Selective Feeding by a Predatory Sea Star Across a Depth Gradient in Northern Patagonia, Chile

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Sea stars often function as keystone predators in food webs of intertidal and subtidal communities, especially in temperate and sub-polar regions. In South America the sea star Cosmasterias lurida is distributed along both the Atlantic and Pacific coasts of Patagonia and is one of the most conspicuous and abundant benthic predators in the shallow subtidal zone (<25 m). Its feeding strategy and prey selection are, however, still poorly known. This study describes the feeding behavior of C. lurida at a site in the Seno del Reloncaví (Chile), assessing its abundance, size and prey selection in the field relative to observed prey abundance and size along a bathymetric gradient. We hypothesized that C. lurida is a generalist predator, feeding on suitable prey according to their availability. However, we found that this predator only consumed a limited number (7 of 48) of potential prey species, primarily the slipper limpets Crepipatella spp. and the mussels Aulacomya ater and Mytilus chilensis. Electivity analysis revealed a clear preference for one mussel (A. ater) but not the other (M. chilensis) as well as depth-dependent selectivity for the slipper limpets, which changed from avoidance to preference with increasing depth. Sea star densities varied with depth, peaking between depths of 5 and 10 m, but the size of sea stars and the size of their prey did not vary significantly along a depth gradient. No significant correlations were found with the most commonly selected prey. These results would indicate that while this predator may be a generalist–opportunist, its feeding behavior is context-dependent and its high selectivity for certain species suggests that this sea star plays a key role structuring subtidal benthic communities in Patagonia.

Keywords: Cosmasterias lurida, benthic ecology, feeding behavior, starfish, predation, dietary preference

INTRODUCTION

Predators strongly affect populations of their prey, but in turn, the availability of prey also regulates the behavior of predators (Sih et al., 1985; Gaymer and Himmelman, 2002; Ross et al., 2003; Navarrete and Manzur, 2008; Skein et al., 2018). In benthic marine communities, sea stars are one of the most active predators and control both directly and indirectly the abundance and distribution of numerous species (Paine, 1966; McClintock, 1994; Saier, 2001; Manzur and Navarrete, 2011; Calderwood et al., 2016; Gianguzza et al., 2016). They have thus been recognized as important...
components of intertidal and subtidal communities, at times even being considered keystone species (Paine, 1966; Dayton, 1985; Gaymer and Himmelman, 2008; Menge and Sanford, 2013).

The effect of sea stars on benthic communities depends on their feeding strategy and the trophic level of their prey in the community (Ross et al., 2003; Menge and Sanford, 2013; Motti et al., 2018). Many sea stars are opportunistic omnivores [e.g., Oreaster reticulatus (Linnaeus, 1758), Martin et al. (2001)], but some species are specialized predators [e.g., Heliaster helianthus (Lamarck, 1816), Meyenaster gelatinosus (Meyen, 1834), Urriago et al. (2011)], herbivores [e.g., Phatariaria unifascialis (Gray, 1840), Phataria pyramidatus (Gray, 1840), Salguero and Bonilla (2010)], or detritus feeders [e.g., Pentaceraster cumingi (Gray, 1840), Salguero and Bonilla (2010) and Hyphalaster inermis (Sladen, 1883), Mironov et al. (2016)]. The prey of sea stars consist of a wide range of organisms including sponges, sea anemones, mollusks, polychaetes, crustaceans, and even other echinoderms (Mutschke and Mah, 2009). Further, cannibalism as well as ontogenetic changes in diet can occur (Verling et al., 2003; Urriago et al., 2012; Baeta and Ramon, 2013; Fernandez et al., 2017; Deaker et al., 2020). For example, Heliaster helianthus (Lamarck, 1822) a keystone predator in rocky intertidal habitats of central Chile, showed ontogenetic changes in habitat and diet composition of prey as it grew. That is, when individuals were recruits inhabit boulders and crevices in the high or mid-high intertidal zones preying on small species, mostly on the periwinkle Austrolittorina araucana (d’Orbigny, 1840) while adults prey on more species including mussels and limpets in the lower intertidal zone (Manzur et al., 2010).

