

Feeding ecology of the siluriform *Pimelodella laticeps* Eigenmann, 1917 in a Pampean stream from Argentina

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Abstract: *Pimelodella laticeps* Eigenmann, 1917 is the most abundant siluriform in the El Pescado Stream, located in the depressed region known as pampean region (Buenos Aires, Argentina). To characterize its feeding habits in this environment, we analyzed the dietary contents on seasonal samplings from May 1991 through August 1993. To evaluate the relative contribution of the different dietary components, the index of relative importance (IRI) was used. *Pimelodella laticeps* preferred sectors with aquatic macrophytes, and predated mainly on organisms from the periphytic and benthic communities. The main food was cyclopoid copepods, which taxa usually have littoral or benthic habitats, and only to a lesser extent are components of the plankton community. The planktonic organisms available in the environment were analyzed by the Olmstead-Tukey-test, which diagram indicated that the dominant items were copepods, ostracods, and chironomids. Chydorid cladocerans, harpacticoid copepods, mayfly larvae, and amphipods also became dominant but less frequently. To understand the feeding strategies of *P. laticeps*, the fish diet was compared with the available plankton as analyzed by a nonparametric multivariate analysis (permutation-based one-way ANOSIM). *Pimelodella laticeps* consumed almost no phytoplankton, while the discriminating items from the latter assemblage were identified as having high values in the environment.

Key words: : Feeding habits, gray catfish, dietary plasticity, Neotropical-plains stream, Pampean region

Resumen: Ecología alimentaria del siluriforme *Pimelodella laticeps* Eigenmann 1917, en un arroyo pampeano de la Argentina *Pimelodella laticeps* Eigenmann 1917 es el siluriforme más abundante en el arroyo El Pescado, situado en la zona deprimida en la región pampeana (Buenos Aires, Argentina). Para caracterizar sus hábitos alimentarios se analizó el contenido del tubo digestivo de peces capturados durante muestreos estacionales (desde mayo 1991 hasta agosto 1993). Se calculó el índice de importancia relativa (IRI) para evaluar la contribución de los diferentes componentes de la dieta. *Pimelodella laticeps* prefirió alimentarse en sectores con macrófitas acuáticas, depredando principalmente sobre organismos de las comunidades bentónicas y perifíticas. Su alimento principal fueron los copépodos ciclopoideos, que por lo general tienen hábitats litorales o bentónicos, y solo ocasionalmente habitan el plancton. Los organismos planctónicos disponibles en el ambiente se analizaron mediante la prueba de Olmstead-Tukey, test que indicó como ítems dominantes a los copépodos, ostrácodos y larvas de quironómidos. Los cladóceros Chydoridae, los copépodos Harpacticoida, las larvas de efemerópteros y los anfípodos también fueron dominantes, pero con una frecuencia menor. La dieta de *P. laticeps* se comparó con los organismos planctónicos disponibles y se analizó mediante un análisis multivariado no paramétrico (ANOSIM de una vía basado en permutaciones) a fin de comprender las estrategias de alimentación. Si bien algunos de los ítems discriminantes de las diferencias encontradas fueron del plancton, y estos se identificaron con valores altos en el ambiente, *P. laticeps* no consumió organismos planctónicos de manera significativa.

Palabras clave: Hábitos alimentarios, bagre gris, plasticidad en la dieta, arroyo de llanura neotropical, región pampeana

INTRODUCTION

Pimelodella laticeps Eigenmann, 1917 (Siluriformes, Heptapteridae) (known as the gray catfish) is a Neotropical species distributed in lentic and lotic environments in Bolivia, Paraguay, southern Brazil, and Argentina (Ringuelet *et al.*, 1967). The southern end of the species distribution is the Sierra de la Ventana hills and

the Chasicó Stream (south of the Buenos Aires province; Menni *et al.*, 1988). It was also found in brackish water of the Samborombón Bay (García & Almirón, 1991) and in the shallow lake El Estado (Grosman & Sanzano, 2003).

Low temperature was identified as the most influential condition determining the southern boundary of Neotropical-fish distribution

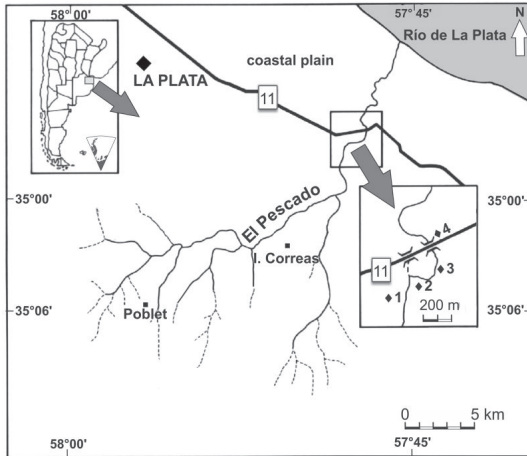


Fig. 1. Sampling sites in El Pescado Stream.

in Argentine freshwater ecosystems, although certain other species have shown better acclimation to colder conditions (Colautti *et al.*, 2015). Baigún *et al.* (2002) attributed the absence of *P. laticeps* in northern Patagonia to its low tolerance to cold temperatures.

