Contribution of biofilm to water quality, survival and growth of juveniles of the freshwater crayfish *Cherax quadricarinatus* (Decapoda, Parastacidae)

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

The effect of biofilm as an alternative food source and/or a complement for improving culture was assayed for early and advanced juveniles of *Cherax quadricarinatus*. For both kinds of juveniles, higher values of survival were seen in the experimental groups provided with either biofilm (B) or a combination of both formulated food and biofilm (B+F), compared to the group only receiving formulated food (F). Such higher survival was associated to a better water quality maintained by biofilm, in terms of low levels of both ammonium and nitrite, together with high levels of pH and dissolved oxygen. As for growing, specific growth rate was higher in the groups fed with formulated food, but only for early juveniles. Considering the crayfish biomass at the end of the experiment (i.e., an integrative index of both survival and growth), the best results were seen in the B+F group, for both kind of juveniles. The main micro-organisms present in biofilm were chlorophyta, xanthophytas, pennate diatoms, cyanobacteria, flagellates, ciliates, rotifers and nematodes. Most of these items were found in the stomach of crayfishes fed on biofilm. The hepatopancreatic levels of total lipids were higher in animals of both B+F and F groups, compared to those of B group, while energetic reserves in the abdominal muscle showed no differences among experimental groups, for any kind of juveniles. Therefore, biofilm could be considered as a good complement for the culture system of *C. quadricarinatus* juveniles, mainly by improving survival through the maintenance of a good water quality. Combination of biofilm and formulated food has shown the best results, in terms of both survival and growth of juvenile crayfish.

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1. Introduction

*Cherax quadricarinatus* (von Martens), or redclaw, is a crayfish species native to the tropical region of Queensland, northern Australia, with high commercial potential. Production of this species on farms, either in Australia or other countries, has been significantly increased during the last decades, due to the advantages for its culture (Medley et al., 1994). According to Jones (1995, 1997), life cycle of *C. quadricarinatus* comprises a relatively long embryonic development (6 to 10 weeks), during which successive larval stages take place within the egg, to finally hatch a first juvenile stage. After two successive molts, juveniles become completely independent from their mother. In farms of Argentina, the culture of this species starts with an indoor incubation and growing stage of about 3 to 4 months long. After hatching, recently independent (early) juveniles are commonly maintained in hatcheries for 2 to 4 weeks. Advanced juveniles (near 1 g of body weight) are later transferred to larger tanks (nurseries) until they achieve a body weight of near 5 g. After that, juveniles are maintained in earthen ponds for 6 to 9 months, until they reach a market weight of 50 g or higher.

*C. quadricarinatus* is a facultative omnivorous species; its diet includes both plant and animal materials, as well as detritus. The gastric mill of *C. quadricarinatus* juveniles has been described as morphologically adapted for the consumption of both detritus and zooplankton (Loya-Javellana et al., 1994). Some studies have shown that in culture conditions, zooplankton leads to a higher crayfish growth than other food sources (Austin et al., 1997; Jones, 1995; Verhoef et al., 1998). Consumption of phytoplankton by crayfish under culture conditions has not been previously reported.

Several formulated diets have been developed to achieve an acceptable survival and growth of *C. quadricarinatus* juveniles, optimizing the percentage of protein, lipids and other nutrients (Campaña-Torres et al., 2006; Cortés-Jacinto et al., 2003; Gutiérrez and Rodríguez, 2010). However, formulated food represents 60% of the total operational costs of aquaculture farms (Tacon, 1999), which can be even higher in intensive cultures.

The term biofilm is referring to a micro-organisms community, associated to an organic matrix formed on a surface submerged in a...
water body. Such micro-organisms are involved in the transference of organic matter through most trophic chains and in the biochemical cycles verified in aquatic ecosystems (Decho, 1990; Meyer-Reil, 1994). Different cultured crustacean species have improved either growth or survival when biofilm was provided to them as a food source (Abreu et al., 2007; Ballester et al., 2007; Thompson et al., 2002, among others). Besides, the water quality of culture systems was frequently improved by the utilization of biofilm (Bratvold and Browdy, 2001; Holl et al., 2011; Thompson et al., 2002, among others). However, only one preliminary study has been reported on a crayfish species ( Cherax destructor) concerning the possible benefit of biofilm (Jones and Thanuthong, 2002).