In southern South America, the sea star Cosmasterias lurida (Philippi 1858) is one of the most abundant benthic predators in shallow subtidal habitats and can be found on both soft sediments and rocky bottoms (Figure 1). C. lurida is widely distributed along the temperate shores of South America, ranging from La Serena (29° 56’ S) on the Pacific coast of Chile to Golfo de San Matías (38° 00’ S) on the Atlantic coast of Argentina (Madsen, 1956; Hernández and Tabladillo, 1985; Clark and Downey, 1992) as well as around the Falkland (Malvinas) Islands, Burdwood Bank, and South Georgia (Vásquez and Castilla, 1984; Hernández and Tabladillo, 1985; Fraysse et al., 2018). Although its bathymetric distribution is large, ranging between the lower intertidal zone and 650 m in depth (Madsen, 1956; Clark and Downey, 1992), highest abundances have been recorded in shallow water habitats (Vásquez and Castilla, 1984; Pastor-de-Ward et al., 2007). Despite its wide geographic and bathymetric distribution, previous work on this species has focused mainly on its reproductive biology (Pastor-de-Ward et al., 2007; Cossi et al., 2015, 2017; Fraysse et al., 2020) and biochemistry (Seldes and Gros, 1985; Maier et al., 1993, 1998; Roccatagliata et al., 1994). However, a wide range of prey items in its diet has been also recorded (Castilla and Moreno, 1982; Vásquez and Castilla, 1984; Pastor-de-Ward et al., 2007; Gordillo and Archuby, 2012), and C. lurida is thought to be an important consumer in the shallow benthic food webs along the Patagonian coast (Adami and Gordillo, 1999; Schjetner et al., 2008; Gordillo and Archuby, 2012; Amsler et al., 2014; Cossi et al., 2015; Fraysse et al., 2018). For example, C. lurida within kelp beds [Macrocystis pyrifera (L.) C. Agardh, 1820] in Tierra del Fuego (Chile) mainly preyed on barnacles [Balanus spp. (Costa, 1778)] and slipper limpets [Crepipatella dilatata (Lamarck, 1822)] but also consumed 25 other species, including other gastropods, other crustaceans, bivalves, ascidians, brachiopods, fish, priapulids, sea urchins, and carrion (Vásquez and Castilla, 1984). In contrast, the main prey items of C. lurida in the shallow soft sediment environments of the Magellan Strait (Chile) were endobenthic bivalves, primarily Ameghinomyxa antiusa (P. P. King, 1832) (Garrido, unpublished data). These differences in diet suggest that this species behaves as a generalist-opportunistic, being able to use different resources depending on the prey availability in the habitat (Ross et al., 2003). However, understanding trophic relationships between predators and prey requires information on both the availability of prey and the preference of the predator. Although spatial associations between predators and prey have been interpreted as preferences, true preference requires an explicit behavior (Singer, 2000; Underwood et al., 2004) where the predator selects a particular prey over others. A proxy of preference, known as electivity, can be estimated as the difference of the relative proportion of prey in the diet compared to the available relative proportion in the local environment (Singer, 2000; Underwood et al., 2004).

Although previous studies (Madsen, 1956; Vásquez and Castilla, 1984; Gordillo and Archuby, 2012) have observed C. lurida feeding on different prey items, Vásquez and Castilla (1984) suggested that this predator is an opportunistic-generalist with little selectivity of prey. In the northern Patagonian zone of Chile, C. lurida, like many higher trophic level sea stars, feeds on a wide range of prey in shallow rocky environments. In this study we test the hypothesis that the diet of C. lurida would reflect prey availability in the environment, i.e., no selectivity of prey. We tested this hypothesis by quantifying sea star abundance, prey abundance, and prey electivity across a depth gradient where pronounced changes in the prey availability.

**MATERIALS AND METHODS**

**Study Site**

Field observations using SCUBA were conducted in 2010 from May to July (austral winter) at Yerbas Buena Bay (41° 40’ 20” S; 72° 39’ 25” W), a cove located in Seno del Reloncavi, a large bay extending 34 km south from Puerto Montt, Chile (Figure 2). Seno del Reloncavi geographically marks the end of Chile’s central valley and the beginning of the Patagonian region and is the first of several large extensions of channels and fjords of the Golfo de Ancud (Soto-Mardones et al., 2009). During our research, tidal level was measured continuously every 10 min at depths of 5 and 20 m with SOLINST pressure sensors, where maximum measured difference between low and high tide was approximately 7 m. All reported depths were corrected to the level of the annual mean tide to be able to compare data taken by SCUBA divers during different tide levels.

**Predator and Prey Abundance**

To quantify sea star densities and assess prey availability at this site, we selected five transects perpendicular to the coastline from...
FIGURE 1 | The sea star Cosmasterias lurida on different substrata in southern Chile. (A) Feeding on mussels on a rocky bottom at 7-m depth; (B) feeding on Crepipatella spp. on a rocky bottom at 8-m depth; (C) occurring at high densities on a rocky bottom at 10-m depth; (D) foraging on a sandy bottom at 9-m depth (A–C – Yerbas Buenas Bay, Seno del Reloncái; (D) – Faro San Isidro, Magellan Strait, Chile).

FIGURE 2 | South America (A) Seno del Reloncái and (B) the study site at Yerbas Buenas Bay (C).
the lowest intertidal level to a depth of 30 m and separated from one another by 10 m. Using a subaqueous GPS connected to a surface buoy with a GPS antenna (Schories and Niedzwiedz, 2012), we georeferenced these transects, allowing future studies to precisely relocate the study area without the need for physical markers in the field. For this study, transects were censused only once, each on a different day between May 1 and June 29 with two objectives: (1) determine the density of C. lurida and (2) assess the availability of potential prey (see details below). For each transect, two divers connected by a rope of 3-m length worked in parallel. While one diver handled the subaqueous GPS and measured depths, the other diver counted all C. lurida between them and recorded photo-quadrants (0.12 m²) using an underwater camera (Nikon D300 inside a Sea and Sea underwater housing) mounted over an aluminum frame that assured that all photo-quadrants were taken from the same distance and were perpendicular to the bottom.

