Ecosystem engineering by burrowing crabs increases cordgrass mortality caused by stem-boring insects

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ABSTRACT: Some studies have shown that the balance between top-down and bottom-up processes is context dependent, but few have tested how biotic interactions can affect this balance. We quantified the attack frequencies by the stem borer moth Haimbachia sp. nov. on the cordgrasses Spartina densiflora and S. alterniflora in 5 marshes of the southwestern Atlantic coastline, located between 36° S and 41° S. We examined whether ecosystem engineering by burrowing crabs Neohelice (Chasmagnathus) granulata, which improves plant performance, increases herbivory incidence by moths. The results show that moths attack an important proportion of Spartina spp. stems (4% to 26%). Moth attack frequencies were higher on low elevation than on high elevation marsh plants, probably due to variations in plant performance across the physical stress gradient. Within marshes, burrowing crab densities and moth attack frequencies were positively correlated, suggesting that ecosystem engineering by crabs may increase moth attacks. Field experiments confirmed this prediction and suggest that the effect of crabs on moth attacks may be driven by nutrient availability. Our results show that stem-boring herbivores can cause significant plant mortality in Spartina spp. marshes, and that ecosystem engineering by burrowing crabs increases this effect. Thus, in this system, biotic factors that improve plant performance lead to an increase in herbivory control.

KEY WORDS: Bottom-up control · Haimbachia sp. nov. · Herbivory · Neohelice (Chasmagnathus) granulata · Spartina spp. · Top-down control · Salt marsh

INTRODUCTION

Ecologists have long debated how resource availability (bottom-up effects) and/or consumer pressure (top-down effects) shape natural plant communities (e.g. Teal 1962, Power 1992, Valiela et al. 2000, Bertness et al. 2008). It is now widely accepted that plant production can be influenced by feedbacks associated with these 2 effects (e.g. Walker & Jones 2001) and that top-down forces can increase with the quality of the environment (e.g. Menge & Sutherland 1976, Prezler & Boecklen 1996, Bertness et al. 2008, Sala et al. 2008). However, ecosystems are complex, and the identification of interactions between regulating forces becomes far more difficult when we attempt to understand how the balance between these opposing effects varies due to indirect interactions (Power 1992, Strong 1992). Such information is crucial to understand the factors that modulate productivity, and their effects on the systems.

Salt marshes are among the most productive systems (Mitsch & Gosselink 2001). In these environments, pri-
Mary productivity is mainly controlled by bottom-up process (physical stress or nutrients) that regulate trophic relationships (Odum & Smalley 1959, Teal 1962, Valiela et al. 2000). However, there is also evidence that consumers may influence marsh productivity (Pennings et al. 1998, Silliman & Zieman 2001, Jefferyes et al. 2006, Sala et al. 2008) and that their effects vary with physical stress gradients across tidal elevation (e.g. Moon & Stiling 2002, Rand 2002, Alberti et al. 2007) and increase in marshes with enhanced nutrient availability (Jefferyes 1997, Silliman & Zieman 2001, Bertness et al. 2008, Sala et al. 2008). Certain organisms can directly and/or indirectly control nutrient availability to other organisms (i.e. ecosystem engineers, Jones et al. 1994), leading to bottom-up control. Nevertheless, little is known on how top-down control of salt marsh plant productivity varies due to the indirect effects of ecosystem engineering species.