Pimelodella laticeps is the most abundant siluriform fish in the El Pescado Stream, where the species inhabits throughout the year. Almirón *et al.* (2000) and López van Oosterom *et al.* (2013) remarked that *P. laticeps* is a permanent species in the El Pescado Stream because of the fish ecologic plasticity facing the marked instability of habitat. Colautti *et al.* (2009) noted that the species is also permanent in the La Chozas Stream; whereas Paracampo (2012) recorded the catfish as a semipermanent species in El Pescado Stream.

El Pescado Stream is located in the ecotone between the subtropical and the Patagonian fish fauna regions. It is inhabited by 55 fish species, with *P. laticeps* being the most abundant from the remote sites down to the mouth of the stream (Almirón *et al.* 2000). Macluf *et al.* (1998) mentioned that the zooplankton found in this stream offered high-quality food for fish, and that *P. laticeps* prefers to feed on the microcrustacean assemblage.

Several *P. laticeps*–population estimations have been made in pampean shallow lakes (Freyre *et al.*, 1966, 1967; Alaimo & Freyre, 1969). Some specific aspects of the *P. laticeps* biology are quite well known—such as the morphometry (Freyre & Mollo, 1987), the structure of the pectoral spines (Bisbal & Gómez, 1986), and the concentration of dissolved oxygen lethal to the fish (Gómez, 1993). In relation to water quality, an investigation on

the fish assemblage from the River Suquia concluded that the genus *Pimelodella* lives only at sites with good water quality (Hued & Bistoni, 2007). After the first studies on diet of *P. laticeps* in pampean environments (Destefanis & Freyre, 1972; Escalante, 1984), the interest in characterizing the eating habits of the gray catfish increased recently (Grosman *et al.*, 2002; Freyre *et al.*, 2003; Diovisalvi *et al.*, 2010; Fernández *et al.*, 2012; López van Oosterom *et al.*, 2013).

The objectives of this study were: to analyze the seasonal variations of the feeding habits of *P. laticeps* in the El Pescado Stream; to compare those results with the available food; and to describe the species' environmental circumstances when the stream was not yet affected by human activity, in order to provide a tool for assessing the current condition of the environment, both there and elsewhere. To fulfill these objectives, we took into account the seasonal analyses of the plankton in the El Pescado Stream performed in the same decade by Macluf *et al.* (1998).

MATERIALS AND METHODS

Study area

El Pescado Stream is located in a depressed area known as Pampean region, within the lower Río de la Plata basin in the Buenos Aires province, Argentina (35° 55' S, 57° 45' W; Frenguelli, 1950) (Fig. 1). This watershed is the northernmost of the four basins in the large plain covering most of the eastern central latitude of Argentina (from 34° 00' to 38° 30' S). This stream flows in an area at the southern border of the Subtropical Zoogeographic Dominion (Ringuélet, 1961; Almirón *et al.*, 2000), and after 36 km is channeled into the coastal plain, from where the channel becomes directed to the Río de la Plata estuary. El Pescado has a basin of 400 km², a slope of 0.6 m/km and develops as a plain stream with a variable flooding frequency; its input coming from groundwater and from numerous tributaries. The surrounding basin is an agricultural area with crops of soybean, corn, wheat, and alfalfa and some livestock.

Four sampling sites with different topographies (Fig. 1) were selected as cited in Almirón *et al.* (2000). The samples for that study and the present one were all collected as indicated here, in 1991 through 1993. Those authors subsequently published the data on the relative abundance of the fish assemblage; the results we present here, from the same original samplings, constitute a retrospective evaluation of the ecologic situation

at that time from the standpoint of the feeding habits of *P. laticeps*, which was only captured at sites 1, 2 and 3, where Almirón *et al.* (2000) described the presence of macrophytes during the same sampling period. *Ceratophyllum demersum*, *Potamogeton striatus*, and *Schoenoplectus californicus* were present during the entire sampling period. Submerged and swamp macrophytes -such as *Alternanthera philoxeroides*, *Hydrocotyle ranunculoides*, *Ludwigia* sp., *Myriophyllum aquaticum*, *M. quitense*, and *Polygonum* sp.- were present during the early spring through the summer, whereas the floating vegetation -such as *Azolla filiculoides*, *Lemna* sp., *Limnobium laevigatum*, *Spirodella intermedia*, and *Ricciocarpus natans*- developed in the later spring.

This evaluation was done within the mentioned environmental context, but with statistical methods for comparative analysis presently available.

Samplings and data analysis

Samplings were performed seasonally from May 1991 through August 1993, except in May 1992 and May 1993 because of floodings. The fish were caught with a seine net (15 m long x 1 m high, with a cod end 1.5 m long, mesh 5–10 mm). The catch effort had the same intensity at each site. The specimens were fixed in situ with 10% formaldehyde and measured (standard length, SL), weighed, and sexed in the laboratory. After dissection of the digestive tracts, the contents were removed and placed in 5% formaldehyde. The minimum sample size was calculated under the assumption of a Poisson distribution as follows: $n = (200cv/r)^2 = (200/r)^2 x^{-1}$, where n is a sample size required for a Poisson variable; r is a desired relative error (as a percent); cv is the coefficient of variation = $1/x$ (Krebs, 1989). The dietary contents were fully analyzed with respect to quality and quantity to the lowest possible categories by observation in counting chambers under the stereomicroscope and optical microscope. The number, weight, and volume of the dietary items were recorded. The volume of the microcrustaceans was obtained by liquid displacement (Escalante, 1982) and the algal volume determined as biovolume (Reynolds, 1984; Lewis, 1976; Wetzel & Likens, 1991; García de Emiliani, 1993). To evaluate the significance of the different dietary components, the index of relative importance (IRI; Pinkas *et al.*, 1971) was used: $IRI = FO(N+V)$, which parameter is a function of the percent frequency of occurrence (FO) and the numerical (N) and volumetric (V) percentages of the various items consumed in the diet.