**Cosmasterias lurida Density in Relation to Depth**

As described above, C. lurida were counted by one of the divers along each transect within the area separating the divers for the length of the transect within each 5-m depth intervals, starting from the surface down to 30 m (i.e., six depth intervals in total: 0–5, >5–10, >10–15, >15–20, >20–25, >25–30 m). The actual area surveyed within each depth range depended on the bottom slope and was calculated using the starting and ending points of the GPS positions at depths of 0, 5, 10, 15, 20, 25, and 30 m and the difference in depths between each interval. In this way following the Pythagorean theorem we calculated the total distance traveled for each depth interval and multiplied it by the transect width (3 m). The density of C. lurida was then calculated as the number of individuals per m² (ind./m²).

**Prey Availability in Relation to Depth**

Along each transect, three photo-quadrants of the bottom (see above), separated by at least 2 m from each other, were taken at each 1 m depth interval along the transects to estimate the prey field (i.e., the relative abundances of epibenthic invertebrates considered to be potential prey items of C. lurida). A total 360 photo-quadrants were thus recorded along the five transects. Photographs were analyzed using the software Coral Point Count with Excel extensions [CPCe 3.6; (Kohler and Gill, 2006)] to superimpose a uniform 10 × 10 grid of points on each image after which the benthic component (algae, sessile and mobile invertebrates, bare substratum) under each point was determined (Figure 3). Nearly all organisms were able to be identified at the species level, and only a few images of poor quality (due to shade, excess particles in the water column or distortion) could not be identified either at the species level or the taxonomic group. In these cases, they were classified as unknown. Any organisms that were identifiable in the photo were also noted. Empty shells of the most abundant mollusks [the mussels *Aulacomya ater* (Molina, 1782) and *Mytilus chilensis* (Hupe, 1854) and the slipper limpets *Crepipatella* spp.] could be easily identified in the images. They were not included in estimates of the relative abundances of epibenthic invertebrates but gave additional evidence of possible important prey items for C. lurida. *Crepipatella* spp. included two cryptic species, *Crepipatella peruviana* (Lamarck, 1822) and *C. dilatata* that could not be distinguished in the field.

**In situ Feeding of Cosmasterias lurida**

Four transects running perpendicular from the shore to a depth of 30 m were also surveyed to observe in situ feeding behavior of C. lurida and determine if there were differences in the prey consumption across this depth gradient. Each transect was again divided into five intervals of 5 m of depth (see above). Within each depth interval we turned over every *C. lurida* encountered and recorded if the sea star was feeding and if so, what prey was being consumed. Observations were divided into five behavioral classes: (1) feeding on mussels (*Aulacomya ater* or *Mytilus chilensis*); (2) feeding on the slipper limpets *Crepipatella* spp.; (3) feeding on other species; (4) stomach extended but without any retained prey (with activity – “W/activity”); and (5) no activity (no feeding – “N/activity”). As the number of observations was not identical between each transect and depth interval, it was standardized as a percentage. In addition, the first 20 sea stars that were observed feeding (i.e., with stomach everted and a prey trapped within) within each depth interval of a given transect were collected and placed in separate mesh bags together with the prey item. The wet weight and disk diameter were measured for each sea star and body length for the associated prey item.

**Statistical Analysis**

**Cosmasterias lurida Density in Relation to Depth**

Differences in density of sea stars among depth intervals were analyzed with the non-parametric Kruskal–Wallis test and pairwise comparisons by a Dunn test, because the data did not comply with the assumptions of normality and homogeneity of the variances, even after testing various transformations.

**In situ Feeding of Cosmasterias lurida**

Feeding activity data of C. lurida fulfilled the assumption of normality (Shapiro Wilk) and homogeneity of variance (Levene’s test). Thus, a two-way ANOVA was used to compare differences in feeding activity with depth interval. We performed a posteriori test (Tukey HSD) for cases that were significant (p < 0.05). The proportion of the species preyed upon by sea stars among depth intervals was visually examined using a Principal Coordinate Analysis (PCO) performed on a Bray-Curtis dissimilarity index matrix estimated from the relative abundance (%) of prey item at each depth. Then, we use a One-way PERMANOVA to test differences in prey proportions along the depth gradient. The data were not transformed to calculate the resemblance matrix. A Pearson correlation was used to determine the relationship between length of the predator and the three most frequently observed prey species consumed by the sea star. Univariate analyses were performed on R software (version 4.0.2; R Development Core Team, 2020) and multivariate analyses were performed on PRIMER software (Version 7.0).

