Short-term feeding response of the mussel *Mytilus chilensis* exposed to diets containing the toxic dinoflagellate *Alexandrium catenella*

Respuesta alimentaria inicial del bivalvo *Mytilus chilensis* expuesto a dietas conteniendo el dinoflagelado tóxico *Alexandrium catenella*

JORGE M. NAVARRO*, ANDREA M. CONTRERAS¹ & ÓSCAR R. CHAPARRO

Instituto de Biología Marina “Dr. Jürgen Winter”, Universidad Austral de Chile, Casilla 567, Valdivia, Chile

¹ Present address: School of Biological Sciences, University of Canterbury, Private Bag 4800, Christchurch, New Zealand; *e-mail for correspondence: jnavarro@uach.cl

ABSTRACT

The short-term feeding response of the bivalve *Mytilus chilensis* was measured using four diets containing different proportions of the toxic dinoflagellate *Alexandrium catenella*. The diets containing the highest concentrations of the dinoflagellate showed the greatest effect on the feeding activity in the mussel, with clearance and ingestion rates significantly reduced during the first hours of exposure. After this period, *M. chilensis* demonstrated a capacity to acclimate to the toxic diets, with feeding parameters reaching values similar to those of untreated control organisms. It was not clear if the negative effect on the feeding behavior was caused by the presence of the paralytic toxin, or due to the larger size of the dinoflagellate cells in comparison with cells of *Isochrysis galbana* used in the control diet. However, parallel studies with diets containing the nontoxic dinoflagellate *Alexandrium affine* of similar size and shape to that of *A. catenella*, suggested the cell size was the main cause for impairment of feeding behavior. The capacity for acclimation to either toxin or cell size by *M. chilensis* makes it a good indicator species for the early detection of harmful PSP events, since its relative insensitivity to the toxin allows it to quickly recover normal feeding behavior and permits it to accumulate PSP in its tissues in a short time.

Key words: *Mytilus chilensis*, feeding response, PSP, dinoflagellates.

RESUMEN

La respuesta inicial del bivalvo *Mytilus chilensis* fue medida bajo cuatro dietas que contenían diferentes proporciones del dinoflagelado tóxico *Alexandrium catenella*. Las dietas que contenían las concentraciones más altas de este dinoflagelado mostraron el mayor efecto durante las primeras horas de exposición. Después de este periodo inicial, *M. chilensis* demostró la capacidad para aclimatarse a estas dietas tóxicas, con parámetros de alimentación que alcanzaron valores similares a aquellos de los organismos controles. No fue claro si el efecto negativo sobre la conducta de alimentación fue causado por la presencia de la toxina paralizante o debido al gran tamaño de las células del dinoflagelado, en comparación al tamaño de las células de *Isochrysis galbana* usadas en la dieta control. Sin embargo, estudios paralelos con dietas conteniendo el dinoflagelado no tóxico *Alexandrium affine*, de similar tamaño y forma a *A. catenella*, sugieren que el tamaño de las células fue la causa principal que afectó la conducta de alimentación. La capacidad para aclimatación ya sea a la toxina o al tamaño celular del dinoflagelado, identifica a *M. chilensis* como una buena especie para la detección temprana de eventos tóxicos producidos por *A. catenella*, ya que debido a su relativa insensibilidad a la toxina permite una rápida recuperación de su conducta de alimentación y la acumulación de toxina en sus tejidos.

Palabras clave: *Mytilus chilensis*, respuesta de alimentación, toxina paralizante, dinoflagelados.

INTRODUCTION

The presence of paralytic shellfish poisoning (PSP) produced by dinoflagellate species in the natural environment has been described as a factor causing a variety of sub lethal effects in marine bivalves. Symptoms identified include closure of the valves, reduction in filtration activity, drops in metabolic rates, increase in mucus production, and erratic cardiac activity (Shumway et al. 1985). One of the most common effects during exposure to paralytic...
toxin is isolation from the immediate environment by firm closure of the valves which implies cessation of filtering activity. Thus for example when *Mytilus edulis* was exposed to the toxic dinoflagellate *Protogonyaulax tamarense*, it demonstrated erratic closure of the valves, while *Mercenaria mercenaria* showed complete closure of the valves (Gainey & Shumway 1988). Reduction of filtering activity due to valve or siphon closure has also been reported for *M. edulis* and *Mya arenaria* (Shumway & Cucci 1987), *M. mercenaria* (Dupuy & Sparks 1968) and *Crassostrea virginica* (Ray & Aldrich 1967, Shumway & Cucci 1987), *M. edulis* and *Mya arenaria* (Shumway & Cucci 1987), *M. mercenaria* (Dupuy & Sparks 1968) and *Crassostrea virginica* (Ray & Aldrich 1967, Shumway & Cucci 1987). Wildish et al. (1998) studying the effects of toxic and non-toxic strains of *Alexandrium tamarense* on the initial feeding response of the oyster *Crassostrea gigas* found a start/stop feeding behaviour with both strains of the dinoflagellate, in contrast with oysters fed with the Isochrysis sp. used as a reference species and with which the oysters did not exhibit this behaviour.

