al., 1984 and Riisgård, 2001 for details). To estimate FR in the Clam treatment we used natural settling rate \((i.e.,\ ln \left( \frac{T_{r_b}}{T_{r_a}} \right))\) in containers from the Control treatment, while FR in the Clam–Crab treatment was estimated using natural settling rate in the Crab treatment.

Since this method is based in the decay of particle concentration, a turbidity increment \((resuspension)\) instead of a reduction from the beginning to the end of a time interval implies that the result of the Eq. (1) is negative \((e.g.\ Fabricius\ et\ al.,\ 1998;\ Grizzle\ et\ al.,\ 2008)\). Thus, given that clams can resuspend material enhancing turbidity in the containers mainly by ejection of faeces and pseudofaeces \((\text{see Graf and Rosenberg, 1997})\) it is possible to find negative FR values. Besides, natural particle settling occur constantly in the containers without filter-feeder. This highlights one of the main limitations of this method \((\text{see Riisgård, 2001})\) as it can also lead to negative values when this settling is high.

2.4. Effects of C. angulatus on T. plebeius shell growth and condition index

To evaluate the effect of C. angulatus on shell growth and condition index (CI) of T. plebeius, a second crab Exclusion/Inclusion experiment was carried out between September 2009 and May 2011. Twenty plastic square cages \((1\ cm\ mesh\ and\ 50\ cm\ side)\) were deployed on the intertidal mud flat. Clams \((120)\) were extracted from the mudflat with a shovel, measured in L and height \((H,\ umbo-ventral\ axis;\ precision \pm 0.01\ mm)\) and marked with a plastic number glued to the periostracum surface with cyanoacrylate \((\text{superglue})\) to be identified later. Six marked clams
between 54 and 66.3 mm L were set into each cage (reflecting the natural density of clams in the area: 36 clams m$^{-2}$, Addino et al., 2010). In 10 of the cages, crabs were then added (3 males and 2 females respecting natural sex proportion and density following Spivak et al. (1996), inclusion treatment). Crab density into cages was evaluated every 15 days and maintained adding crabs if necessary.

At the end of the experimental time all cages were recovered (except one from exclusion treatment) and all alive and marked clams were collected accounting for a total of 50 clams (nearly 3 clams per cage for all treatments). Final L and H (i.e. L and H at the end of the experiment) were measured and CI was calculated using the dry weight of soft tissue (dried at 80 °C to constant weight) and the L of clams with the following equation (Clausen and Riisgård, 1996):

$$\text{Cl} = \text{soft tissue L} - b$$

where b (2.93) is the slope of the regression curve between L and soft tissue for these clams. The growth increments in L and H (i.e. the increase in size (L and H), hereafter L increment and H increment) of each clam were calculated as: final L − initial L and final H − initial H (mm), respectively. Also their relationship with initial L and H, respectively, were evaluated by simple regression models. Given that there were no relationships, except for H increments only in exclusion treatment (see Section 3.3), mean L increment and mean H increment were evaluated in further statistical analysis. To perform a more powerful test we looked for differences in mean CI, L increment and H increment between replicates (cages) into each treatment by ANOVA analysis (three separate analysis). As differences were not found (see Table 1 data were pooled into each treatment. Thus, we evaluated the null hypothesis of no difference in 1) mean CI, 2) mean shell L increment and 3) mean shell H increment between treatments through t$_c$ test (3 analysis in total, Zar, 1999).

### 3. Results

#### 3.1. Displacement of C. angulatus in the mudflat

The number of crabs h$^{-1}$ crossing around a fixed point showed differences between tidal moments being higher during flooding tide (df = 171, F = 82.2, p < 0.001; mean crabs h$^{-1}$ (SD): stable = 2.5 (3.4), flooding = 18.7 (12.2), ebbing = 2.9 (4.2)). If we extrapolate these 10 min of observation to the total time period in which tide is flooding (around 5 h in our study site; M. Addino, pers. obs.) nearly 95 crabs are potentially crossing above the siphon holes of a clam (a fixed point) in this time period.

#### 3.2. Effect of C. angulatus on suspended material and filtration activity of T. plebeius

Turbidity and Chl $a$ were not different between Crab and Control at time 0, but they were both higher in Crab treatment 40 min after the beginning (Fig. 2A and B) indicating particle resuspension. Mean shell length (L) was not different between treatments Clam and Clam–Crab (df = 10, t$_c$ = 1.62, p = 0.14) and there was no relationship between L and filtration rate (FR) of clams (df = 4, F = 1.54 and 0.11, p = 0.28 and 0.75; for Clam and Clam–Crab respectively). Mean FR was different between treatments Clam and Clam–Crab being higher in the latter in the last time interval (Fig. 3) showing a positive response to enhanced food supply. Statistical data for turbidity, Chl $a$ and FR are summarized in Table 2.