**The Dietary Electivity of Cosmasterias lurida**

The dietary electivity analysis was performed using a selection index of relative prey availability (Pearre, 1982). The coefficient “c” [which is a correction of V coefficient of Yule (X²)]
FIGURE 3 | Image analysis of photo-quadrats (example from 10-m depth) with red crosses indicating the matrix of uniform 10 × 10 points used to assess potential prey abundance. The mussel *Aulacomya atra* (*Aa*), the sea urchin, *Arbacia dufresnii* (*Ad*), and the slipper limpets *Crepipatella* spp. (*Cr*) are easily identified in the image.

(Kendall, 1952; Kendall and Stuart, 1973) was calculated for each item by depth interval using the following formula:

\[
C = \left( \frac{X^2}{n} \right)^{1/2} = \sqrt{n \left( \frac{|a_d b_e - b_d a_e| - n/2}{a b d e} \right)}
\]

where:
- \(n\): Number of total individuals feeding,
- \(a_d\): Number of a given prey item in the diet,
- \(b_d\): Available number of the other prey items in the environment,
- \(a_e\): Available number of a particular prey item in the environment,
- \(a\): Sum of \(a_d\) and \(a_e\),
- \(b\): Sum of \(b_d\) and \(b_e\),
- \(d\): Sum of \(a_d\) and \(b_d\),
- \(c\): Sum of \(a_e\) and \(b_e\).

The C index ranges from −1 to +1 with the value 0 indicating no selection, i.e., the prey was consumed according to its availability. A value of 1 is absolute preference (maximum selectivity), whereas a value of −1 indicates that the predator avoids the prey completely. Those extreme values are called absolute associations (Kendall, 1952). The significance of prey selectivity index C was tested using a Chi-square test (Pearre, 1982).

**RESULTS**

**Distribution and Density of *Cosmasterias lurida***

*Cosmasterias lurida* was restricted from 2 to 25 m of depth at Yerbas Buenas Bay (Figure 4) and was not observed deeper even though rocky substratum continued to a depth of 35 m. Densities reached peak abundance in the >5–10-m interval (0.93 ± 0.56 ind/m²) where they were over twice the levels observed in any other interval. In contrast, densities were very low in the >20–25-m interval, less than 10% observed overall. Sea star abundance was significantly different among depth intervals [Kruskal–Wallis, \(H(5, \ n = 140) = 39.4; \ p < 0.001\); Dunn method].
Bathymetric Distribution of Potential Prey Species

Forty-eight species of benthic invertebrates considered as potential prey items were identified in the survey (Supplementary Material) and covered 56% of the bottom at this site with the remaining surface being shell fragments and sand (31%), bare rock (10%, including bedrock and cobbles), and mud (4%). The percent covers of the general taxa were 16% sessile gastropods, 15% echinoids, 11% algae, 3% bivalves, 3% holothuroids, 3% crustaceans, 2% anthozoa, 1% polychaetes, 1% porifera, <1% polyplacophores, <1% ascidians, <1% brachiopods, <1% bryozoans, <1% hydrozoans, and <1% asteroids. The most abundant potential prey species at the site were the slipper limpets Crepipatella spp., the mussels Aulacomya ater and Mytilus chilensis, the barnacles Notobalanus flosculus (Darwin, 1854), Elminius kingii (Gray, 1831), Balanus laevis (Bruguère, 1789), and Austromegabalanus psittacus (Molina 1788), and the sea urchins Pseudechinus magellanicus (Philippi, 1857), Arbacia dufresnii (Blainville, 1825), and Loxechinus albus (Molina, 1782).

Unlike most gastropods, Crepipatella spp. are sessile. They were most abundant between depths of 0 and 10 m (18–20% cover) but decreased thereafter, falling to 3% within the deepest interval where sea stars occurred (>20–25 m; Figure 5). Other sessile species were mussels and barnacles. They were also more abundant in shallow depth intervals and, indeed, were not observed below 15 m, except for the barnacle A. psittacus, which was recorded only in some transects below the 15 m but in low abundance. The two mussel species, A. ater and M. chilensis, were less abundant than Crepipatella spp., reaching only 1% and 11% cover, respectively, in the shallowest depth interval (Figure 5). Barnacles were relatively abundant (13% in this same depth interval) but mainly consisted of small individuals (<1-cm diameter). Among mobile animals, the sea urchins P. magellanicus, A. dufresnii, and L. albus were the most abundant and were recorded in all depth intervals (Figure 5). P. magellanicus was the most abundant sea urchin species and occurred primarily in depths from 10 to 15 m. A diverse group of mobile small gastropods (<1 cm) was observed at the site but even collectively they were not abundant (Figure 5, Supplementary Material, and Table 1).