The relative abundance of each item was obtained and the feeding habits compared between the different seasons through analysis of the gut-content-abundance values by means of a nonparametric multivariate analysis (a permutation-based one-way ANOSIM; Clarke & Warwick, 2001). In this way, a similarity-percentage analysis (SIMPER) was used to identify the so-called discriminating items that most highly contributed to accounting for the observed similarity (or dissimilarity) between samples. This method uses the Bray-Curtis measure of similarity by comparing, in succession, each sample in one group with every sample in another set. Finally, in order to understand the feeding strategies of *P. laticeps*, the fish's diet was compared with the plankton availability recorded by Macluf *et al.* (1998). This plankton community was analyzed by means of the Olmstead-Tukey method (Sokal & Rolf, 1979), a nonparametric association test that allows to determine dominant, frequent, rare, and common items. This information was also used to compare by multivariate analysis (ANOSIM) between the plankton available in the environment and the consumed items of this assemblage. SIMPER was also used to identify the discriminating items accounting for this similarity (or dissimilarity). Before both analyses, the rare species were discarded from the general matrix, and the data were then transformed to $\log(x + 1)$ to reduce the contribution of the most highly abundant species. For these comparisons, fish and plankton samples were obtained at the same time (as explained in reference to the study area above).

RESULTS

Sampling results

The specimens of *P. laticeps* were captured only at sampling sites 1, 2 and 3 with the presence of *S. californicus* and thus with a well-developed epiphytic community. The capture abundances were 228 (Site 1), 349 (Site 2), and 24 (Site 3) individuals. One hundred thirty fish were examined for diet analyses; including 15 juveniles (SL: 29.3–41.5 mm; weight: 0.3–1 g), 62 females (SL: 39.4–78.2, weight: 0.9–8.2 g), and 53 males (SL 35.3–62 mm; weight: 0.6–3.6 g). The minimum sampling was 73 stomachs.

Fish diet

The diet of *P. laticeps* resulted in 1,470 consumed prey from 52 taxa. Tables 1a and 1b illustrate the seasonal variations in the composition of that diet and provide taxonomic details of the items therein.

The year-round diet of *P. laticeps* contained mainly cyclopoid copepods, present in 18% of the examined stomachs; with *Ectocyclops phaleratus*, *Macrocyclus albidus*, and *Tropocyclops prasinus* -typical periphytic or benthic species- along with *Acanthocyclops robustus* -the only planktonic species- being recorded.

The benthic organisms were the second most highly consumed. The chironomid larvae appeared in 15% of the analyzed stomachs, while the mayfly nymphs (Ephemeroptera) were present in 10% and ostracods in 9%, followed by cladocerans in 7%. The Chydoridae cladocerans, which are common in the periphyton and sparsely represented in the plankton, were the most abundant in the diet. Moreover, the harpacticoid copepods were present in 6% of the stomachs along with the amphipods. The chrysophytes were found in 5% of the stomachs in winter 1993, but they were absent in the planktonic samples. The insects and macrophyte fragments both remained at a 4% frequency. Cyanobacteria, chlorophytes, euglenophytes, pyrrophytes, charophytes, rotifers, nematodes, amphipods, acari, tardigrades, molluscs, and fish remains were present at a frequency of fewer than 2% of the analyzed stomachs. It is noteworthy that cyanobacteria, chlorophytes, euglenophytes, pyrrophytes, and rotifers were however present in the planktonic samples. Table 2 shows the percent frequency of occurrence, number and volume, and the index of relative importance (IRI) of the main groups found in the diet of *P. laticeps*.

The multivariate analysis showed that the diet composition changed during the different sampling dates (ANOSIM, $R = 0.45$ $p < 0.05$); and when the data were compared according to seasonality, significant differences between seasons were observed (ANOSIM, $R = 0.27$ $p < 0.05$). Table 3 lists the percentages of dissimilarity between seasons (SIMPER). Data in this table indicate that cyclopoid copepods, chironomids, and ostracods were the responsible for the differences in the diet between seasons and that cyclopoid copepods were the most highly consumed item; this statement is especially true for autumn when this season was compared with the others and for summer when the latter was compared with spring and winter.

Among the 1,470 total preys consumed during the year, cyclopoid copepods (copepodites and adults) dominated the diet during autumn, whereas calanoid copepod nauplii were abundant in the plankton in the same season. The latter appear in 100% of the stomachs analyzed, rep-

resenting 87% of the food consumed. The second most frequent food item was the ostracods and chironomid larvae, with those two taxa appearing at a somewhat lower proportion. Other prey had a lower representation in the diet (1%). The algae were not consumed during this season, except for *Surirella ovalis* and *Oedogonium* sp.