The current study was aimed at evaluating the effect of biofilm, as an alternative and/or complementary food source, on survival and growth of both early and advanced juveniles of C. quadricarinatus. The effect of biofilm on water quality maintenance was also evaluated.

2. Materials and methods

2.1. Experimental design

C. quadricarinatus ovigerous females were purchased from a local dealer (Pinzas Rojas SRL, Tucumán, Argentina). A short-time after hatching early juveniles (N = 288, mean weight 0.10 ± 0.05 g), corresponding to the first independent juvenile instars, were selected for the experiment. Besides, another group of early juveniles was fed every day with a commercial diet (TetraColor®; 47.5% crude protein) for one month, in order to obtain advanced juveniles (N = 234, mean weight 0.98 ± 0.43 g) for a second experiment.

Each experiment comprised 30 days, beginning by early February (early juveniles) or by early March (advanced juveniles); all the experiments were carried out in Buenos Aires (34° 36 W, 58° 23 S). A zero water exchange system was used, only adding dechlorinated water, were employed as experimental units (N=9 to each group of juveniles). A number of either 32 early juveniles or 26 advanced juveniles were assigned to each aquarium, yielding a density of 133 and 111 animals/m², respectively. These densities were similar to others previously used for juveniles of the same species (Jones, 1995; Masser and Rouse, 1997). A mean water temperature of 25±2 °C and a pH of 7.8 ± 0.5 were maintained throughout the experimental period.

The following experimental groups, by triplicate, were run for each group of juveniles:

- B juveniles only fed on biofilm.
- B+F juveniles fed with either biofilm and formulated fish meal (TetraColor®; 47.5% crude protein).
- F juveniles only fed on formulated fish meal (same as above).

Formulated meal, successfully used in previous studies on C. quadricarinatus (Chaulet et al., 2008; Viau and Rodríguez, 2010) was given every day at a percentage of 10 (early) or 5 (advanced) of juvenile biomass, in specially feeding trays, designed according to Wasselesky et al. (2006). Briefly, each tray was composed by a cylindrical plastic base (15 cm de diameter), coupled to a vertical PVC pipe (30 cm long, 1.5 inch diameter) for loading food. Two trials were placed in each aquarium, in order to ensure that all juvenile could easily access to the food provided.

Biofilm used to feed juveniles was developed during 30 d before starting the experiment, on plastic nets (50 x 27 cm, 1 mm mesh size) vertically placed in aquaria containing a heterotrophic medium, prepared from an initial inoculum taken from natural body water and added as 15% of the aquaria volume water. To help biofilm development a little portion (50 mg/aquarium/day) of the commercial diet was added, as a source of nitrogenous and phosphorous. The biofilm was considered as mature when chlorophyll a reached a concentration near 5 μg/cm², according to Thompson et al. (2002). At this time, 4 plastic sheets were transferred to each aquarium assigned to both B and B + F groups. Group F received the same number of plastic sheets with no biofilm formed, that were also periodically replaced during the experiment, to avoid biofilm formation. Additionally, aquaria of all groups received some PVC pipes as refuges (32 of 1 in. and 26 of 1.5 in. diameter, for each aquarium of early and advanced juveniles, respectively), which were daily withdrawn for cleaning them of any biofilm eventually formed.

2.2. Biofilm monitoring

Weekly, three small samples (4 x 4cm) were cut from the plastic nets of each aquarium corresponding to both B and B + F groups, at 20 cm deep, in order to determine the biofilm dry weight and chlorophyll a concentration, as well to characterize the micro-organisms presented. Additionally, samples of sheets were also taken from the F group, in order to check if any biofilm formation took place.