In situ Feeding Activity of Cosmasterias lurida

Regarding the feeding activity survey of C. lurida, significant differences were found in the interaction between the prey items and the depth intervals ($F = 8.512; df = 9, 48; P < 0.001$) (Table 2). Regardless of the depth interval, approximately half the sea stars sampled were not feeding, and no individuals of C. lurida were found feeding at depths below 20 m (0–5 m = 55%; >5–10 m = 46%; >10–15 m = 43%; >15–20 m = 57%; >20–25 = 0%; Figure 6). Among those that were feeding (0–5 m = 42%; >5–10 m = 45%; >10–15 m = 52%; >15–20 m = 31%; Figure 6), only seven species of the 48 taxonomic groups recorded in the photo-quadrats were observed as prey (Supplementary Material). Moreover, the principal prey observed were limited to just four species [the two species of Crepipatella (51%), Aulacomya ater (36%), and Mytilus chilensis (8%)] with three other taxa making up the balance (holothuroid, crustacean, gastropod – each <2%; Figure 6). Regarding the proportion of
FIGURE 5 | Prey availability – Abundance (percent cover) at different depth intervals of common invertebrates that are potential prey of the sea star Cosmasterias lurida in Yerbas Buenas Bay.

TABLE 1 | Abundance (percent cover) of the potential prey of the sea star Cosmasterias lurida in photo-quadrats at depths down to 25 m (no sea stars were observed below this depth).

| Taxa/Depth (m) | 0–5 | 5–10 | 10–15 | 15–20 | 20–25 |
|---------------|-----|------|-------|-------|-------|
| %E | %P | %E | %P | %E | %P | %E | %P | %E | %P |
| Asteroidea   | 0.30 | –  | 0.46 | –  | 0.79 | –  | 0.22 | –  | 0.24 | –  |
| Bivalvia     | 11.05 | 75.73 | 1.16 | 64.10 | 1.70 | 25.00 | 0.19 | 1.73 | 0.02 | –  |
| Gastropoda   | 23.88 | 12.86 | 23.40 | 30.77 | 15.78 | 73.33 | 12.23 | 96.55 | 3.66 | –  |
| Holothuroidea| 0.07 | –  | 2.93 | –  | 4.28 | 1.67 | 5.22 | 1.72 | 3.79 | –  |
| Echinaidea   | 5.42 | –  | 22.32 | –  | 24.06 | –  | 16.52 | –  | 7.55 | –  |
| Crustacea    | 13.15 | 1.41 | 0.02 | 1.28 | 0.05 | –  | 1.07 | –  | 0.26 | –  |
| Polychaeta   | 0.65 | –  | 0.12 | –  | 0.12 | –  | 0.51 | –  | 3.04 | –  |
| Porifera     | –  | –  | 0.07 | –  | 0.73 | –  | 1.28 | –  | 1.41 | –  |
| Polyplacophora| 0.67 | –  | 0.32 | –  | 0.14 | –  | 0.11 | –  | 0.02 | –  |
| Anthozoa     | –  | –  | 0.09 | –  | 0.98 | –  | 3.31 | –  | 2.96 | –  |
| Ascidiae     | –  | –  | –  | –  | 0.20 | –  | 0.01 | –  | –  | –  |
| Brachiopoda  | –  | –  | –  | –  | –  | –  | –  | –  | –  | –  |
| Bryozoa      | –  | –  | –  | –  | –  | –  | –  | –  | –  | –  |
| Opisthobranchia| –   | –  | 0.03 | –  | –  | –  | 0.07 | –  | 0.06 | –  |
| Hydrozoa     | –  | –  | –  | –  | –  | –  | –  | –  | –  | –  |
| Ochrophyta   | 2.70 | –  | –  | –  | –  | –  | –  | –  | 0.02 | –  |
| Rhodophyta   | 6.98 | –  | 9.55 | –  | 4.72 | –  | 9.40 | –  | 21.36 | –  |
| Mud, Sand, Rock, Shell, Cobble/Pebble| 33.24 | –  | 38.72 | –  | 46.15 | –  | 49.35 | –  | 54.98 | –  |
| Shadow       | 1.59 | –  | 0.58 | –  | 0.21 | –  | 0.20 | –  | 0.13 | –  |
| Unidentified | 0.30 | 10.00 | 0.23 | 3.85 | 0.09 | –  | 0.22 | –  | 0.38 | –  |

%E, Percent cover of potential prey; %P, Percentage of each taxa in diet of C. lurida; “–,” Percentage <1%.

the species preyed upon by C. lurida PERMANOVA showed significant differences among depth intervals (Table 3). The posteriori Pair-wise test indicated that the most similar depth intervals were >10–15 and >15–20 dissimilar depth intervals were 0–5 and >15–20 m (Table 3). Those differences are better visualized in the PCO analysis, where the shift in the importance of the major prey items (A. ater and Crepitipatella spp.) shift with depth can be seen (upper panels of Figure 7) as well as the lack of importance of the other mussel (M. chilensis) and other prey items (lower panels of Figure 7). With regards to the size of sea stars [mean disk diameter ± SD: 31.0 ± 4.5 mm and the size of prey sizes (mean length ± SD: 57.0 ± 26.0 mm)] that
they consumed, neither varied significantly with depth, and there were no significant correlations between predator and prey size for the three most commonly selected prey: *A. ater* \( (r = 0.238, n = 99, p > 0.05) \); *M. chilensis* \( (r = -0.383, n = 25, p > 0.05) \); *Crepipatella* spp. \( (r = 0.207, n = 134, p > 0.05) \).