During winter, benthic organisms were the principal components of the diet. The diatoms *Aulacoseira distans*, *Melosira varians*, *Navicula* sp., and *Ulnaria ulna* were the most frequent items, those being registered in 43% of the examined stomachs. The cyclopoid copepods *E. phaleratus*, *M. albidus*, and *T. prasinus*; the chydoridcladoceran *Coronatella rectangula*, chironomid larvae; and the ephemeropteran nymphs followed in frequency.

The highest number of total items consumed was recorded in spring ($N = 43$). In this season the prey with the highest absolute frequency were the chironomids and cyclopoid copepods (65.6% each), followed by amphipods (53.1%), chydoridcladocerans (46.84%), and harpacticoid copepods (40.6%) in the analyzed stomachs (Table 1b). Furthermore, a wide variety of algae including diatoms, chlorophytes, euglenophytes, pyrrophytes, and charophytes were also found (Table 1a).

In summer the fish also preferred feeding on cyclopoid copepods, chironomid larvae, and ostracods; with those respective taxa being found in 69%, 55%, and 39% of the stomachs analyzed. Harpacticoid copepods, represented at a frequency of 22%, were also often consumed (Table 1b).

Food availability

The plankton community in El Pescado Stream offered a wide range of available food comprising 103 phytoplanktonic species (cyanobacteria, chrysophytes, chlorophytes, euglenophytes, and pyrrophytes) and 152 zooplanktonic species (testate amoebae, ciliates, rotifers, copepods, and cladocerans) (Macluf *et al.* 1998).

In the Olmstead-Tukey diagrams (Figs. 2 and 3), the dominant dietary items were copepods, ostracods, and chironomids. Chydorids, harpacticoids, mayfly nymphs (Ephemeroptera), and amphipods were also highly prevalent, but somewhat less frequent. Among the occasional available food items were algae at high frequencies (chlorophytes and diatoms), with *Daphnia* sp., the remains of macrophytes and insects (dipteran adults and larvae), scales, and acari (not shown) being less frequent.

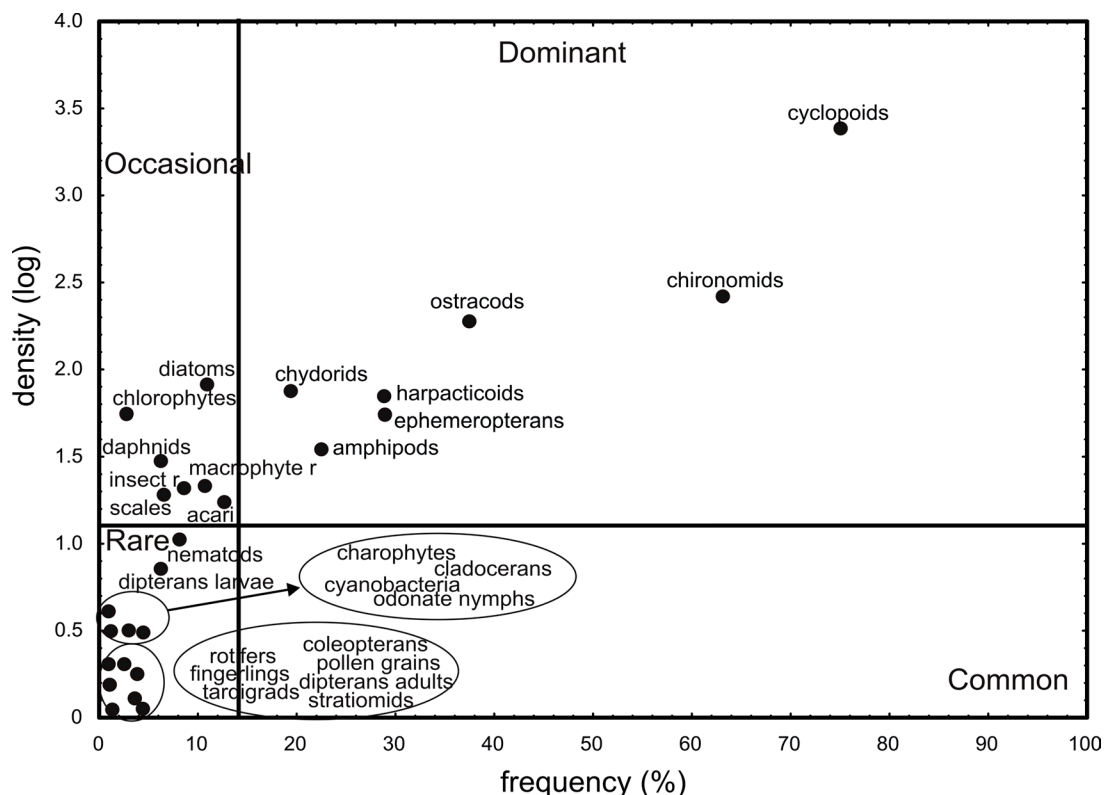


Fig. 2. Olmstead-Tukey diagram showing the distribution and abundance of the dietary items of *Pimelodella laticeps* expressed as taxonomic groups when recognizable (otherwise stated as remains or materials). The items in the occasional and rare sectors of the graph are indicated by abbreviations. In the figure, the logarithm of the item density is plotted on the ordinate as a function of the percent frequency on the abscissa. Those taxa found at both a low frequency and density are arbitrarily classified as rare; at a high frequency but low density as common; at a low frequency but high density as occasional; and at both a high frequency and density as dominant.