Chile has large populations of filter feeding bivalve species inhabiting its southern regions which are susceptible to contamination by toxins produced by blooms of *A. catenella*. During the last decade, these harmful algae blooms (HABs) have increased in frequency, intensity, and area affected including the coastal zone from Castro (42°29' S, 73°48' W, Chiloé Island) to south of the Beagle Channel (55°07' S, 68°36' W). From 1991 to the present, several cases of red tides formed by *A. catenella* occurred in southern Chile, with the largest in the summer-fall of 2002 having maximum concentrations of 7.8 x 10^5 cells L^-1 (Clement et al. 2002) and toxicity values of above 8500 μg of STX/100 g bivalve tissue. It is therefore relevant to evaluate the effect of diets containing PSP-producing dinoflagellates on the filtration activity and the subsequent level of toxicity acquired by *M. chilensis*.

**MATERIAL AND METHODS**

*Collection site and experimental mussels*

Adult individuals of *M. chilensis* were collected in August 2004, from culture ropes at the Yaldad Experimental Station in southern Chile (43°08' S, 73°44' W). Individuals of about the same lengths were chosen (5.38 ± 0.16 cm), having dry tissue weights of about 0.75 ± 0.20 g (termed “standard size” in the present study). The mussels were transported to the laboratory, where they were acclimated in four tanks of 20 L volume (25 mussels per tank) for one week at a temperature of 14 ºC and seawater salinity of 30 ‰. During this period they were fed “ad libitum” with the microalga *Isochrysis galbana* and provided with constant aeration from compressed air bubblers. Seawater was changed every 48 h.

*Preparation of diets*

Monoclonal non-axenic *Alexandrium catenella* (strain ACC02) was employed as food in a series of feeding experiments. This strain was isolated in Chile’s Aysen Region in 1994 and was kindly donated by the Centro Regional de Análisis de Recursos y Medio Ambiente (CERAM) of the Universidad Austral de Chile. *Alexandrium catenella* was cultivated in 0.45 μm filtered seawater enriched with “L1” algae culture medium (Guillard 1995), at 14 ºC and 30 ‰ S. *Isochrysis galbana* was cultivated using f/2 medium (Guillard 1975). The experiments were carried out using both species of algae harvested in their exponential growth phase.

To emulate the organic fraction of the natural suspended particulate matter, sediment was used for preparation of the diets. It was collected from the upper centimeter of the substrate on the Yaldad tidal flat, sieved to 40 μm, rinsed with distilled water, and ashed in a muffle furnace at 500 ºC for 12 h to eliminate the organic fraction.
After ashing, the sediment was again sieved to eliminate sediment aggregations.

Experimental diets (1.7-2.0 mg·L⁻¹; dry weight) were prepared, with different dry weight proportions of *A. catenella*, *I. galbana* and sediment, in order to provide a range of diets containing different levels of toxicity as listed in Table 1. The control diets were prepared by mixing *I. galbana* with sediment or with pure *I. galbana*. The experiments were carried out consecutively with each diet, each time using seven 15 L experimental aquaria containing numbers of mussels in accordance with the duration of each experiment (two mussels per measurement). Four aquaria contained mussels fed with the diet containing *A. catenella*, and the other three with mussels fed the control diet. Each day each aquarium received an amount of food representing 3% (23 mg day⁻¹) of the dry body weight of the experimental mussels, delivered continuously using a Masterflex 7519-05 peristaltic pump at the temperature and salinity cited above. The concentration of saxitoxin in each diet was calculated on the basis of the results of Navarro et al. (2006), who found a mean concentration of 3.84 ± 1.81 fmole cell⁻¹ in the same strain of *A. catenella* (ACC02) and under the same culture conditions.