The salt marshes occurring between southern Brazil (32° S) and northern Argentinean Patagonia (42° S) are dominated by the cordgrasses *Spartina densiflora* and *S. alterniflora* (see Isacch et al. 2006) and by the burrowing crab *Neohelice granulata* (species formerly known as *Chasmagnathus granulatus*, see Sakai et al. 2006 for taxonomic revision). This burrowing crab is an efficient ecosystem engineer (sensu Jones et al. 1994, see Gutierrez et al. 2006, Daleo et al. 2007, Fanjul et al. 2007). In marshes, crabs excavate and maintain semi-permanent open burrows (e.g. Spivak et al. 1994) that can extend up to 1 m into the sediment, with surface openings of up to 10 cm diameter and with a density of up to 60 burrows m⁻² (Iribarne et al. 1997, Bottos et al. 2005). Burrow maintenance results in sediment removal of up to 2.5 kg m⁻² d⁻¹ (Iribarne et al. 1997), which increases soil nitrification-denitrification ratios due to enhanced soil oxygenation (Bottos et al. 2005, Gutierrez et al. 2006, Fanjul et al. 2007). These activities can boost plant productivity by facilitating mycorhizal formation (Daleo et al. 2007) and by increasing nutrient availability (Fanjul et al. 2008). These positive effects of crabs on plants could be transferred through the food web to other species. In fact, by increasing the density of reproductive stems, crabs positively affect granivorous species (Canepuccia et al. 2008), which increase their use of areas with higher densities of reproductive stems (marsh birds: Cardoni et al. 2007, rodents: Canepuccia et al. 2008). Thus, we may expect a similar process with herbivorous species. Given that most marshes are nitrogen-limited (Valiela & Teal 1974, Adam 1993), and that plants of higher quality as food item are more attractive to several herbivores (Moon & Stiling 2002), we hypothesized that the positive effect of burrowing crabs on plants will also indirectly favor the establishment of plant consumers, intensifying herbivore pressure.

Stem-boring insects are common marsh herbivores that can have important impacts upon their host plants (e.g. Stiling & Strong 1983, 1984, White et al. 2005). The stem borer moth *Haimbachia* sp. nov. (Crambidae: Crambinae) is commonly found in SW Atlantic *Spartina densiflora* and *S. alterniflora* marshes. The larvae overwinter inside grass stems, and when the adult moth emerges in spring and summer, it makes a characteristic hole through the stem (Fig. 1). The overwintering larvae feed on more than half of the central tissue of the stem, eventually killing the stem as a result of the damage (A. Canepuccia pers. obs.; Fig. 1). Thus, this moth can potentially have an important negative effect on *Spartina* spp. Stem borer larvae are restricted in their mobility (see Rathcke 1976, Stiling & Strong 1983), thus, mature females usually oviposit on high-quality hosts to maximize the fitness of their brood (Mayhew 1997, Moravie et al. 2006). In this context, we predict that higher nutrient availability due to burrowing crab activities (Bertness 1983, Fanjul et al. 2008) will promote attacks of *Haimbachia* sp. nov on *Spartina* spp. plants. Hence, this multi-species scenario (plant: *Spartina* spp., crab: *Neohelice granulata*, and moth: *Haimbachia* sp. nov.) may provide a model system to investigate how the intensity of herbivore pressure varies with other biotic drivers, and if this is a widespread pattern across some of the most extensive SW Atlantic coastal marshes. Thus, we examined: (1) the attack patterns on stems of *Spartina* spp. produced by moth larvae in salt marshes across marsh tidal height and throughout several SW Atlantic coastal marshes, (2) whether positive effects of *N. granulata* on *S. densiflora* indirectly increase herbivory pressure of *Haimbachia* sp. nov. on *S. densiflora*, and (3) given that the underlying mechanism could involve nutrient availability, whether increasing nutrient supply (bottom-up effect) could lead to increased herbivory pressure from moth larvae feeding on salt marsh plants.

![Fig. 1. *Spartina alterniflora* and *Haimbachia* sp. nov. (a) Longitudinal section of an attacked stem, with a moth larva inside, showing the area consumed by the larva. (b) The hole (arrow) produced by the moth when it emerges as an adult.](image)
MATERIALS AND METHODS

Study area. The study was carried out in 5 Argentinean marshes located between the mouth of the La Plata River estuary (Ria de San Clemente, 36° 22' S) and the northern part of Patagonia (San Antonio Bay, 40° 84' S) including some of the most extensive SW Atlantic coastal marshes (see Isacch et al. 2006; Fig. 2). Studied marshes are located at San Clemente (36° 22' S), Mar Chiquita coastal lagoon (37° 32' S), Bahía Blanca (38° 41' S), Bahía Anegada (39° 48' S), and Bahía San Antonio (40° 84' S, Fig. 2). The cord grasses *Spartina alterniflora* and *S. densiflora* and the perennial glasswort *Sarcocornia perennis* (formerly known as *Salicornia ambiguа*) cover 70% of these marshes (Isacch et al. 2006). The Mar Chiquita coastal lagoon is dominated by *S. densiflora*, while the other sites are inhabited by both cordgrasses. Given logistic and economic constraints, we performed the experiments to evaluate the effects of burrowing crabs on stem borer moth–*Spartina* spp. interactions with *S. densiflora* at the San Clemente and Mar Chiquita marshes (Fig. 2). In the studied marshes, *S. alterniflora* occupies sites at a lower marsh elevation than *S. densiflora*, which occurs at a wide range of intertidal elevations including the upper marsh edge (Isacch et al. 2006). Thus, for comparative reasons, we divided marshes into 4 intertidal levels according to their elevation: the lowest, occupied by *S. alterniflora*, and 3 upper levels occupied by *S. densiflora* (hereafter named ‘high,’ ‘middle,’ and ‘low’). The lowest marsh remains under water more frequently than the middle marsh, which develops at 50% of the higher tidal altitude. High marsh flooding occurs only during the highest spring or storm-induced tides.