Comparison between fish diet and food availability

The relative abundances of phytoplankton and zooplankton found in the environment were different from those in the diet (ANOSIM, $R = 0.87$ $p < 0.05$ and $R = 0.99$ $p < 0.05$, respectively). *Pimelodella laticeps* consumed almost no phytoplankton, while the discriminating items from this assemblage were identified as having high values in the environment (Table 4). The dissimilarity between the zooplankton taxa available and those consumed was also high, with a smaller number of items indicated as discriminating because of their high values in the environment but being almost absent in the diet (rotifers and nauplii larvae) and the cyclopoid copepods for having high values in the diet (Table 4).

DISCUSSION

In the present study, *P. laticeps* was captured

only at sites with dense mats of *Schoenoplectus californicus* covering the shores throughout the year; which macrophytes likely provide refuge and food, even during floods. The relationships between fish and submerged plants in warmer climatic regimes still remains incompletely studied (Jeppesen *et al.*, 2005); and according to some authors, fish communities associated with plants appear to be characterized by a high proportion of omnivorous species (Winemiller, 1990; Branco *et al.*, 1997) that, for their part, have different effects on other trophic levels than those predicted by the trophic-cascade theory (Lazzaro, 1997; Pace *et al.*, 1999).

Our analysis indicated that in El Pescado Stream *P. laticeps* mainly predated upon organisms from the periphytic and benthic communities. The main food was cyclopoid copepods, which taxon usually has littoral or benthic habitats and, to a lesser extent, is also a part of the plankton community (Reid, 1985). Among these

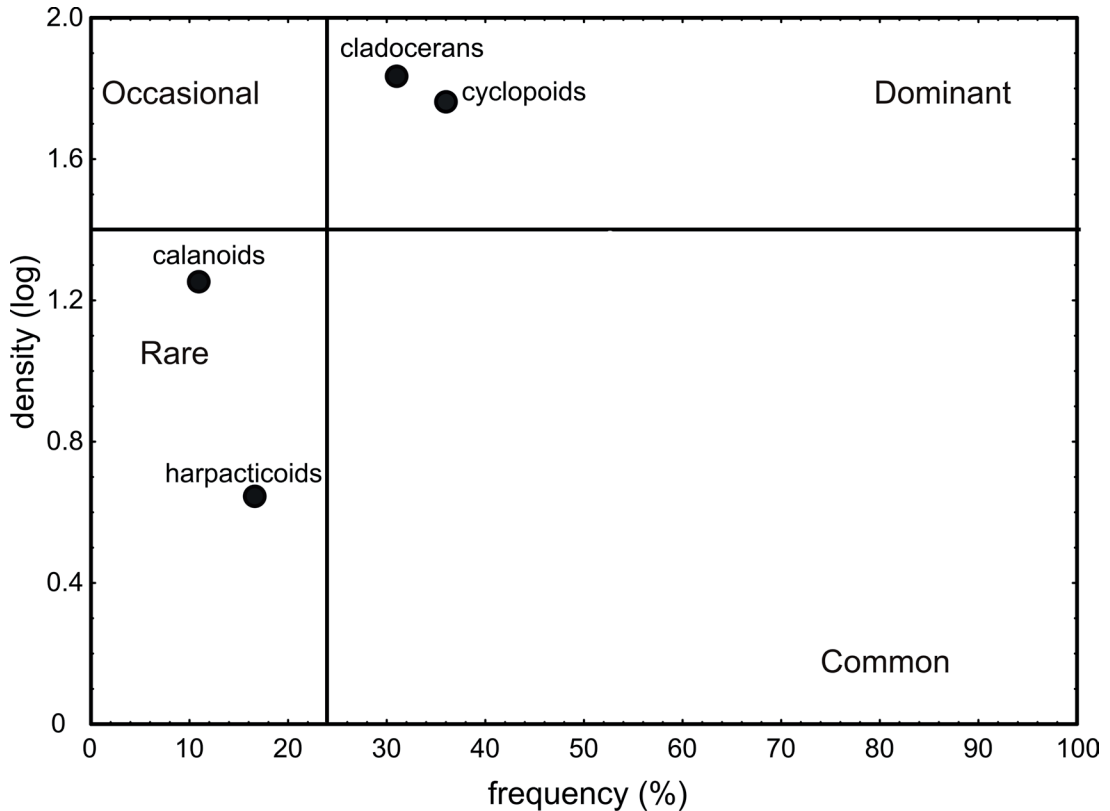


Fig. 3. Olmstead-Tukey diagram showing the offer of microcrustacean zooplankton in El Pescado Stream. For the axes and definition of sectors cf. legend of Fig. 2.

species, *Acanthocyclops robustus* lives mainly in the limnetic zones of large ponds and lakes (Dussart & Fernando 1990); *Ectocyclops phaleratus* is an uncommon benthic species, inhabiting mainly the littoral zone and only occasionally the water column (Dussart, 1969; Reid, 1985); *Macrocyclus albidus* lives among macrophytes, reaching its maximum development during summer (Pennak 1966); and *Tropocyclops prasinus* is a common species within the periphytic and benthic communities (Dussart, 1969), in warm waters, tolerating brackish water, and rarely behaving as planktonic (Margalef, 1983). Chironomid larvae and mayfly nymphs, which taxa are common in lotic and lentic Neotropical environments where the food availability is ample (Marchese, 1984; Kaisin, 1989; López van Oosterom *et al.*, 2013), were found in the diet of *P. laticeps* at high frequencies during spring.

