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Feeding preference of the native sea urchin Arbacia dufresnii in an environment invaded by the kelp Undaria pinnatifida

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Abstract: In Argentina, the invasive kelp Undaria pinnatifida has modified the benthic community of Nuevo Gulf, northern Patagonia, where it was first recorded and is currently showing advanced invasion. Arbacia dufresnii, an abundant generalist omnivore sea urchin, has fully incorporated U. pinnatifida into its diet. Since sea urchins can influence the structure and dynamics of benthic macroalgal assemblages, determining the importance of U. pinnatifida in the feeding preferences of A. dufresnii complements the evaluation of the modifications caused by this species. Thus, this study aimed to assess the feeding preference of A. dufresnii in an environment invaded by U. pinnatifida. We conducted feeding experiments to determine whether A. dufresnii feeds on different developmental stages of U. pinnatifida, and preference experiments of this invasive species against six macroalgal species abundant in the environment. Arbacia dufresnii consumed both blades and sporophylls of U. pinnatifida at all the developmental stages evaluated, showing a preference for blades over sporophylls. Arbacia dufresnii showed preference for U. pinnatifida only when it was offered together with species that possess defense mechanisms against herbivory. The lack of preference for other macroalgae species over U. pinnatifida suggests that this invasive species may alter resource use and thereby affect the trophic structure of native communities. These findings complement previous diet studies, corroborating the feeding plasticity of A. dufresnii, as well as its ability to adapt to newly available resources. This research enhances the understanding of the impacts caused by invasive macroalgal species through feeding ecology studies of native grazers.

Key words: Food preference, echinoid, invasive kelp, Patagonia

Resumen: Preferencia alimentaria del erizo de mar Arbacia dufresnii en presencia del alga invasora Undaria pinnatifida. En Patagonia Argentina, el alga invasora Undaria pinnatifida fue registrada por primera vez en el Golfo Nuevo, donde ha modificado la comunidad bentónica, presentando actualmente un grado de invasión avanzada. El equinoideo Arbacia dufresnii es un omnívoro generalista que ha incorporado completamente el alga invasora a su dieta. Dado que los equinoideos pueden influir en la estructura y dinámica de los ensambles de macroalgas bentónicas, determinar la importancia de U. pinnatifida en las preferencias alimentarias del equinoideo complementa la evaluación de las modificaciones causadas por esta especie. Con este fin se realizaron experimentos de única opción para determinar si A. dufresnii se alimenta de diferentes estadios de U. pinnatifida, y experimentos de preferencia de esta especie invasora frente a seis especies de macroalgas. El equinoideo consumió los diferentes estadios de U. pinnatifida, tanto el esporofilo como de la fronde, mostrando preferencia por esta última. Arbacia dufresnii mostró preferencia por U. pinnatifida únicamente cuando fue ofrecida junto a especies que poseen defensas contra la herbivoría. La falta de preferencia por las otras especies de macroalgas sugiere que U. pinnatifida alteraría el uso de los recursos afectando la estructura trófica de las comunidades nativas. Estos hallazgos complementan los estudios existentes de la relación U. pinnatifida - A. dufresnii, corroborando la plasticidad alimentaria del equinoideo así como su capacidad para adaptarse a los nuevos recursos disponibles. Esta investigación aporta comprensión a los impactos causados por U. pinnatifida mediante estudios de ecología alimentaria de los herbívoros nativos