Attacks by stem borer moths on *Spartina* spp. After adult moths emerged at the end of the summer (March 2008), we sampled *S. densiflora* and *S. alterniflora* stems to estimate the attack frequency of moth larvae. Given that herbivore damage on plants often varies spatially across tidal gradients (Rand 2002, Alberti et al. 2007), we collected 10 random samples of plants (containing 30 adult stems each) at each marsh level described above for each marsh location. All stems were cut at ground level, and then were longitudinally cut and classified as attacked (stems with larvae or evidence of their activity: holes, tunnels, or sawdust, Fig. 1) and not attacked (stems without larvae or evidence of their activity). We estimated the frequencies of moths attack on *S. densiflora* and *S. alterniflora* as the ratio between the number of moth-attacked stems and the total number of stems in each sample. A Tukey test following an analysis of variance (ANOVA; Zar 1999) was used to evaluate the null hypothesis of no differences in the proportions of attacked stems between marsh elevations.

Does the effect of burrowing crabs on *Spartina densiflora* increase the attacks by stem borer moths? The burrows of crabs and crab activity can locally have positive effects on plant growth (Bertness 1985, Kristensen 2008; see also Daleo et al. 2007 for this crab species). We evaluated whether there was a relationship between crab density and the proportion of stems attacked by moth larvae across marsh sites. To evaluate the possible relationship between crab density and the proportion of stems attacked by moth larvae, we collected plant samples in the low marsh, where crab densities are most variable. During the summer of 2008 (after adult moths emerged), we collected 30 randomly chosen stems inside each of 50 quadrats (0.5 × 0.5 m) with varying crab densities in the low *S. densiflora* zone at each study site. The proportions of attacked stems were assessed as described above. To estimate crab densities, we counted the total number of crab burrows in each plot (a good estimator of crab densities, see Iribarne et al. 1997, Bortolus & Iribarne 1999). Correlation analysis (Zar 1999) was employed to examine the relationship between crab density and moth attack at each marsh site.