Ostracods were another major item recorded in the diet. These crustaceans increased their relative abundance from spring to summer and then autumn, but decreased in winter. These diet variations were coincident with the seasonal

fluctuations in ostracod abundance, and thus availability, in the environment; as had been observed in other temperate streams within the area (Sampóns, 1988).

The diet of the specimens of *P. laticeps* captured in El Pescado Stream was similar to that from La Choza Stream (López van Oosterom *et al.*, 2013) as regards invertebrate larvae, probably in relation with the presence of macrophytes. In contrast, some differences arose when comparing main food items. Cyclopoid copepods and chironomid larvae constituted the main food of *P. laticeps* in El Pescado, while mayfly larvae and ostracods were found to serve as complementary food items, as had been pointed out in La Choza Stream by López van Oosterom *et al.* (2013). Cyclopoid copepods are widespread and play a major role in the trophic dynamics of freshwater ecosystems (Hopp *et al.*, 1997). Grossman *et al.* (2002) mentioned the degree of relevance of planktonic microcrustaceans in feeding the fish of the shallow lake San Miguel del Monte, even in clearly benthic species such as the siluriforms *P. laticeps* and *Corydoras paleatus* (Jenyns, 1842).

Table 1a. Absolute frequency (AF), relative frequency (RF) and density (N) of vegetable items in the diet composition of *Pimelodella laticeps* in each season.

	AUTUMN			WINTER			SPRING			SUMMER		
	AF	RF	N	AF	RF	N	AF	RF	N	AF	RF	N
CYANOPHYTA												
<i>Anabaenopsis</i> sp.							3.12	0.19	1			
<i>Oscillatoria</i> sp.				4.35	1.60	3						
CHRYSOPHYTA												
<i>Actinopterychus</i> sp.										1.96	0.07	1
<i>A. undulatus</i>							3.12	0.19	1			
<i>Amphora</i> sp.										1.96	0.07	1
<i>Denticula</i> sp.							3.12	0.19	1			
<i>Eunotia</i> sp.				4.35	1.60	1						
<i>Gomphonema</i> sp.							3.12	0.19	1			
<i>G. herculeana</i>							3.12	0.19	2			
<i>Aulacoseira distans</i>				8.69	11.23	21						
<i>Melosira varians</i>				8.69	2.67	5	3.12	0.19	1			
<i>Navicula</i> sp.				4.35	0.53	1	3.12	0.19	1			
<i>Surirella ovalis</i>	3.84	0.05	1				3.12	0.19	1	1.96	0.07	1
<i>Ulnaria ulna</i>				8.69	24.63	45	6.25	0.39	2			
CHLOROPHYTA												
<i>Closterium</i> sp.							6.25	0.78	4			
<i>Oedogonium</i> sp.	3.84	0.05	1							1.96	0.07	1
<i>Pediastrum boryanum</i>							3.12	0.39	2			
<i>P. duplex</i>							3.12	0.39	2			
<i>Scenedesmus quadricauda</i>							6.25	0.39	2			
<i>S. spinosus</i>							3.12	0.39	2			
<i>Spirogyra</i> sp.							3.12	1.4	10			
EUGLENOPHYTA												
<i>Lepocinclis</i> sp.							6.25	0.78	4			
<i>Phacus</i> sp.							3.12	0.19	1			
<i>P. longicauda</i>							3.12	0.19	1			
CHAROPHYTA												
							3.12	0.78	4			
PYRROPHYTA												
<i>Peridinium</i> sp.							3.12	0.19	1			
MACROPHYTES												
remains	3.84	0.05	1	17.40	5.35	10	15.60	4.27	22	11.80	1.19	18
pollen grains				4.35	0.53	1	3.12	0.58	3			

Planktonic pyrrhophytes and diatoms were rarely predated by *P. laticeps*. The low intake of planktonic items during summer could be attributable to the low water levels that make the boundaries between communities imprecise (Escalante, 1982, 1983; Menni, 2004). Nevertheless, several algal groups appearing in winter and spring were not present in the diet.

A comparison between available and consumed food indicated a high preference of *P. laticeps* for cyclopoid copepods, which species are eurytopic and commonly occur in the periphyton or benthos. The periphytic cyclopoids found in the

diet were also present in the planktonic samples obtained at the macrophyte-development sites. *Acanthocyclops robustus* was recorded in the digestive tract but not in the plankton samples during the studied period.