**Prey Electivity**

The electivity \( (C) \) of different prey types changed by depth interval (Table 4 and Figure 8). In shallower depth intervals, the sea star had a high positive electivity for the mussel *A. ater* in depths of 0–15 m, especially in the >5–10-m depth interval \( (C = 0.72) \). It also had a slight but not significant positive electivity for the other mussel, *M. chilensis*, but only in the >5–10-m depth interval. Electivity changed dramatically below 10 m where there was a moderately high electivity for *Crepipatella* spp., which increased with depth \( (C = 0.28 \text{ and } 0.44 \text{ in the } >10–15\text{-m and } >15–20\text{-m depth intervals, respectively}) \). This contrasts sharply with the shallowest depth intervals with the significantly negative electivity value \( (C = -0.33) \) in the 0–5-m interval, which indicated that *Crepipatella* spp. was avoided there, and \( C = 0.062 \) for the >5–10-m depth interval, where this gastropod was not selected by the sea star. With regard to other prey, the index was close to 0 in the shallowest depth interval, indicating that they were consumed to the degree of their availability in the environment, but in all other depth intervals, the indices were negative as these other prey were avoided to

### Table 2 | Results of two-way ANOVA with item prey and depth interval as fixed factors.

| Source                      | DF | SS    | MS    | F     | P     |
|-----------------------------|----|-------|-------|-------|-------|
| Prey item                   | 3  | 604.547 | 201.516 | 23.664 | <0.001 |
| Depth interval              | 3  | 12.797 | 4.266 | 0.501 | 0.683 |
| Prey item × depth interval  | 9  | 652.391 | 72.488 | 8.512 | <0.001 |
| Residuals                   | 48 | 408.75 | 8.516 |       |       |
| Total                       | 63 | 1678.484 | 26.643 |       |       |

### Table 3 | PERMANOVA partitioning and analysis of prey item per depth intervals (4) from Yerbas Buenas Bay, based Bray–Curtis dissimilarities.

| Source                  | df | SS    | MS    | Pseudo-F | P(perm) |
|-------------------------|----|-------|-------|-----------|---------|
| Depth interval          | 3  | 16415.4 | 5471.8 | 7.6       | 0.001   |
| Residual                | 12 | 8688 | 724 | –         | –        |
| Total                   | 15 | 25103.4 | –     | –         | –        |

| Pair-Wise Test | t   | P(perm) | Similarities (%) |
|---------------|-----|---------|------------------|
| Depth         |     |         |                  |
| 0–5 vs. >5–10 | 0.9 | 0.572   | 59.7             |
| 0–5 vs. >10–15| 2.1 | 0.049   | 35.4             |
| 0–5 vs. >15–20| 4.2 | 0.028   | 12.9             |
| >5–10 vs. >10–15| 1.9 | 0.087   | 55.9             |
| >5–10 vs. >15–20| 6.9 | 0.033   | 33.1             |
| >10–15 vs. >15–20| 1.6 | 0.15    | 68.3             |

Pseudo F statistic were calculated for each term using direct analogs to univariate expectations of mean squares (EMS); p-values were obtained using 9999 permutations under a reduced model. Each term is identified as contributing either a fixed or random component to the overall model. The Pair-Wise Test below indicates similarities (%) among depth intervals.
FIGURE 7 | Principal Coordinates Analysis (PCO) showing prey distinctly selected according to depth. Symbols represent different depth intervals (m); ⋆ = 0–5, ▲ = >5–10, ◆ = >10–15, ■ = >15–20. The size of the circles represent the proportion of the main species preyed upon by Cosmasterias lurida.

TABLE 4 | Electivity values (C index) for feeding in the sea star Cosmasterias lurida.

| Depth interval (m) | Prey item     | C index | Chi-square value |
|-------------------|---------------|---------|------------------|
| 0–5               | Crepipatella spp. | −0.33*** | 0.0016           |
|                   | Aulacomya ater  | 0.439*** | 0                |
|                   | Mytilus chilensis | 0.003   | 0.7515           |
|                   | Other prey*     | −0.013  | 0.6228           |
| >5–10             | Crepipatella spp. | 0.062   | 0.3663           |
|                   | Aulacomya ater  | 0.72*** | 0                |
|                   | Mytilus chilensis | 0.165** | 0.0125           |
|                   | Other prey*     | −0.338*** | 0.0002         |
| >10–15            | Crepipatella spp. | 0.28*** | 0.0026           |
|                   | Aulacomya ater  | 0.258*** | 0.0012           |
|                   | Mytilus chilensis | 0.034   | 0.193            |
|                   | Other prey*     | −0.437*** | 0              |
| >15–20            | Crepipatella spp. | 0.44*** | 0.0001           |
|                   | Aulacomya ater  | 0.009   | 0.3537           |
|                   | Mytilus chilensis | −       | −                |
|                   | Other prey*     | −0.474*** | 0              |