To experimentally evaluate the association between crabs and the proportion of stems attacked by moth larvae, we performed a caging experiment manipulating crab densities from February 2005 to May 2007.
Thirty permanent cages (plastic mesh squares 1 x 1 x 0.5 m; mesh size: 1.5 cm, which allowed free movement of moths but excluded crabs) were deployed at an intermediate elevation at Mar Chiquita. We selected this marsh elevation because crab densities are lower than at the lower marsh level, and thus it was relatively easier to keep them from invading the exclusion treatments. It was also easier, and resulted in fewer undesired effects, to increase crab burrow density than to decrease it. We buried cages 20 cm into the sediment to prevent crabs from burrowing beneath them. We cut roots and rhizomes with a shovel around the perimeter of each cage to a depth of 20 cm to avoid resource reallocation through clonal integration. Cages were randomly assigned to 3 different treatments: (1) crab inclusion: 10 cages in which we used a corer to make 30 artificial burrows (4 cm diameter, 1 m deep) and included 1 crab (3 to 4 cm carapace width) in each, i.e. 30 ind. m⁻² (similar to field densities, Botto et al. 2005); (2) burrows without crabs: 10 cages with 30 artificial burrows but without adding crabs, to independently evaluate the effect of burrows; (3) full exclusion: 10 cages in which we removed crabs from natural burrows by hand and then filled the burrows with sediment from an adjacent area. Treatments were maintained weekly to ensure that every burrow was active in treatment (1), that every burrow was not active in treatment (2), and that no crab invaded treatment (3) during the study period (for similar design see Bortolus & Iribarne 1999, Bortolus et al. 2004, Canepuccia et al. 2008). After more than 2 yr (from February 2005 to May 2007), we collected 30 random Spartina densiflora stems cage⁻¹ to determine the proportion of attacked stems as described above. To fit the assumption of linearity and normality, the proportion of attacked stems from the Mar Chiquita study site were square transformed (Zar 1999). A Tukey test after significant ANOVA (Zar 1999) was used to evaluate the null hypothesis of no differences between treatments in the frequency of stems attacked by moths. Does increased nutrient supply lead to an increase in herbivory pressure by stem borers? Given that burrowing organisms usually increase soil nutrients (Bertness 1985, Fanjul et al. 2007, 2008), we performed a second experiment to evaluate whether the frequency of Spartina densiflora stems attacked by moths increased with soil nutrient levels, at San Clemente and Mar Chiquita marshes from December 2005 to April 2007. Quadrats (0.7 x 0.7 m) were randomly selected at the S. densiflora marshes of San Clemente (14) and Mar Chiquita (16). Plots were randomly assigned to 1 of 2 treatments: (1) nutrient addition (n = 7 and 8, respectively) and (2) control (no nutrients added; n = 7 and 8, respectively). Nutrient addition plots received 30 g of fertilizer pellets containing 29% N, 5% P, and 5% K every month (see Emery et al. 2001, Daleo et al. 2008 for similar procedures). At the end of the experiment, after adult moth emergence in summer, we randomly cut 30 stems of S. densiflora in each plot to determine the proportion of attacked stems. The null hypothesis of no differences in the proportion of attacked stems between treatments was evaluated with t-tests (1 for each site, Zar 1999).

**RESULTS**

**Attacks by stem borer moths on Spartina spp.**

Overall, the analysis of 10400 Spartina spp. stems showed that moth larvae attacks varied from 4 to 14% of S. alterniflora stems and between 6 and 26% of S. densiflora stems depending on the marsh (Fig. 2). The larvae make a characteristic hole between the second and third stem internodes for drainage of excretions and cached food (Fig. 1). At least half of each stem with larvae or pupae inside was dry, whereas stems that were alive and not attacked by moth larvae were entirely green. All attacked stems from which the larvae had already emerged as moths were completely dry. The proportion of attacked stems varied with marsh elevation. Three of the 5 marshes sampled (San Clemente, Mar Chiquita, and Bahía Blanca; Fig. 2) showed a much higher proportion of attacked stems in samples collected from the low marsh than in those collected from the high marsh (Tukey test after ANOVA, Fig. 3a). The other 2 marshes (Bahía Anegada and Bahía San Antonio) showed lower moth infection than the 3 marshes indicated above, and showed no differences between marsh elevations (Fig. 3a).

**Does the effect of burrowing crabs on Spartina densiflora increase attacks by stem borer moths?**

There were positive relationships between crab densities and the proportion of attacked stems in all S. densiflora marshes (Fig. 3b). The experiment manipulating crab densities confirmed that stem damage is related to the presence of crabs. Cages with crabs and burrows had, on average, more than twice the proportion of attacked stems of S. densiflora than cages from which crabs and their burrows were excluded (Tukey test, p < 0.05 after ANOVA, F₂, 27 = 4.67, p = 0.018, Fig. 4). Instead, the proportion of attacked stems in cages with only burrows (without crabs) was not different from the attack proportion observed in cages with crabs and burrows, and in cages without crabs and burrows (Tukey test, p > 0.05, Fig. 4).
Fig. 3. *Spartina* spp. and *Haimbachia* sp. nov. across 5 southwestern Atlantic coastal marshes. (a) Proportion of stems of *S. alterniflora* and *S. densiflora* attacked by the moth at different marsh elevations. (b) Relationship between the proportion of stems of *S. densiflora* attacked by the moth, and crab densities in the low marsh (n = 50, see also Fig. 2). In this and subsequent figures: limits of the box: 25th and 75th percentiles, vertical lines: minima and maxima, black dot inside the box: median value. Different letters indicate differences (p < 0.05) by Tukey test after ANOVA; na: absence of plants at that marsh elevation.
Does increased nutrient supply lead to an increase in herbivory pressure by stem borers?