Palabras clave: Preferencia alimenticia, equinoideo, algas invasoras, Patagonia

INTRODUCTION

Invasive species are globally recognized as one of the five major direct drivers of environmental change, alongside land and sea use change, direct exploitation of organisms, climate change, and pollution (Roy et al., 2023). The magnitude and types of impacts of the invasion vary among different invasive species and across ecosystems and regions. In the marine environment, the predominant ecological impacts of invasive macroalgae are space monopolization and altered community composition (Davidson et al., 2015). Specifically, the macroalga Undaria pinnatifida (hereafter referred to as Undaria), native to northeast Asia (Akiyama & Kurogi, 1982), is in the top 10 most widespread invasive algae species worldwide (Kulhanek et al., 2011; Seebens et al., 2023). In Argentina, this species was first recorded in Nuevo Gulf, northern Patagonia, in 1992 (Casas & Piriz, 1996). Since its introduction, it has spread over most of the Argentine coast, with the densest kelp beds in northern Patagonia, with densities occasionally higher than 4 kg·m⁻² (Dellatorre *et al.*, 2014). Undaria is an annual species with a heteromorphic life cycle typical of the Laminariales, with alternating gametophytic (microscopic) and sporophytic (macroscopic) generations (Choi et al., 2007). Prior to the Undaria invasion, macroalgal communities in Nuevo gulf were characterized by small species (Ulva sp., Codium sp., Dictyota dichotoma, among others), which did not exceed 50 cm in height (Boraso & Zaixso, 2008). Because of its invasive features, the presence of Undaria has modified the benthic community of Nuevo Gulf (Orensanz et al., 2002; Raffo, 2014), causing a decrease in the populations of native macroalgae (Casas et al., 2004), an increase of 800% in biomass availability for herbivores (Casas et al., 2008; Raffo et al., 2015), and an increase in the populations of some invertebrates. For example, in the presence of Undaria, the abundance of the sea urchin Arbacia dufresnii has increased three times (Irigoyen et al., 2011).

Arbacia dufresnii is the most common sea urchin in the northern Patagonian coast of Argentina (Zaixso & Lizarralde, 2000; Brogger et al., 2013). Depending on the availability of food in the environment, this species can be considered as a herbivore, carnivore, or generalist omnivore (Vásquez et al., 1984; Penchaszadeh & Lawrence, 1999; Newcombe et al., 2012; Zárate, 2014; Castro et al., 2022). Particularly, in Nuevo Gulf, Undaria is consumed and assimilated in the tissues of A. dufresnii, suggesting a complete incorporation of the invasive kelp into the sea urchin diet (Castro et al., 2022). However, questions such as whether the sea urchin prefers to feed on Undaria or other macroalgae available in the environment have not yet been resolved. Considering that sea urchins can impact the structure and dynamics of benthic macroalgal assemblages in coastal habitats (Scheibling & Anthony, 2001; Valentine & Johnson, 2003), determining the relevance of Undaria in their feeding preferences may contribute to evaluating the magnitude of the impacts caused by the invasive species.

Feeding preference occurs when a consumer actively selects prey based on specific behaviors (Singer, 2000). This behavior is typically assessed by comparing the consumption of a prey item when presented alone versus when presented alongside another prey item (Underwood & Clarke, 2005). These studies are often useful to evaluate the role of grazers such as sea urchins in structuring benthic communities, but also to understand the interactions between consumers and prey (Jackson & Underwood, 2007; Aguilera, 2011). However, feeding preference studies performed in the field are challenging due to multiple variables acting simultaneously, making any interpretation uncertain. The aim of this study was to assess the feeding preference of the native sea urchin Arbacia dufresnii from an environment invaded by Undaria pinnatifida. To achieve this, we performed feeding experiments at controlled aquarium conditions, using Undaria at different developmental stages to determine whether the sea urchin can graze on the invasive kelp throughout the year. Additionally, we evaluated the feeding preferences of A. dufresnii comparing Undaria with the six most abundant macroalgae species available in the environment.