To determine dry weight, the first sample was placed in stove at 60 °C until constant weight, estimating by difference the biofilm dry biomass at a precision of ± 0.1 mg. In the second sample, chlorophyll a concentration was determined after removing fresh biofilm from plastic with a soft brush; biological material was then filtered through a micro-fiberglass GF/F (Munktell®, 47 mm diameter, 0.7 μm pore size). Finally, the filter was placed in a glass vial containing 8 ml 60% ethanol in total darkness, for 24 h. Chlorophyll a concentration was determined spectrometrically at 665 and 750 nm, before and after the acidification with HCl 0.1 N, according to the equations proposed by Marker et al. (1980).

To characterize micro-organisms, the biofilm of the third sample was gently removed and preserved in a glass vial filled with 20 ml of distilled water and formaldehyde at 4% final concentration. Later, three aliquots were separately and systematically analyzed under binocular microscope (Nikon SMZ 645, 100 to 1000× magnifications), to identify the main items presented. Determination of genera was made according previous studies (Canter-Lund and Lund, 1995; Da Cunha, 1913; Hartley, 1996; Kramer and Lange-Bertalot, 1986, 2004; Lee et al., 2000).

2.3. Variables measured on juvenile crayfish

At the end of the experiment, both survival and body wet weight of all juveniles (precision ± 0.1 mg) were determined, after drying the external surface of animals. After that, the following variables were calculated:

- Specific growth rate (SGR) = [(In Wf − In Wi)/T] × 100, where Wf and Wi are the final and initial body weights, respectively, and T is the time elapsed from the beginning of the experiment.
- Biomass, expressed as g/m²: at the end of the experiment, body weight of all animals belonging to a same experimental group were summed, and then divided by the surface of the three aquaria assigned to each group.

Additionally, 15 juveniles from each group, either early or advanced, were cryo-anaesthetized and the stomach (pro-ventriculus) dissected to determine their content. For this purpose, the stomach content from each dissected juvenile was carefully inspected under binocular microscope (Nikon SMZ 645, 100 to 1000× magnifications), in order to identify the micro-organisms presented. Eventually, samples were photographed under microscope, to further verify the identity of any item. In addition, at the end of the assay both abdominal muscle and hepatopancreas were dissected from both early
(N = 10) and advanced (N = 10) juveniles, to determine the levels of glycogen (according to Van Handel, 1965), lipids (according to Folch et al., 1957; Fring and Dunn, 1970) and total proteins, by the method of Lowry et al. (1951).

2.4. Water quality monitoring

The following variables were weekly monitored throughout the experiment: temperature (precision ± 0.5 °C), pH (Jenco Model 6350 pH meter, precision ± 0.01), dissolved oxygen (Digital Oxygen Meter, precision ± 0.01 mg/L), ammonium (Wiener® kit), nitrates and total hardness (Aquanalitica® kits, for low detection range). Besides, a sample of 20 mL of water culture was taken every 15 d from each aquarium, to determine chlorophyll a concentration, following the same procedure indicated above for biofilm.

2.5. Statistical analysis

Biofilm dry weight, chlorophyll a concentration, as well as water quality parameters were analyzed by a two way-ANOVA, taking experimental group and time of sampling as factors. Lilliefors and Levene tests were used for checking normality and homogeneity of variance, respectively. To analyze survival, SGR, biomass or energetic reserve levels of juveniles at the end of the experiment, a one way-ANOVA was used. Angular transformation was used for survival. When pertinent, Tukey test was used for multiple comparisons of means. Both data normality and homogeneity of variances were always checked. A 5% significance level was always considered (Sokal and Rohlf, 1995).