Although a marked preference for copepods and a lower opting for cladocerans were observed, *P. laticeps* showed a notable degree of food flexibility as the fish's diet changed according to food availability and in response to environment alterations. The choice of food became adapted not only to availability in the environment throughout seasons, as had been found in

Table 1b. Absolute frequency (AF), relative frequency (RF) and density (N) of animal items in the diet composition of *Pimelodella laticeps* in each season.

	AUTUMN			WINTER			SPRING			SUMMER		
	AF	RF	N	AF	RF	N	AF	RF	N	AF	RF	N
ROTIFERA												
<i>Filinia longiseta</i>							3.12	0.78	4	0.98	0.65	1
<i>Keratella americana</i>							3.12	0.19	1	0.98	0.65	1
unidentified remains	3.84	0.05	1									
COPEPODA												
Cyclopoida (unidentified)	100	86.7	1,610	26.1	11.8	22	62.5	13.6	70	58.9	64.5	98
<i>Acanthocyclops robustus</i>	3.84	1.67	31							1.96	0.73	11
<i>Ectocyclops phaleratus</i>							3.12	0.78	4			
<i>Macrocyclops albidus</i>										5.88	10.3	156
<i>Tropocyclops prasinus</i>										1.96	0.99	15
Harpacticoida	19.3	0.38	7	8.69	1.6	3	40.6	6.4	33	21.6	1.92	29
CLADOCERA												
Cladocera fragments				13.0	3.74	7	3.12	0.39	2	3.92	0.13	2
Chydoridae (unidentified)	3.84	0.32	6	13.0	7.49	14	40.6	11.8	61	7.84	0.26	4
<i>Allona pulchella</i>							3.12	0.78	4			
<i>A. rectangula</i>							3.12	0.78	4			
Daphnidae (unidentified)										9.8	1.06	16
<i>Daphnia</i> sp.				4.35	1.6	3				1.96	0.07	1
OSTRACODA	57.7	6.19	115				31.25	4.27	22	39.2	6.68	101
AMPHIPODA												
<i>Hyalella curvispina</i>	7.64	0.1	2	17.4	3.74	7	53.1	6.21	62	7.84	0.4	6
ACARINA	15.4	0.21	4				12.5	1.36	7	13.7	0.46	7
TARDIGRADA				4.35	0.53	1				1.96	0.07	1
INSECTA												
Diptera (unidentified larvae)	3.84	0.05	1	8.69	1.07	2	3.12	0.19	1	7.84	0.4	6
Chironomidae (larvae)	77	2.96	55	26.1	11.2	21	65.6	26.8	138	54.9	5.49	83
Stratiomyidae (larvae)	3.84	0.05	1									
Coleoptera				4.35	0.53	1	9.37	0.58	3			
Odonata (fragments)	3.84	0.05	1	4.35	0.53	1	6.25	0.38	2			
Zygoptera (nymphs)	3.84	0.05	1							1.96	0.13	2
Ephemeroptera (unidentified)	38.5	0.59	11	26.1	6.42	12	37.5	3.69	19	31.4	1.45	22
<i>Hexagenia</i> sp.							9.37	0.57	3			
Insect fragments	7.69	0.1	2	13.0	1.6	3	18.8	6.4	33	13.7	1.12	17
MOLLUSCA												
<i>Heleobia</i> sp.				4.35	1.6	1						
PISCES							3.12	0.19	1	1.69	0.07	1
<i>Cnesterodon decemmaculatus</i> scales	3.84	0.16	3	8.69	1.07	2				17.7	1.72	26
NEMATODA	11.5	0.21	4				6.25	0.38	2	9.8	0.46	7

Tabla 2: Percent frequency of occurrence (FO), number (N) and volume (V), and the index of relative importance IRI = FO (N+V) of the main groups found in the diet of *Pimelodella laticeps*.

	FO	N	V	IRI
Copepods	30.83	77.61	14.50	2,839.75
Chironomid larvae	19.69	7.63	53.43	1,202.27
Ephemeropterans	12.66	1.75	22.80	310.80
Ostracods	11.92	6.23	1.30	89.76
Amphipods	6.99	1.23	7.69	62.35
Algae	10.10	3.15	0.05	32.32
Cladocerans	8.81	3.25	0.14	29.86

Table 3: Percentages of dissimilarity in diet composition of *Pimelodella laticeps* among seasons, and items found as discriminants of those differences indicating in which season dominated in relative abundance.

	Autumn	Winter	Spring	Summer
Autumn	0			
	80.00			
Winter	cyclopoids & chironomids (> in autumn)	0		
	71.82			
Spring	cyclopoids & ostracods (> in autumn) chironomids (> in spring)	chironomids & cyclopoids (> in spring)	0	
	47.67			
Summer	cyclopoids & chironomids (> in autumn) ostracods (> in summer)	81.16 cyclopoids & ostracods (> in summer)	76.14 cyclopoids & ostracods (> in summer) chironomids (> in spring)	0

Table 4. Percentages of similarity for phytoplankton and zooplankton as source of food available and consumed by *Pimelodella laticeps*, percentages of dissimilarity between availability and diet, and items found as discriminants of those differences for having high availability or for being highly consumed.