METHODOLOGY

Adults of A. dufresnii (N =112 in all experiments) were randomly collected by scuba diving from Punta Cuevas, Nuevo Gulf ($42^{\circ}46'44''S$; $64^{\circ}59'52''W$) in different seasons, according to the developmental stages of the Undaria life cycle reported by Casas *et al.* (2008) for the study area: i) winter (August 2014), when the sporophytes were starting to grow, Undaria formed dense stands and the mean densities were maximum (*growth stage*); ii) spring (November 2014), when sporophylls were mature, showing

the highest average of biomass (maturity stage); and iii) summer (January 2015), when the kelp was senescent (senescence stage). In winter, we also collected entire Undaria individuals and the most frequent and abundant macroalgal species, such as Dictyota dichotoma, Lomentaria clavellosa, Codium vermilara, Ceramium sp., Ulva spp., and a Corallinales (rhodolith) species. After collection, sea urchins were transported to the Experimental Aquarium of the Centro Nacional Patagónico (CCT CONICET CENPAT), Puerto Madryn, Chubut, Argentina, where they were maintained on a 12 h light:12 h dark photoperiod with a salinity constant with a mean value of 34 and temperature between 14 and 16 °C. Water quality was maintained within optimal parameters. Undaria and the other macroalgae were manually cleaned to remove visible epiphytes or animals and separately preserved in containers with flowing seawater.

Two experiments were carried out. One experiment was aimed to evaluate the consumption of the different structures of Undaria (blades and sporophylls) at the different developmental stages (growth, maturity and senescence) (Undaria experiment). The other experiment was aimed to determine the feeding preference of sea urchins between Undaria and the other macroalgae species available in the environment (feeding preference experiment). Experiments were carried out in 90-L containers with water recirculation, biofilter and air pump to ensure good water oxygenation, for three days. Each container was subdivided into eight equal compartments (15x15x15 cm) using a plastic screen of 1 mm of mesh for water circulation. Prior to the start of the experiments, sea urchins were starved for three days to promote uniform nutritional state. Each individual was measured (test diameter, caliper accuracy of 0.1 mm), and

a one-way ANOVA (α =0.05) was performed to corroborate the homogeneity of individual sizes across treatments and trials. The mean (± SD) diameter of the sea urchin was 28.35 mm (± 3.8 mm).

Undaria experiment

To evaluate the consumption of the different structures of *Undaria* (blade and sporophyll) at the different developmental stages (growth, maturity, and senescence), six trials were conducted (two structures × three developmental stages), using two containers per trial. Sea urchins were randomly assigned to one part of each container (n=7 per container) and fed with a monospecific diet of blade or sporophyll (~ 2 g). The excess of water on the kelp surface was first dried using a salad spinner (high speed centrifugal force, 20 turns), following a modification of the method used by Hay *et al.* (1986). Blades and sporophylls were weighed at the beginning and at the end of the three-day experiment.

The remaining part of the container was used to evaluate the autogenic change of the kelp (i.e. Undaria structure in the absence of the sea urchin). To consider the potential variability in kelp weight attributable to autogenetic changes (Manly, 1993, Trowbridge, 1995), the consumption of Undaria was calculated considering the Undaria weight lost in the absence of the sea urchin (control) (Cacabelos et al., 2010). The consumption of Undaria was calculated using the following formula (1): Weight consumed (WC) = (WC)(Wi - Wf) - (WCi-WCf), where Wi is the initial blade or sporophyll wet weight and Wf is the final wet weight (after consumption). WCi is the initial wet weight of the control and WCf is the final wet weight of the control. In the cases where consumption had a negative value close to zero, this value was corrected to zero because it may be attributed to a scale error.

Differences in consumption between the different *Undaria* developmental stages were tested by using a nested ANOVA, using the container (1 and 2) factor nested within the developmental stage factor (growth, maturity and senescence) for each structure (blade or sporophyll).

Feeding preference experiment

To determine whether A. dufresnii chose Undaria blades over sporophylls or other macroalgae available in the environment, a feeding preference experiment was conducted following the methodology proposed by Underwood & Clarke (2005). The experiment consisted of two treatments: i) a no-choice treatment, aimed to estimate the quantity of each macroalgae species consumed when it was offered individually to the sea urchin, and ii) a **choice treatment**, aimed to estimate the consumption when two species were offered. Thus, preference was defined when the consumption of one item in the no**choice treatment** was lower than that in the choice treatment, while the consumption of the other item remained unchanged.