3. Results

3.1. Biofilm

For both early and advanced juveniles, no significant difference (p > 0.05) was observed in the dry weight of biofilm adhered to plastic sheets, neither between groups B and B+F, nor among times of sampling (Fig. 1). Concerning chlorophyll a concentration, no significant (p > 0.05) differences were found throughout the assay between B and B+F groups, but significant (p < 0.05) differences were seen among sampling time, depending on the kind of juveniles. For early juveniles, a significant (p < 0.05) increase of chlorophyll a concentration was seen at the first week, gradually decreasing toward the end of the experiment (Fig. 1). For advanced juveniles, though, a significant (p < 0.05) decrease occurred from day 0 to day 15 (Fig. 1).

Identification of the micro-organisms presented in the biofilm adhered to plastic nets provided to groups B and B+F is indicated in Table 1, for both early and advanced juveniles. Main micro-organisms identified in most samples were: both filamentous and aggregated cyanobacteria (such as Phormidium sp and Aphanocapsa sp), pennate diatoms (such as Achnanthes exiguum, Nitzschia spp, Navicula spp and Comphomnesia spp), chlorophytes (Oedogonium spp.), xanthophytes (Characiopsis sp), ciliates (such as Vorticella sp, Euplotes sp, Coleps sp), flagellates, rotifers and nematodes. No micro-organisms were found in any sample taken from the control plastic sheets placed on the aquaria of group F.

3.2. Juveniles

3.2.1. Survival and growth

Results of both survival and growth of juveniles are shown in Table 2. For both groups of juveniles, survival was significantly (p < 0.05) higher in groups with biofilm (B and B+F) than in group fed with formulated feed (F), in which survival was relatively low (40% and 60% in early and advanced juveniles, respectively, Table 2). For early juveniles, SGR was significantly (p < 0.05) higher in the groups receiving formulated feed (B+F and F) than in the group only receiving biofilm (B), while no significant differences (p > 0.05) were detected for the advanced juveniles. Biomass showed the highest value in the B+F group, compared to the remaining groups. For early juveniles, significant differences (p < 0.05) were detected between B+F and any of the remaining groups, while for advanced juveniles significant differences (p < 0.05) were only detected between B+F and F (Table 2).

3.2.2. Stomach content

Stomach samples taken from the groups B and B+F showed micro-organism items very similar to those detected in the biofilm provided to the same groups, indicating that both early and advanced juveniles had fed on it (Table 1). However, no nematodes were detected in the stomach of early juveniles, and also in only 33% (N = 5) of those stomachs some rotifers were detected; while both items were found in the stomach of all advanced juveniles. Stomach taken from the F group showed some kind of micro-organisms, such as filamentous cyanobacteria, pennate diatoms, chlorophytes and xanthophytes (Table 1), but in a very lower extent than that observed in groups B and B+F.

3.2.3. Energetic reserves

No significant (p > 0.05) differences among groups of early juveniles were detected in either hepatopancreatic or muscle glycogen...
Table 1

Main micro-organisms detected in both biofilm adhered to artificial substrates and stomachs of either early (EJ) or advanced (AJ) C. quadricarinatus juveniles sacrificed at the end of the 30-d experiment. Experimental groups are indicated as B (only biofilm supplied as food), B+F (both biofilm and formulated diet provided, or F (only formulated food given). (−): absence, (+): low frequency, (+ +): median frequency, (+ + +): high frequency.

<table>
<thead>
<tr>
<th>Division</th>
<th>Family</th>
<th>Gender/specie</th>
<th>Juvenile stage</th>
<th>Presence in biofilm at different sampling days</th>
<th>Presence in stomach</th>
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<td>B 8 15 22 30 B+F 8 15 22 30 F 8 15 22 30</td>
<td>B 30 B+F 30 F 30</td>
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<tr>
<td>Cyanophyta</td>
<td>Oscillatoriaeae</td>
<td>Phormidium sp.</td>
<td>EJ</td>
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(p<0.05) lower than values observed in the remaining groups; while hardness seen by day 15th was significantly (p<0.05) higher in the F group compared to B or B+F group (Figs. 3 and 4).