	Phytoplankton	Zooplankton
Similarity in the availability (%)	26	68.5
Similarity in the diet (%)	71.6	58
Dissimilarity between availability and diet (%)	89.1	86.9
Discriminant items for having high availability	Diatoms, chlorophytes and euglenophytes	Rotifers and nauplii larvae
Discriminant items for being highly consumed	None	Cyclopoid copepods

other species (Townsend & Winfield, 1985), but also to floodings occurring in El Pescado Stream during both sampled autumns (Almirón *et al.*, 2000). These environmental disruptions were reflected later in the diet by the presence of plant remains that had been washed into the stream, a situation that was also observed in cichlid species (Jiménez-Badillo & Nepita-Villanueva, 2000).

Food availability seems to be in some way a determining factor of food preference, since fish diet generally is a reflection of the abundance of food items found in the environment (Jiménez-Badillo & Nepita-Villanueva, 2000).

Aquatic vegetation provides a high diversity of habitats in which the organisms that are food

for predator fish develop (Giorgi *et al.*, 2005). Seasonal changes in a fish's diet are generally the result of food availability (Wootton, 1998; Zavala-Camin, 1996). Wolf *et al.* (1988) found that certain groups of insects exhibit a higher abundance during the months of highest precipitation, thus offering a wider range of prey. In contrast, periods of drought were characterized as intervals of scarcity or low supply of both food and habitat resources (Lowe-McConnell, 1987; Wootton, 1998). Resource diminution under such conditions is characterized by a decrease in the total available water surface, which shrinkage increases competitive interactions while decreasing the allochthonous food availability, mainly

in the form of insects (Wolf *et al.*, 1988; Zavala-Camin, 1996).

Destefanis & Freyre (1972) considered that the feeding strategy of *P. laticeps* was mainly planktivorous and that the fish occasionally fed on benthic organisms without significant alimentary preferences. Their results, however, were referred to a small sampling from a shallow pampean lake. In contrast, other authors have placed this catfish species among the micro- and meso-animalivorous fish, among those commonly dwelling in waters with abundant vegetation (Ringuelet, 1975).

Unlike the findings with other members of the family Heptapteridae living in highland environments (Haro *et al.*, 2001) or in rivers and streams with forest galleries (Araujo Lima *et al.*, 1995), allochthonous material was not observed in the diet of *P. laticeps*.

Differing types of feeding behavior were recorded in other species of the genus *Pimelodella*. For example, unlike *P. laticeps*, *P. Gracilis* Valenciennes, 1835 preferred benthic, planktonic, and pleustonic organisms -such as insect larvae, copepods, and rotifers; but not filamentous algae- thus indicating that this species is related to those three communities (Oliveros 1980). Other species in the family -such as *Pimelodus maculatus* (Lacépède, 1803) (Bonetto *et al.*, 1963; Nomura *et al.*, 1972; Alonso, 1978), *Pimelodella hasemani* Eigenmann, 1917 and *Rhamdia sebae* (Valenciennes, 1840) (Saul, 1975), *Pimelodella chagresi* (Steindachner, 1877) and *Rhamdia wagneri* (Günther, 1868) (Angermeier & Karr, 1984), and *Rhamdia cf. hilarii* and *Rhamdella minuta* (Lütken, 1875) (Uieda, 1983)- are considered omnivorous. In most of those species diet, aquatic invertebrates and especially insects (e. g., mayfly nymphs, chironomid larvae, trichopterans, and larval and adult coleopterans) dominate.

The diet of *Pimelodella australis* Eigenmann, 1917 was studied by González-Bergonzoni (2011) and Masdeu (2011), who classified it two possible functional groups: (1) omnivorous-benthivorous with inclusion of zooplankton, and (2) omnivorous-benthivorous with a herbivorous tendency. Teixeira *et al.* (2009) also placed *P. australis* in the omnivore-benthic-planktivorous trophic group residing in temperate and subtropical lakes.

In view of all the above considerations, *P. laticeps* appears to be a species with a high degree of dietary plasticity; as organism's trophic position depends of the available energy sources and the distribution of resources models its ecologic

niche (Odum & Barrett, 2005). The diet is a key determinant that provides information on the interrelationships between foraging behavior, distribution, and physiologic needs along with the abundance of food. Generalist species feed upon a broad spectrum of food in terms of the number of species and their microhabitats, with no marked preference for a particular food source; whereas specialists maintain a diet restricted to a relatively small number of food items (Gerking, 1994). The occurrence of generalist or specialist species in a particular habitat appears to be influenced mainly by the dynamics of their food resources. A predominance of generalist species inhabiting rivers has been recognized because of the ephemeral character of that habitat (Lowe McConnell, 1987; Araujo Lima *et al.*, 1995). A large number of species have the ability to change their eating habits in response to seasonal and/or spatial variations in the availability of food. This flexibility means that those species have the ability to take advantage of a given food source at different times or places (Gerking, 1994; Chemes *et al.*, 2010). In conclusion, because at the time of this study this area had been proposed as a provincial natural reserve, when that environment was in a relatively pristine state, the results obtained by this investigation can be regarded as a critically relevant tool for pursuing the objective of studying the El Pescado Stream more extensively now, when its condition is no longer pristine. The information obtained on the *P. laticeps* diet and the species' response to food availability and habitat distribution could be used in future studies evaluating the change in environmental status that characterizes this and other related lotic environments -some of which may still have remained pristine- within the pampean region of Argentina.

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