For the no-choice treatment, all macroalgae collected in winter were dried with a salad spinner (20 turns for laminar species and 40 turns for filamentous species). Two containers subdivided into eight compartments were used for each trial, as previously described for the *Undaria* experiment. Two grams (initial wet weight) of the same sections of each macroalgal species was always offered to avoid variation. After three days, the remains of macroalgae were removed, dried with the salad spinner, and weighed to obtain the final wet weight. The consumption of *Undaria* blades and sporophylls and each macroalgal species was calculated with the formula described above (1).

For the choice treatment, two containers were used for each trial (algae-Undaria pair). Four parts of each container were used for individual feeding of four sea urchins with 1 g of the macroalgae (Undaria sporophylls, Corallinales species, D. dichotoma, L. clavellosa, C. vermilara, Ceramium sp., or Ulva spp.) and 1 g of Undaria blades (growth stage). The remaining four parts were used as control (same amount of feed but in the absence of sea urchins). The total weight of feed offered during the no-choice treatment was equal to that offered at the choice treatment to avoid effects due to different encounter rates (Underwood & Clarke 2005). After three days of trials, macroalgae were dried, weighed and the consumption was calculated with the formula described above (1).

Preference was analyzed with a nested ANOVA, using the container (1 and 2) as a factor nested within the treatment (no-choice or choice) factor for each trial (Martins *et al.*, 2014). All analyses were performed in INFOSTAT (Di Rienzo *et al.*, 2019).

RESULTS

Undaria experiment

Arbacia dufresnii consumed both the blades and sporophylls of Undaria at all the developmental stages (Fig. 1). The consumption of blades did not differ significantly between developmental stages (F=0.74, p>0.05), whereas that of sporophylls was greater during the maturity stage than during the senescence stage (F=3.93, p<0.05).

Feeding preference experiment

The feeding preference experiment showed that, when *Undaria* was at the growth stage, *A. dufresnii* individuals preferred to feed on blades rather than on sporophylls. The weight of *Undaria* blades consumed was higher during the choice treatment than during the nochoice treatment (weight consumed: F=39.35, p<0.0001). Only 25% of sea urchins consumed sporophylls during the choice treatment (Fig. 2a).

Concerning the preference between Undaria blades and the other macroalgae available in the environment in winter, A. dufresnii selected Undaria blades over the Corallinales macroalgae and D. dichotoma. For the Corallinales species, the wet weight consumed was lower during the choice treatment (F=5.01, p<0.05, Fig. 2b), while D. dichotoma was not consumed during this treatment (Fig. 2c). In both cases, the Undaria wet weight consumed did not vary between treatments (Undaria- Corallinales: F=0.44, p>0.05; Undaria-D. dichotoma: F=1.40, p>0.05; Fig. 2 b,c).

Arbacia dufresnii did not show preference for either Undaria or Lomentaria clavellosa (Fig. 2d). The Undaria wet weight consumed showed no differences between treatments (F=0.41,p>0.05); however, there was a significant difference in the consumption of L. clavellosa (F=109.12, p<0.0001), but it was almost 100% consumed during both treatments. Regarding the consumption of Codium vermilara, A. dufresnii fed higher amounts during the choice treatment than during the no-choice treatment (F=7.87,p=0.01), while the consumption of *Undaria* did not vary between treatments (F=0.004, p>0.05) (Fig. 2e). In addition, in the Undaria-Ceramium sp. trial, no differences in the weight consumed were observed between treatments (Undaria: F = 0.78, p>0.05, Ceramium sp: F=3.31, p>0.05, Fig. 2f). Similarly, no preference in consumption was observed between Undaria and Ulva spp. and no differences between treatments were recorded (Undaria weight consumed F=0.73, p>0.05 and *Ulva* spp. weight consumed F=1.57, p>0.05, Fig. 2g).