No significant difference (p>0.05) were observed in the ammonium and nitrite concentrations neither between B and B+F treatments nor among times of sampling, for any kind of juveniles (Figs. 3 and 4), being the corresponding values below the recommended limits for culturing of the studied species (Boyd, 1982; Jones, 1997). However, during the first two weeks the F group of early juveniles showed a significant (p<0.05) increase of both ammonium and nitrite concentrations, with respect to the other two groups, reaching by day 15th values exceeding the recommended limit for culture of the studied species (Fig. 3). Besides, dissolved oxygen showed a significant (p<0.05) decrease in the F group of early juveniles after the first week, together with a marked increase in mortality (near 25%). This fact led us to completely exchange the water of the aquaria at day 22th, to avoid further mortality. After water replacement, levels of ammonium, nitrite and hardness decreased near initial values. However, at the end of the experiment nitrite concentration increased significantly (p<0.05) in the F group, reaching again the value recorded at day 15th (Fig. 3).

For advanced juveniles, the F group showed a significant (p<0.05) increment of ammonium concentration by the first week of the experiment, with respect to the remaining groups, exceeding the recommended limit. By the third week, ammonium levels returned to values similar to the initial one, but at day 30th a new significant (p<0.05) increment was observed (Fig. 4). Nitrite concentration reached its maximum value (p<0.05) at day 15 and up to the end of the assay (Fig. 4).

Finally, the concentration of chlorophyll a in the culture water, for both kinds of juveniles, showed a gradual increment during the experiment. This increment was statistically significant just at the end.
of the experiment, with respect to the initial value, but only for the groups B and B+F (Figs. 3 and 4). The decrease of chlorophyll $a$ seen in the F group of early juveniles was correlated to the water replacement made at day 22th.

4. Discussion

For early juveniles of *C. quadricarinatus*, formulated diet yielded a growth rate (SGR) higher than produced by biofilm as a single diet. However, survival was significantly improved by the presence of biofilm. Moreover, the best values of either SGR or survival were seen in the combination of both formulated food and biofilm as food sources. Correspondingly, the highest value of biomass (an integrative index of both survival and growth) was seen in the later group, at the end of the experiment. Juveniles feeding on both sources would be incorporating a broader spectrum of nutrients, while a high survival would be likely favored by a better water quality associated to the presence of biofilm, as discussed below. Advanced *C. quadricarinatus* juveniles showed a similar trend than early juveniles concerning survival, but growth did not show significant variations among treatments, probably due to the relatively high within-groups variability (standard errors higher than mean by 30%, Table 2).

The only previous report concerning biofilm effects on crayfish species, made on *C. destructor* by Jones and Thanuthong (2002), showed that early juveniles fed on both pellet food and biofilm, grew better than those juveniles only fed with pellets. Shrimp and prawn species, though, have received more attention to this respect. For instance, incorporation of biofilm to the intensive culture of both juveniles and post-larvae of *Litopenaeus vannamei* (Bratvold and Browdy, 2001; Moss and Moss, 2004; Zhang et al., 2010), and also juveniles of the shrimp *Penaeus monodon* (Stuart et al., 2006), improved both weight gain and survival. Biofilm also showed to

**Fig. 3.** Mean values (±SE) of chlorophyll $a$ and several chemical parameters in the water used for culturing early juveniles of *C. quadricarinatus* during 30 d. Experimental groups (B, B+F and F) as defined in Table 1. Asterisks indicate significant ($p<0.05$) differences between F and the other groups. Water in the F group was completely replacement at day 22th (see text).
contribute with several additional nutrients to post-larvae of *Penaeus esculentus*, reared at high densities (Burford et al., 2004). This kind of benefits was also found in other species of high economic value, such as the prawn *Macrobrachium rosenbergii* (Tidwell et al., 1998) and the shrimp *Farfantepenaeus paulensis* (Ballester et al., 2007).