**Toxin analyses in soft tissues**

The toxin content in the soft tissues of *M. chilensis* was calculated by pooling the tissues of two individuals (5 g approx.) from each experimental aquarium on days 0, 1, 2, and 3 of the experiment. The same procedure was carried out on individuals from control aquaria to verify complete absence of toxin. Toxin content of the tissues was quantified using the electrophysiological test of Vélez et al. (2001), where HEK 293 cells (human embryonic kidney cells) expressing STX-sensitive rat skeletal muscle Na channels were patch clamped in the whole-cell configuration. The equivalent STX concentration was estimated using calibration curves obtained by external perfusion with known concentrations of purified STX; these curves were generated using a stepped series of increasing concentrations of STX-dihydrochloride (US Food and Drug Administration, Office of Seafood). According to Velez et al. (2001), there is a correlation of 0.96 between the mouse bioassay and the electrophysiological test.

**Clearance rate**

The clearance rate (CR) was estimated using mussels from the seven experimental aquaria twice a day, until the test mussel showed a feeding response which did not differ to the test mussels, over periods of 0, 6, 24 30, 48, 54, and 72 h. The clearance rate was determined with the four experimental diets, using a particle concentration that varied between 1.7 and 2.0 mg L⁻¹ dry weight, temperature of 14º C and salinity of 30 ‰. The clearance rate experiments were carried out in a static system that was homogenized by aeration, in which the decrease in particle concentration was periodically monitored in each experimental aquarium. This was carried out using an Elzone.

**Table 1**

<table>
<thead>
<tr>
<th>Diet condition</th>
<th>Total diet (mg d⁻¹)</th>
<th>Organic matter (mg d⁻¹)</th>
<th>A. catenella (mg d⁻¹)</th>
<th>I. galbana (mg d⁻¹)</th>
<th>Sediment (mg d⁻¹)</th>
<th>A. catenella (cells d⁻¹)</th>
<th>Toxicity (pmol d⁻¹)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Diet 1 (10 % A. catenella)</td>
<td>23</td>
<td>9.5 (41.4 %)</td>
<td>2.3</td>
<td>6.9</td>
<td>13.8</td>
<td>489,362</td>
<td>1,879</td>
</tr>
<tr>
<td>Diet 2 (30 % A. catenella)</td>
<td>23</td>
<td>8.3 (36.3 %)</td>
<td>6.9</td>
<td>2.3</td>
<td>13.8</td>
<td>1,468,085</td>
<td>5,637</td>
</tr>
<tr>
<td>Diet 3 (50 % A. catenella)</td>
<td>23</td>
<td>12.4 (54.0 %)</td>
<td>11.5</td>
<td>2.3</td>
<td>9.2</td>
<td>2,446,809</td>
<td>9,396</td>
</tr>
<tr>
<td>Diet 4 (100 % A. catenella)</td>
<td>23</td>
<td>20.5 (89.0 %)</td>
<td>23.0</td>
<td>0.0</td>
<td>0.0</td>
<td>4,893,617</td>
<td>18,792</td>
</tr>
<tr>
<td>Control (Diets 1-2-3)</td>
<td>23</td>
<td>10.3 (44.7 %)</td>
<td>0.0</td>
<td>10.8</td>
<td>12.2</td>
<td>0.0</td>
<td>0.0</td>
</tr>
<tr>
<td>Control (Diet 4)</td>
<td>23</td>
<td>20.9 (90.7 %)</td>
<td>0.0</td>
<td>23</td>
<td>0.0</td>
<td>0.0</td>
<td>0.0</td>
</tr>
</tbody>
</table>
model 180XY particle counter equipped with a counting tube having a 120 μm aperture. These tests were carried out over a period of three hours, with measurements made every 30 minutes, and in every case replacing the food consumed with new food. To test any sedimentation of the cells during the feeding measurements, a control aquarium without the presence of mussels was run. The clearance rate (L/h) was calculated following Coughlan (1969).

Ingestion rate

A known volume of each experimental diet (in triplicate) was concentrated on 47 mm diameter Whatman GF/C glass fiber filters, which had been previously washed, ashed and tared. Blank filters as well as filters with retained diets were rinsed with isotonic ammonium formate to remove salt and prevent cell lysis. The filters were dried at 80 °C for 24 h, weighed, ashed at 450 °C for 3 h and reweighed after cooling in a desiccator, to calculate total and organic content of the diets. The concentration of food used (1.7-2.0 mg L-1) was below the threshold of pseudofeces production (Velasco & Navarro 2002) so that the ingestion rate could be calculated as the product of the clearance rate and the amount of material contained in each experimental diet.

Statistical analyses

The response of *M. chilensis* to the experimental diets was analyzed using STATISTICA 4.2 software. The means were compared using one way analysis of variance (ANOVA), followed by a “post-hoc” Tukey test, using a log(x) transformation of the data when these did not comply with a posteriori assumptions (normality and homogeneity of the variance). The relation between clearance rate and *A. catenella* cell concentration was carried out using regression analysis.