DISCUSSION

Arbacia dufresnii consumed both blades and sporophylls of Undaria at all developmental stages. These results complement previous diet studies evaluating stomach contents and stable isotopes, which determined that the sea urchin consumes and assimilates the kelp in both spring and summer, when the sporophytes are mostly mature and senescent, respectively (Castro et al., 2022). Although sporophylls were consumed in all trials, preference experiments indicated that A. dufresnii preferred feeding on Undaria blades rather than on sporophylls at the growth stage. Similar results have been found in the native crab Pugettia producta, which also prefers blades over sporophylls in California, USA (Thornber et al., 2004). Previous studies have demonstrated that

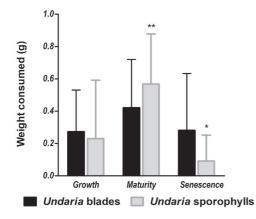


Fig. 1: Weight (g) of blades and sporophylls of *Undaria* pinnatifida consumed by Arbacia dufresnii in the feeding experiment at each developmental stage of the invasive macroalga (growth, maturity and senescence). * Lowest and ** Highest, statistically significant value.

the chemical composition of Undaria can change according to the tissue, developmental stage, and season, this latter related to abiotic factors, such as temperature or illumination (Skriptsova et al., 2010; Ank et al., 2014; Skriptsova, 2016; Arijon et al., 2021; 2023; Solana, 2022). Among chemical compounds, phenolic compounds are known to act as chemical defenses under conditions of natural herbivory (Haavisto et al., 2017) and have been found at higher amounts in Undaria sporophylls than in blades (Solana, 2022). Therefore, the preference of A. dufresnii for blades at the growth stage found in this study could be related to the higher concentration of chemical compounds in sporophylls, which may decrease their palatability and digestibility (Cronin & Hay, 1996, Van Alstyne et al., 2001). Preferences of native species to incorporate an invasive species into their diet may vary according to its developmental stage (Thornber et al., 2004). Future research involving preference experiments between blades and sporophylls of Undaria at different developmental stages could provide more information to understand consumption patterns of native sea urchins on the invasive kelp Undaria.

Undaria blades were also preferred over the brown seaweed Dictyota dichotoma. This result is in accordance with previous studies showing that although D. dichotoma is frequently present in the stomach contents of A. dufresnii, it is not assimilated into its tissues (Castro et al., 2022). Previous studies have shown that many herbivores reject species of the genus Dictyota, presumably due to the presence of deterrent secondary

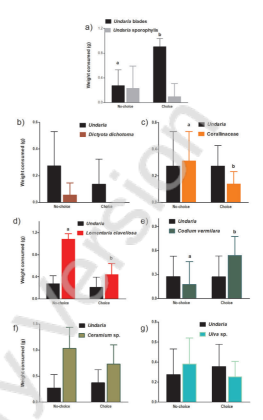


Fig. 2: Weight consumed (g) by Arbacia dufresnii during the no-choice and choice treatments of the feeding preference experiment. a) Undaria blades - sporophylls. b) Undaria - Dictyota dichotoma. c) Undaria - Corallinales. d) Undaria - Lomentaria clavellosa. e) Undaria - Codium vermilara. f) Undaria - Ceramium sp. g) Undaria-Ulva sp.

metabolites (Hay et al., 1986; Pereira et al., 2000; Paul et al., 2001; Souza et al., 2008; Spiers et al., 2021). Dictyota dichotoma is also an exotic species in the study area but, unlike Undaria, its introduction occurred much earlier (Orensanz et al., 2002; Raffo, 2014). This suggests that the presence of specific compounds in D. dichotoma may explain why it is not a food source for A. dufresnii, despite the long- term availability of the macroalgae in the environment, and the known dietary plasticity of the sea urchin (Penchaszadeh & Lawrence, 1999; Castro et al., 2022). The exclusive consumption of D. dichotoma in the no-choice treatment likely indicates that it was ingested solely because it was the only food source available. Regarding the Corallinales macroalgae, the sea urchin also showed preference for feeding on Undaria, probably because the calcification of coralline algae serves as a physical defense against herbivory (Litter et al., 1983).