Although total dry weight of biofilm remained constant in all the experimental groups where it was assayed, chlorophyll a concentration decreased throughout the experiment. During the first two weeks of the assay, autotrophic organisms like chlorophytes, xanthophytas, diatoms, flagellates and cyanobacteria were observed, probably contributing to the decrease observed in nitrogenous compounds in water. However, during the second half of the experiment some heterotrophic groups (mainly ciliates, nematodes and rotifers) proliferated, probably grazing on the autotrophic groups and therefore causing the decrease in chlorophyll a observed during this period. According to Da Silva et al. (2009) and Whal (1989), biofilm is formed following a settling pattern, being the relatively high size microorganisms (such as ciliates, nematodes and rotifers) the items colonizing biofilm during its advanced stage of formation. On the other hand, the gradual increment in chlorophyll a observed in the culture water throughout the experiment, mainly in the groups provided with biofilm, could be likely due to the proliferation of autotrophic organisms in the column water from the biofilm already formed, as suggested in previous studies on biofilm systems (Decamp et al., 2002; Moss and Moss, 2004; Thompson et al., 1999).

Both early and advanced juveniles of *C. quadricarinatus* were seen actively grazing on the biofilm formed on the artificial substrates used. From the analysis of stomach content, the main consumed items were chlorophytes, xanthophytas, diatoms, filamentous and aggregated cyanobacteria; only in stomach of some bigger early juveniles rotifers were also seen, but no nematodes were found. Instead, advanced juveniles were able to ingest all the items above mentioned, including nematodes and rotifers, which were found in the stomach of all these juveniles. The development of all the feeding apparatus (such as mandibular appendices and gastric mill), needed for eating and processing relatively big size zooplankton, seems to gradually
complete during the early juvenile instars of C. quadricarinatus (Loya-
Javellana et al., 1994). Some studies found that highest growth rate of 
F. paulensis juveniles was related to the consumption of biofilm colon-
ized by nematodes, a substantial source of nutrients for shrimps 
(Ballester et al., 2007; Pissetti, 2005). The presence of certain autotro-
phic organisms in the stomach of some juveniles of the F group could 
be the consequence of the proliferation of such micro-organisms in 
water, probably favored by the accumulation of organic matter due 
to dead animals or exuviae, as seen in previous studies (Decamp 
et al., 2002; Moss and Moss, 2004).

Total lipid levels, determined at the end of the assay, showed 
a higher accumulation of lipid in the hepatopancreas of both early 
and advanced juveniles fed on formulated diet, alone or in combina-
tion with biofilm. In addition, a higher glycogen level was observed 
in advanced juveniles fed with that food. The hepatopancreas of crust-
aceans is in fact the main storage organ for both lipids and carbohy-
drates (Hernández-Vergara et al., 2003; Verri et al., 2001). The 
relatively high caloric value of the commercial food employed 
(254 kcal/100 g, Chaulet et al., 2008) could be therefore explaining 
the higher caloric reserve levels seen in the hepatopancreas of juve-
niles fed with formulated food. Nevertheless, these levels of energetic 
reserve (expressed as mg/g of tissue) did not show a clear correlation 
with the growth index calculated (SGR): as mentioned before, the 
maximum value of SGR, for both kinds of juveniles, was seen in the 
B + F group, while maximum values of lipid reserves were found in 
juveniles only fed with formulated food (F group). These results are 
suggesting that factors other than the level of energetic reserves 
(such as vitamins, prebiotics and other micronutrients) are involved 
in growing of juveniles, and also that such factors are related to bio-
film consumption. Consumption of plant material has been previously 
suggested to enhance growth of C. quadricarinatus juveniles (Jones, 
1997). Interestingly, protein levels in both hepatopancreas and mus-
cle were similar in all groups of juveniles, indicating that no serious 
nutritional deficit seemed to be occurred in such juveniles that only 
had biofilm as food source.