RESULTS

The *M. chilensis* in the control group showed a clearance rate (CR) which was stable over time. The mussels of the toxin-contaminated group demonstrated erratic behavior (Fig. 1A) when given the diet containing the lowest proportion of *A. catenella* (10 %). More stable behavior was observed in mussels exposed to diets containing 30 and 50 % *A. catenella*. In these, an initial feeding reduction on the diet containing the toxic dinoflagellate was observed (Fig. 1B and 1C), followed by a period of acclimation to the diet, and then reaching the values of the controls. With the diet consisting only of *A. catenella*, the *M. chilensis* clearance rate became significantly lower (P < 0.05) than the controls during the first 54 h. The mussels acclimated to this diet by 72 h, again reached the rate of the control group (Fig. 1D). Thus the effect of *A. catenella* on lowering the feeding rate of *M. chilensis* was more intense and more extended in time as the proportion of *A. catenella* increased in the diet, which represented an inverse relation between the capacity of *M. chilensis* to acclimate, and the concentration of *A. catenella* in the diet. The highest average clearance rate was observed in the control mussels (hour 0), while the lowest average clearance rate, 0.31 L h⁻¹, was observed in experimental individuals fed (toxic) diet 4 at the beginning of the experiment (hour 0).

The relationship between clearance rate and the number of cells of *A. catenella* supplied in the diets (Fig. 2A and 2B), indicated a significant negative correlation (P < 0.05) at hours 0 and 24 (R² = 0.75 and 0.96, respectively). In the sampling at 54 h. there was no significant relationship (P > 0.05) between these variables (Fig. 2C). Variation in ingestion rate (IR) follows exactly the same trends as those described previously for clearance rate (CR).

The toxic content in the tissues of the individuals fed during 72 h with diets 1 and 2 were not significantly different from each other, and remained below the safety limit for human consumption (80 μg STX eq 100 g tissue⁻¹). In contrast, the mussels fed with diets 3 and 4 reached toxin levels significantly higher than with the other two diets, and showed values above the safety limit for human consumption (diet 3 = 236.8 μg STX eq 100 g tissue⁻¹; diet 4= 314.5 μg STX eq 100 g tissue⁻¹), with values significantly different than those obtained with diets 1 and 2, but not significantly different between each other (Fig. 3).
Fig. 1: Clearance rate of *M. chilensis* (standard animal) fed with diets containing different concentrations of *A. catenella* (see Table 1 for diet composition). Values are means ± standard error.

Tasa de aclaramiento de *Mytilus chilensis* (animal estándar) alimentado con dietas conteniendo diferentes concentraciones de *A. catenella* (ver Tabla 1 para composición de las dietas). Valores corresponden a medias ± error estándar.
Fig. 2: Clearance rate of *M. chilensis* (standard animal) in relation to cell number of *A. catenella* supplied in the diets. Values are means ± standard error.

Tasa de aclaramiento de *M. chilensis* (animal estándar) en relación al número de células de *A. catenella* entregado en las dietas. Valores corresponden a medias ± error estándar.
The genus *Mytilus* has been described as generally insensitive to dinoflagellates containing paralytic shellfish toxin (Bricelj & Shumway 1998), experiencing only a small reduction in their filtering rates although high levels of toxins accumulate in their tissues during short periods of exposure. Contrasting behavior has been described for other filter feeding bivalves such as the oyster *Crassostrea gigas* (Bardouil et al. 1993) and the clam *Mya arenaria* (Bricelj et al. 2005). Filtration activity of these species is sensitive to PSP toxins; hence they do not accumulate high concentrations of toxins, presumably due to behavioral and physiological mechanisms used to avoid exposure to the toxic dinoflagellate cells (Gainey & Shumway 1988).

Similarly to the results obtained with diet 1 for *M. chilensis*, Li et al. (2001, 2002) observed no negative effects of diets containing small amounts of paralytic toxin on clearance rates of the mussel *Perna viridis*. Bardouil et al. (1996) showed that the inclusion of 10% toxic *A. tamarense* in a diet produced a 20% reduction in the clearance rate of *Crassostrea gigas*. The initial negative effect of our diets 2, 3 and 4 on the clearance and ingestion rates of *M. chilensis*, was similar to the behavior described by Wildish et al. (1998). These authors described a start/stop behavior in the clearance rate of the oyster *C. gigas* when fed toxic and nontoxic strains of *Alexandrium tamarense*, suggesting that periods of over 48 h were required to achieve acclimation to these diets. Bardouil et al. (1996) demonstrated a complete inhibition of filtering activity in *C. gigas* in the first six hours of experimentation when using a diet composed solely of toxic *A. tamarense*. Dupuy & Sparks (1968) mentioned that a period of two weeks was required for *C. gigas* to return to normal feeding on *A. catenella* which was greater than the time observed in the present study with the mussel *M. chilensis*. This supports the contention that the mytilids have a high capacity for acclimation to diets containing toxic dinoflagellates (Lesser & Shumway 1993, Bricelj & Shumway 1998).