In the "Undaria - Lomentaria clavellosa" trial, no preference for either macroalga was observed, but L. clavellosa was fully consumed during both the no-choice and choice treatments of the experiment. Previous studies have shown that L. clavellosa is found in high abundance in the stomach contents of A. dufresnii and is assimilated into the muscle (Castro et al., 2022). This inconsistency may be attributed to the insufficient amount of food offered to the sea urchins. Therefore, future experiments providing sea urchins with a greater quantity of food are necessary to definitely confirm the absence of preference. Likewise, in the "Undaria - Codium vermilara" trial, it was not possible to determine preference. In the choice treatment, A. dufresnii consumed more C. vermilara, while Undaria consumption did not vary between treatments. Species of the genus Codium have been frequently observed in the stomach contents of the native sea urchin and are relevant assimilated items, contributing more than 50% to gonadal and muscle tissues according to isotopic values (Castro et al., 2022). Various feeding experiments on different Codium species have shown that sea urchins consume cylindrical macroalgae more rapidly than flat-shaped macroalgae (Scheibling & Anthony, 2001; Lyons & Scheibling, 2007). However, despite the higher feeding rate, sea urchins obtain less energy from these cylindrical macroalgae (Lyons & Scheibling, 2007), suggesting the need to supplement with other items of higher nutritional quality.

When Undaria and Ulva sp. were offered together, we observed that A. dufresnii consumed both alternately, showing no preference between the two laminar macroalgae. Numerous reports have indicated that the green macroalgae Ulva sp. are often preferred by sea urchins (e.g. Navarrete et al., 2008; Souza et al., 2008; Cyrus et al., 2015) and other herbivores such as gastropods (e.g. Van Alstyne et al., 2009; Angell et al., 2012). These preferences have been attributed to their high nutritional value (Lawrence, 1975; Van Alstyne et al., 2009), and, in the case of A. dufresnii, it is significantly assimilated into muscle tissues (Castro et al., 2022). The similarity in the consumption between Undaria and Ulva sp. could be considered in nutritional value tests under the hypothesis that the invasive kelp Undaria is nutritionally similar to native species of the genus Ulva.

In summary, the native sea urchin Arbacia dufresnii did not exhibit strong feeding preference behavior, supporting evidence that this species has significant dietary plasticity and can adapt to newly available resources in the environment (Castro et al., 2022). The lack of preference for the most common macroalgae species over Undaria, even those with known high nutritional value, suggests that this invasive species may alter resource use and thereby affect the trophic structure of native communities. Similarly, this lack of preference has also been experimentally observed in other native herbivores (Thornber et al., 2004; Jiménez et al., 2015; Cardoso et al., 2020; Mabey et al., 2022). Therefore, our results also suggest that the successful distribution of Undaria is likely driven by factors other than herbivory escape (Cardoso et al., 2020; Mabey et al., 2022). On the other hand, our results differ from the experiments of Teso et al. (2009), where A. dufresnii did not feed on sporophylls, which could be due to the 10 years that have passed between studies. Santamaria et al. (2022) have recently suggested that native herbivores progressively learn to consume invaders. Hence, the differences associated with more than a decade of variation in the timing of macroalgal invasion also reinforce findings that Undaria contributes more significantly to the diet of A. dufresnii in sites with a longer history of invasion (Castro et al., 2022). The present research contributes to better understanding the magnitude of impacts caused by invasive macroalgal species through feeding ecology studies of native grazers.

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