Several studies reported high protein level for biofilm micro- 
organisms, i.e., ranging from 20 to 40% (dry weight basis) in diatoms 
(Martinez-Fernández et al., 2006; Renaud et al., 1999), to more than 
50% (dry weight basis) in nematodes (Schlechtriem et al., 2004). Be-
sides, several micro-organisms of biofilm present a high proportion 
of amino-acids, vitamins, carotenoids, esteroids and polysaturated 
fatty acids, these latter intensively produced by cyanobacteria, flagel-
lates, ciliates and nematodes (Meyers and Latscha, 1997; Schlechtriem 
et al., 2004; Zhukova and Kharlamenko, 1999). Burford et al. (2004), 
by using stable isotopes, have reported that both carbon and nitrogen 
coming from micro-organisms of biofilm were assimilated by shrimp 
post-larvae under intensive culture, contributing in a great ex-
tent (39–53%) to their nutritional requirements. With the same tech-
nique, Abreu et al. (2007) observed that micro-organisms of biofilm 
can contribute with 49% of carbon and 70% of the nitrogenous needed 
for growing of F. paulensis early juveniles. Avnimelech (2000) deter-
mined that micro-organisms presented in culture tanks increased 
the efficiency to protein conversion around 45%, by incorporating in-
organic nitrogenous dissolved in water to microbial protein, which is 
farther ingested by cultured animals, therefore reducing the use of 
formulated food.

A clear benefit of biofilm to both survival and growth of cultured 
species is related to water quality. Several bacteria and microalgae 
presented in biofilm are able to actively uptake ammonium and other 
nitrogenous waste, to synthesize protein that can be consumed by 
either fish or crustaceans, therefore contributing to both water 
quality maintenance and nutrition of cultured animals (Azim et al., 
2001; Bratvold and Browdy, 2001; Ramesh et al., 1999). Other studies 
have shown that nitrogenous uptake by biofilm micro-organisms can 
help reduce the proliferation of pathogen bacteria in cultures (Alabi 
et al., 1999; Thompson et al., 2002). In the current study, both 
ammonium and nitrite concentrations were maintained at a very 
low and constant level in the aquaria where biofilm was added, 
while those aquaria only receiving formulated food showed increased 
levels of both toxic metabolites, frequently above the permissible 
levels for aquaculture. These relatively high levels were associated, 
especially for early juveniles, to an increased mortality. Therefore, 
observed results strongly suggest that absorption and/or transforma-
tion of nitrogenous waste compounds by biofilm micro-organisms 
greatly improve the survival of juveniles of the studied species.

According to Thompson et al. (2002) biofilm developed on the 
wall of shrimp culture tanks was able to maintain the water concen-
tration of both ammonium and phosphorous at low levels, especially 
once chlorophyll a reaches values around 5 μg/cm²; this chlorophyll 
mainly corresponding to both pennate diatoms and cyanobacteria 
presented in biofilm. For both kinds of C. quadricarinatus juveniles, 
chlorophyll a in biofilm ranged from 4 to 10 μg/cm², in correlation 
with low levels of both ammonium and nitrite in water. On the con-
trary, in those aquaria with no biofilm added, high and fluctuating 
levels of nitrogenous waste were verified. These later aquaria, com-
pared to those provided with biofilm, also showed lower values of 
both pH and dissolved oxygen, as expected from a weak development 
of an autotrophic community helping to regulate such critical vari-
ables. In this sense, the gradual increment in chlorophyll a levels 
seen in the water column of group F could indicate the presence of 
some autotrophic micro-organisms in such compartment, but their 
contribution to the control of nitrogenous waste was not certainly 
so efficient as that of micro-organisms associated to the biofilm de-
grown in groups B and F + H. However, both pH and dissolved oxy-
gen in F groups showed moderate fluctuations, in all cases below the 
recommended limits for the studied species (Jones, 1997; Masser and 
Rouse, 1997).

5. Conclusion

We conclude that the addition of biofilm to culture of C. quadricar-
inatus juveniles represents a clear advantage for both survival and 
growth of animals. Biofilm is clearly involved in the maintenance 
of water quality (essential for a good survival), and perhaps also as 
complementary food source, therefore improving the growth of juveniles, 
especially early ones. By using biofilm in a zero water exchange sys-
tem, a significant reduction in the cost of production could also be 
atained.

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