Values near 0 indicating no selectivity and values of +1 and −1 indicating absolute preference and avoidance, respectively. No values could be calculated for the depths below 20 m as no sea stars were observed feeding there. * Other prey included all prey observed as the diet of Cosmasterias lurida. ***P < 0.01, **P < 0.05.

DISCUSSION

The impact of any predator on community structure and organization depends on multiple factors, for example, the degree of selectivity (generalist vs. selective predators) when choosing their prey, prey diversity, trophic level of the prey together...
FIGURE 8 | Availability of potential prey (percent cover) in the environment compared to their occurrence in the diet (percentage) of the sea star Cosmasterias lurida in Yerbas Buenas Bay.

with their defense mechanisms or prey recruitment dynamics. Predator behavior is especially relevant because selective predators can have strong indirect effects on competition among prey, producing drastic changes in community structure, especially when having preferences for competitive dominants (Paine et al., 1985; Castilla, 1999; Rettig and Smith, 2021). In many marine ecosystems sea stars are considered to be keystone species (Paine, 1966, 1969; McClintock and Lawrence, 1985; Menge and Sanford, 2013), and indeed, this general ecological concept arose from early studies on sea star ecology (Paine, 1966, 1969). However, this original concept has evolved into a holistic vision and is now used in a more complex conceptual framework to be able to define an entire ecosystem as a “keystone species complex” (Hermosillo-Nunez et al., 2018). In many cases, prey diversity (species and size), system productivity, prey recruitment dynamics, and predator preferences have all been important factors determining their impacts (Menge and Sanford, 2013). Because sea stars prey strongly on bivalves, mussels are often a principal component of the diet of many asteroids, especially in shallow waters (Castilla and Crisp, 1970; Tokeshi, 1991; Tokeshi and Romero, 1995; Gaymer et al., 2001a,b; Gaymer and Himmelman, 2002; Gil and Zaixso, 2008; Lamare et al., 2009). For example, *Heliaster helianthus*, described as a keystone predator in intertidal communities along the central coast of Chile, limits the lower distribution of the mussel *Perumytilus purpuratus* (Lamarck, 1819), which can monopolize the rocky surface when *H. helianthus* is absent (Paine et al., 1985). The dominance of filter-feeding gastropods, the slipper limpets *Crepipatella* spp. (52% of all feeding observations), in the diet would then seem, at first glance, to go against this trend, suggesting that *C. lurida* does not appear to be a predator that specializes on mussels but is instead a generalist. However, the electivity analysis across depth showed clearly that one of the mussel species, *Aulacomya ater*, was preferred in shallower zones. Strikingly, the other mussel species, *Mytilus chilensis*, was never preferentially consumed by this sea star, and this difference could possibly explain why this mussel occurred in higher abundance than the preferred species. Regardless, this pattern goes against the generalization that sea stars invariably prefer feeding on mussels. The subtle differences that might drive the distinction between these two very similar species remain unknown and worthy of future investigation.

The large proportion of gastropod prey in the diet of *C. lurida* was, however, not surprising for several reasons. First, unlike most gastropods, slipper limpets are sessile and thus cannot move to escape predators as observed in many other systems where mobile gastropods are common (McClintock, 1985; Bryan et al., 1997; San-Martin et al., 2009). Second, it was the most abundant substratum occupier at this site and thus was the most available item in the prey field. Therefore, there was a high overall electivity index for this group, especially at greater depths (5–20 m) where
alternative invertebrate prey were either scarce (e.g., mussels) or highly mobile (e.g., sea urchins).

Selective removal of species by sea stars can structure benthic communities (Paine, 1966; Gaymer and Himmelman, 2008; Manzur et al., 2010), and _C. lurida_ may be acting as a keystone predator in Patagonian rocky subtidal ecosystems. Indeed, the dominance of the slipper limpets _Crepipatella_ spp. at our study site (reaching 100% in some photo-quadrats) may rely on the selective predation of _C. lurida_ on the mussel _A.atra_, a potential competitor for space. Manipulative experiments would be needed to establish any such role of this predator. For example, the very low abundance of mussels below depths of 5 m could be due simply to competitive displacement by _Crepipatella_ spp. or by predation from _C. lurida_, which occurred at its highest local densities at depths between 5 and 15 m. Either mechanism could allow the slipper limpets to monopolize the substratum. Alternatively, physiologically restricted could limit mussels to inhabit shallower subtidal environments. However, both _A. ater_ and _M. chilensis_ have been recorded at depths up to 25 m in other localities (Solis and Lozada, 1971; Lorenzen et al., 1979; Cazzaniga, 1990; Zagal et al., 2001), making this latter possibility unlikely.