#### 3.3. Effects of C. angulatus on T. plebeius somatic and shell growth

Mean condition index (CI) did not show differences between treatments (mean CI (SD) = 0.0039 (0.0008) and 0.0041 (0.0008) for inclusion and exclusion, respectively, df = 44, t$_c$ = −0.91, p = 0.37). There was no relationship between the initial L and the L increments of clams in any treatment (df = 15 and 29, F = 0.3 and 0.93, p = 0.59 and 0.35, for inclusion and exclusion, respectively). Increments in height (H) showed a negative linear relationship with the initial H only in exclusion treatment (df = 15 and 29, F = 0.03 and 12.9, p = 0.86 and 0.0014; for inclusion and exclusion, respectively, Fig. 4). Also, the mean shell L increment as well as H increment was not different

### Table 1

Comparison of CI and shell length increment and height increment between replicates in to each treatment of the second Exclusion/Inclusion experiment.

| Treatment | Variable      | df | F     | p-Value |
|-----------|---------------|----|-------|---------|
| Inclusion | CI            | 5  | 2.79  | 0.11    |
|           | Shell L increment | 5  | 1.11  | 0.42    |
|           | Shell H increment | 5  | 1.14  | 0.41    |
| Exclusion | CI            | 7  | 2.06  | 0.10    |
|           | Shell L increment | 7  | 0.72  | 0.66    |
|           | Shell H increment | 7  | 1.32  | 0.29    |
between treatments (L increment: \( df = 41.99, t_c = -1.46, p = 0.15 \); H increment: \( df = 42.97, t_c = -0.66, p = 0.51 \)). However, both shell L and H increments showed smaller variances in the inclusion treatment (Levene test, \( df = 43, F = 2.68 \) and 6.44, \( p = 0.043 \) and 0.015 for L and H increment respectively; Fig. 5A and B).

4. Discussion

Our results showed that as expected *C. angulatus* present higher migration in the intertidal flat during flooding tide following the tidal water movement. Turbidity and Chl a were enhanced by crabs after 40 min under experimental conditions supporting our suggestions that crabs may act as biodiffusers. Parallel to those changes in the sediment–water interface, FR of clams was higher when crabs were present indicating that there was no negative effect on *T. plebeius*. Besides, mean clam CI and mean shell growth increments were not affected by crab presence. However, shell increments both in length as in height presented smaller maximum values in crab inclusions.

The different consumer modes generate different modifications in the sediment–water interface (Nogaro et al., 2009). Bioturbators may also modify resuspension by water turbulence (Davis, 1993; Cadée, 2001) and all together generate different effects on coexisting filter-feeding organism. In previous studies, *C. angulatus* is not commonly considered a bioturbator in the Mar Chiquita coastal lagoon (but see Martinetto et al.

### Table 2

| Variable                  | Treatments compared | Time (min) | df   | t-value | p-value |
|---------------------------|---------------------|------------|------|---------|---------|
| Turbidity                 | Crab–Control        | 0          | 8    | 1.08    | 0.31    |
|                           |                     | 40         | 8    | 2.88    | 0.02*   |
| Chl a                     | Crab–Control        | 0          | 8    | 1.19    | 0.27    |
|                           |                     | 40         | 8    | 3.79    | 0.005*  |
| Filtration rate (l h\(^{-1}\)) | Clam–Clam–Crab   | 0–10       | 12   | 1.28    | 0.23    |
|                           |                     | 10–20      | 12   | -0.13   | 0.89    |
|                           |                     | 20–30      | 12   | 0.65    | 0.53    |
|                           |                     | 30–40      | 12   | 3.33    | 0.006*  |

* Denotes significant differences.
filtration activity is not physiologically regulated but it is determined by the physical capacity of filtration apparatus and the environmental prevalent food concentration (Hawkins et al., 1996; Jørgensen, 1996). Nevertheless, enhanced food acquisition may have costs due to pseudofaeces production and thus, a negative impact (e.g. Bayne et al., 1987; Ellis et al., 2002). Therefore, the lack of knowledge about the costs associated to the higher FR of *T. plebeius* makes it difficult to conclude if this effect is really positive for this clam. However, even when turbidity generated by crabs is almost ten times greater than in the natural conditions (this study); there was no negative effect on clams, such as stuck of filtration apparatus which is commonly predicted (Rhoads and Young, 1970; Wilson, 1991) for this kind of interaction.

Negative effects that mobile species generate on sessile organisms can be mediated by direct contact (e.g. Lomovasky et al., 2006) or more frequently by indirect mechanisms (e.g. Murphy, 1985; Nakaoka, 2000; Mermillod-Blondin and Rosenberg, 2006). In the case of *T. plebeius* and *C. angulatus* we suggested that crabs negatively affect the clam’s growth by indirect mechanisms that change clams’ behaviour. However, in our study the second inclusion/exclusion field experiment showed no effect of *C. angulatus* on mean clams CI neither on mean clam shell growth increments. Nevertheless, only for clams with initial height between 17.5 and 20.5 mm the height growth increment was smaller in inclusion treatment. That is, in smallest clams the height increment was diminished. Besides, the maximum values of shell increment, both in length and in height were smaller in inclusion treatment which leads to smaller variances. This means that when crabs are present, shell growth may be partially limited leading to smaller maximum clam sizes. This result may also be explained by changes in clam behaviour (e.g. escape behaviour, feeding interruption) and consequent energy cost. However, on average this effect is negligible in the case of the interaction between *C. angulatus* and *T. plebeius*.

Filtration capacity is adapted to the prevalent natural food concentration in the environment and thus, this trait maximizes resource use allowing bivalves to almost completely exploit their growth and/or reproductive potential (Jørgensen, 1996). We found that the crabs can increase clams’ resources by resuspension under experimental conditions without generating negative effects on clams such as clogging of filtration apparatus. In addition, we found no effect of crab presence on clam’s growth. Thus, our findings show that clams can exploit the increased food availability (i.e., Chl a) generated by biological resuspension and are also adapted to the disturbance that the crabs can make over siphon holes. In conclusion, contrary to the generalized outcome proposed for this kind of interaction, in this case crabs do not have a negative effect on *T. plebeius*.