**Fig. 3:** STX equivalent of *M. chilensis* (standard animal) exposed to diets containing different concentrations of *A. catenella*. For diet 4 (days 1 and 2) there is no information. Values are means ± standard error.

STX equivalente en *M. chilensis* (animal estándar) expuesto a dietas conteniendo diferentes concentraciones de *A. catenella*. Para dieta 4 (días 1 y 2) no existe información. Valores corresponden a medias ± error estándar.
A number of authors have related the reduction of filtration activity to the sensitivity of bivalves to the paralytic toxin (Twarog & Yamaguchi 1974, Shumway & Cucci 1987). The capacity of acclimation in *M. chilensis* may also be related to the origin of the population studied (Yaldad Bay, Chiloé), which was exposed in 2002 to one of the largest blooms of *A. catenella* ever occurring in Chile, when concentrations reached 7.8 x 10^5 cells L^-1 (Clement et al. 2002), accompanied by toxin levels of 8554 ug STX eq 100 g *M. chilensis* tissue^-1 (Llanchipal Health Service). Twarog et al. (1972) compared resistance to STX of various bivalve species, measuring the blocking of action potentials of the sodium channels on a cellular level. These authors concluded that *Mytilus edulis* was the most resistant species to the toxin, noting that previous exposure of the individuals to toxic algal blooms was related to differences in the responses measured. An example of this was the study by Shumway & Cucci (1987) who suggested that individuals of *M. edulis* from areas which had never experienced toxic blooms were more sensitive to PSP than individuals originating in areas which undergone repeated toxic algal blooms. These results coincide with those described by Bricelj et al. (2005), where individuals of the clam *Mya arenaria* exposed to recurrent toxic events experienced a natural mutation of an amino acid which prevented the union of the saxitoxin with the pore of the sodium channel, thus permitting normal conduction of nervous impulses of the cells and conferring greater resistance to the toxin in these individuals. Nevertheless, in the case of *M. chilensis* this behavior seems to be more related to the effect on the feeding mechanism of the bivalve in relation to the larger size of the dinoflagellate cell in comparison with that of feeding on Isochrysis (*A. catenella* = 32-36 μm, *I. galbana* = 4-5 μm) than to the toxicity of the dinoflagellate. Indeed, preliminary results of a parallel study carried out to evaluate the capacity for pre-ingestive selection by *M. chilensis* exposed to a mixed diet (1:1 by weight) of *I. galbana* and the non-toxic dinoflagellate *Alexandrium affine* (which has a similar size and shape to *A. catenella*) also showed a significant initial reduction in the clearance rate compared to controls only fed with Isochrysis. This response of *M. chilensis* could be also related with the stimuli produced by chemical cues from the dinoflagellate species, as has been suggested by Wildish et al. (1998), who concluded that PSP was not the cause of the initial inhibition of the feeding response since both toxic and nontoxic strains of *A. tamarense* caused start/stop feeding in the oyster *C. gigas*.

This insensitivity of *M. chilensis* to PSP allows it to accumulate toxin in its tissues in a short period of time, suggesting it would be valuable as an indicator species for early detection of harmful algal blooms. The increasing frequency of blooms of *A. catenella* in southern Chile make it essential to continue studies which help define the responses of *M. chilensis* to these blooms in the long term. This is of importance considering the ecological relevance of the species as well as its commercial export value based on the high volume of these mussels produced in artificial cultures which are undergoing continued expansion in southern Chile.

ACKNOWLEDGEMENTS

We especially thank G. Urrutia and M. Maturana for their valuable help during the experiments, to M. Seguel for providing the *A. catenella* strain (ACCO2). We also thank Marco Cordova from the laboratory of Marine Toxins from Universidad de Chile for performing electro-physiological analysis of *Alexandrium catenella* cells. This study was supported by a research grant to JMN (FONDECYT 1030340).

LITERATURE CITED


BRICELJ VM, L CONNELL, K KONOKI, SP MACQUARRIE, T SCHEUER, WA CATTERALL & VL TRAINER (2005) Sodium channel mutation
leading to saxitoxin resistance in clams increases risk of PSP. Nature 434: 763-767.