In both *Spartina densiflora* marshes (San Clemente and Mar Chiquita), nutrient addition resulted in a 3-fold higher proportion of attacked stems (respectively: mean ± SD = 21.48 ± 10.82; 5.57 ± 3.91) than in control plots (respectively: 6.43 ± 4.76, \( t_{12} = 3.37, p = 0.0056; \) 1.62 ± 2.45, \( t_{16} = 2.56, p = 0.0041 \), Fig. 5).

**DISCUSSION**

Our results show that in several representative marshes of the SW Atlantic coastline, the moth *Haimbachia* sp. nov. is an important herbivore on *Spartina* spp. We also show that the positive effects of the crab *Neohelice granulata* on plants cascaded up, resulting in increased moth attack frequencies on *S. densiflora*. Thus, part of the positive effect of crabs on *S. densiflora* results in an indirect negative effect on the plants. Therefore, crabs, acting as ecosystem engineers, alter the energy flows between trophically interconnected organisms, leading to an increase in herbivore incidence.

Stem borers can be very common in marshes. In northwest Florida (USA) marshes, for example, at least 40% of the total available stem internodes of *Spartina alterniflora* can be perforated by larvae of different insect species (Stiling & Strong 1983). In *S. alterniflora* collected along the Louisiana (USA) coast, 20% of the stems supported at least 1 larva, whereas an additional 17% exhibited evidence of prior insect feeding (White et al. 2005). Throughout SW Atlantic coastal marshes, *Haimbachia* sp. nov. attacked an important fraction of *S. densiflora* and *S. alterniflora* stems (from 4% to 26%, depending on marsh elevation and site). Stem borers can have negative effects on marsh plants (e.g. Klopatek & Stearns 1978). Indeed, our results show that by the end of the life cycle of the larvae (summer), at least half of the length of each attacked stem from which moths had not emerged yet (larvae or pupae) was dry, whereas all attacked stems from which moths had already emerged were completely dry. Given that across SW Atlantic coastal marshes *Spartina* spp. do not die in winter but grow all year (stems live for more than 1 yr, e.g. Vera et al. 2009), these observations suggest that feeding by moth larvae has an impact on the salt marsh community by increasing the availability of dry stems.

Our results also showed that in marshes with a high proportion of attacked stems, moth attack was more frequent on *Spartina alterniflora* than on *S. densiflora* (see Fig. 3). This pattern may be due to different plant palatability or moth preference (e.g. Goranson et al. 2004). However, this pattern could also result from differences in physical stress across the tidal gradient. *S. alterniflora* occurs at a lower elevation of the marsh than *S. densiflora* (Isacch et al. 2006). Indeed, the damage by moths on *S. densiflora* marshes was also higher in the low marsh compared to the high marsh. Plants that grow at higher tidal elevations are usually exposed to limiting hypersaline soil conditions as a result of pore water evaporation, and low nutrient availabil-
ity, which can have profound impacts on salt marsh plant performance (Bertness et al. 1992, Emery et al. 2001, Goranson et al. 2004) and consumers (Moon & Stiling 2000, 2002, Rand 2002). Thus, environmental factors might play an important role in mediating marsh plant–moth interactions.