The decrease in abundance of _Crepipatella_ spp. below a depth of 20 m corresponded to the low abundance of sea stars also observed at these depths, suggesting that it is not predation that is determining the lower limit of these species. However, the spatial distribution and abundance of asteroids often tracks that of their prey (Gaymer and Himmelman, 2002; Himmelman et al., 2005), and the low abundance of all invertebrates suggests that other ecological processes (e.g., low recruitment) are structuring benthic assemblages at deeper depths, which were dominated primarily by octocorals, mainly _Primnoella_ sp. (Garrido, unpublished data), and another calyptraeid gastropod, _Calyptraea aurita_ (Reeve, 1859) (Holtheuer et al., 2018), below depths of 20 m.

The depth-dependent selectivity that we observed in our study contrasts strikingly with the only other detailed study of the feeding ecology of this species (Vásquez and Castilla, 1984). That study, conducted in southern Patagonia, concluded that _C. lurida_ was a generalist, consuming prey species according to their availability in the environment. Their assessment of selectivity was, however, based on a much simpler, non-parametric analysis, that is determining the lower limit of these species. However, the spatial distribution and abundance of asteroids often tracks that of their prey (Gaymer and Himmelman, 2002; Himmelman et al., 2005), and the low abundance of all invertebrates indicates that other ecological processes (e.g., low recruitment) are structuring benthic assemblages at deeper depths, which were dominated primarily by octocorals, mainly _Primnoella_ sp. (Garrido, unpublished data), and another calyptraeid gastropod, _Calyptraea aurita_ (Reeve, 1859) (Holtheuer et al., 2018), below depths of 20 m.

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the diet of *C. lurida* even though they occurred in relatively high abundance at our study site. One explanation is that sea urchins, including one of the species in our study (*Loxechinus albus*), can have effective risk-avoidance behaviors to sea star attacks (Urriago et al., 2011, 2012).

Prey electivity could also change depending on the size of predators and prey (Paine, 1976; Barbeau and Scheibling, 1994). However, we did not find a predator–prey size relationship between *C. lurida* and any of its three primary prey species. Similarly, observations on feeding behavior in another Chilean sea star, *Meyenaster gelatinosus*, did not indicate preferences for any particular size of the sea urchin *L. albus*, the dominant invertebrate species at that study site (Dayton et al., 1977). In contrast, *Leptasterias polaris* (Müller and Troschel, 1842) and *Asterias vulgaris* (Packard, 1863), two common subtidal sea stars in eastern Canada, had differences in their use of prey (species and size of bivalves) that may facilitate their coexistence (Gaymer et al., 2001a). On the Pacific coast of North America, the selection of different sizes of prey (mussels) appears to maximize the energy intake in the competing sea stars *Leptasterias hexactis* (Stimpson, 1862) and *Pisaster ochraceus* (Brandt, 1835) (Menge and Menge, 1974). Thus, the presence of competitors might select for intraspecific differences in size to partition the trophic niche. This is not the case, however, for our study system, because no other predatory sea stars co-occur with *C. lurida* (Garrido, unpublished data).

This study shows the importance of assessing the environmental context to describe the trophic behavior of predators. Laboratory studies about dietary preference indicate that asteroids show preferences for some prey offered, but they did not consider the relative abundance of prey in the environment (Castilla and Crisp, 1970; Castilla, 1972; Rochette et al., 1994; Pratchett, 2007). On the other hand, field studies usually show generalist behaviors regardless of environmental variability. Species that have been considered as generalists are frequently much more selective when the environmental offer is evaluated (Gaymer and Himmelman, 2008). Detailed and realistic knowledge of the trophic habits of species is a critical element for constructing food webs (Martinez, 1993), which, in turn, help to predict effects of perturbations on communities (Pérez-Matuz et al., 2017). In addition, anthropogenic activity, such as coastal urbanization that results in the destruction of habitats, needs to be considered as it can significantly affect sea star populations and their impact on the trophic structure and function of subtidal communities (Chan et al., 2018). In this sense, this study is a key contribution to understanding food web dynamics of the benthic marine communities of northern Patagonia.

**DATA AVAILABILITY STATEMENT**

The original contributions presented in the study are included in the article/Supplementary Material, further inquiries can be directed to the corresponding author.

**AUTHOR CONTRIBUTIONS**

IG: conceptualization, validation, formal analysis, writing, and visualization. LP: conceptualization, investigation, and writing – review and editing. IJ: conceptualization, validation, and writing – review and editing. DS: conceptualization, investigation, validation, formal analysis, and writing. All authors contributed to the article and approved the submitted version.

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**SUPPLEMENTARY MATERIAL**

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fmars.2021.636208/full#supplementary-material

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The remaining authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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