Environmental conditions for plant growth are generally harsh in salt marshes (e.g. Pezeshki 2001). Ecosystem engineering by burrowing organisms usually changes the primary limiting resource, potentially having strong effects on the conditions in harsh habitats (e.g. Jones et al. 1997). For example, in alpine meadows, mounds from burrowing plateau zokors *Myospalax baileyi* enhance the biomass of adjacent plants (Wang et al. 2008). Also, in the arid Negev Desert Highlands of southern Israel, digging by the crested porcupine *Hystrix indica* enhances the growth of annual plants (Alikon 1999). In cool desert environments, the added moisture from spring recharge in ground squirrel *Spermophilus townsendii* burrows can increase plant productivity (Laundré 1998). Also in salt marshes of the east coast of North America, the burrowing activity of mud fiddler crab *Uca pugnax* (Bertness 1985) and the burrowing activity of *Neohelice granulata* in SW Atlantic coastal marshes increases the production of *Spartina* spp. (Daleo et al. 2007). The most likely mechanisms of enhanced *Spartina* spp. growth include that crab burrows increase sediment drainage and oxygen concentrations (Bertness 1985, Gutierrez et al. 2006, Daleo et al. 2007, Fanjul et al. 2008), which in turn may reduce phytotoxin concentrations (e.g. Smith et al. 1991), an effect that might be even more important in the highly anoxic low marsh. In addition, burrow structure may enhance nutrient availability, possibly because burrows act as detritus traps (Botto et al. 2006), and may also accumulate ammonium (Fanjul et al. 2008) from crab excretions and cached food (Bertness 1985,Fanjul et al. 2008, Kristensen 2008). Consequently, the positive effects of crab activities on plant productivity (Bertness 1985, Daleo et al. 2007) could indirectly facilitate primary consumers. Indeed, on the large scale of our sampled marshes, there is a positive relationship between crabs and moth spatial distribution. Overall, moths attacked more than 50% of *S. densiflora* stems in areas with high crab densities and less than 10% in areas with lower crab densities. Our experiment evaluating this pattern confirmed that this association was driven by crabs and not by an underlying factor affecting both species (moths and crabs). Plants growing in cages with crabs (and their burrows) had more than twice the proportion of attacked stems than plants in cages from which crabs (and their burrows) were excluded. The proportion of attacked stems in cages with burrows only (no crabs) showed intermediate values that did not differ from either of the other treatments. This intermediate attack frequency in stems from cages with only burrows could mean that the effect of burrowing crabs on moths is mediated by 2 mechanisms: (1) crab burrows (which can increase sediment drainage and oxygen concentrations and (2) crab activities which can increase nutrient availability from excretions and cached food, Kristensen 2008. Nutrient additions also increased moth attack frequencies, suggesting that crab effects on moth attacks may be driven by nutrient availability. As a consequence, the positive bottom-up effects of crabs on *Spartina* spp. improve their quality as a food resource and make these plants more susceptible to attacks by moths.

Thus, crabs have positive effects on plant growth mediated by ecosystem engineering processes (Daleo et al. 2007) that can cascade to upper trophic levels (Cardoni et al. 2007, Canepuccia et al. 2008, this study). However, crabs are also herbivores, and hence, they can have negative effects on plants (Bortolus & Iribarne 1999, Costa et al. 2003, Bortolus et al. 2004, Alberti et al. 2007), particularly on plants of marsh-mudflat edges, or in isolated plants compared to those surrounded by neighbors (Alberti et al. 2007). The balance between those negative and positive effects by crabs upon plant growth is regulated by the physical characteristics of the local environment where interactions take place (Daleo & Iribarne 2009). While crab presence increases sediment aeration and nutrient availability, these positive effects are most important in stressful zones (e.g. where oxygen availability is low, Daleo & Iribarne 2009). However, in zones where the naturally high porosity allows better oxygenation, there can be an increase in the negative effects of crabs (Daleo & Iribarne 2009). This might explain the weaker relationship between crab densities and moth attack frequencies in the 2 salt marshes located at the southern distribution of our study sites (and the lack of differences between marsh elevations). These marshes develop on larger grain size sediments than those located to the north (Daleo & Iribarne 2009), which may increase the negative effect of crab herbivory.

Bottom-up and top-down forces act on plant production simultaneously, and the balance between the 2 ways of regulation depends on the environmental context. Studies have shown how top-down factors can increase together with the quality of the environment in a wide variety of harsh ecosystems (e.g. mountain, Prezler & Boecklen 1996; goldenrod community, Uriarte & Schmitz 1998; salt marshes, Bertness et al. 2008, Sala et al. 2008). In this context, our study shows how an ecosystem engineering crab positively affects plant productivity (e.g. Daleo et al. 2007), indirectly increasing top-down control by modulating herbivore inci-
ence. Therefore, given that ecosystem engineers that ameliorate physical stress are a common component of harsh environments (see Wright & Jones 2004, 2006), our work suggests the possibility that in these systems, biotic factors that increase plant performance will lead to an increase in herbivory control.

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LITERATURE CITED


Klopopate JM, Stearns FW (1978) Primary productivity of emergent macrophytes in a Wisconsin freshwater marsh ecosystem. Am Midl Nat 100:320–332


Power ME (1992) Top-down and bottom-up forces in food webs: Do plants have primacy? Ecology 73:733–746


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