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## Aquatic Diptera assemblages in four sympatric *Eryngium* (Apiaceae) phytotelmata in flowering and senescent times

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### ABSTRACT

This study addresses how Diptera assemblages differ in four sympatric phytotelmata. The *Eryngium* species retain water in their axils forming a phytotelm and this situation leads to the speculation that dipteran assemblages may differ, mainly when the plants begin to grow and retain more water in their axils, and when becoming senescent. Diptera from *Eryngium horridum*, *Eryngium stenophyllum*, *Eryngium* aff. *serra* and *Eryngium elegans* were sampled. This study revealed that the *Eryngium* spp. are used by Diptera differentially and their populations show seasonal changes, with peaks in the autumn. Diptera were represented by immature stages belonging to six aquatic and seven semi-aquatic families. The most important were Chironomidae, Culicidae, Perisclididae and Ceratopogonidae. Two species of Chironomids were the most abundant. Culicidae, Perisclididae and Ceratopogonidae were more selective than Chironomidae. Saprophages were the most abundant where the fine detritus feeders were the dominant group. Perisclididae was the main free-swimming predator.

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seasonality; Perisclididae

### Introduction

Phytotelmata are puddles retained in terrestrial and epiphytic plants, which harbour many species of invertebrates and small vertebrates. The most common and studied phytotelmata are tree holes, bamboo internodes, leaf axils of Bromeliaceae and pitcher-plants (Frank and Lounibos 1983); however, it is known that more than 1500 species of plants provide suitable habitats where aquatic organisms develop (Fish 1983).

*Eryngium* L. (Apiaceae) is a cosmopolitan genus of perennial, rhizomatous herbs and terrestrial plants that includes more than 200 species in temperate and tropical regions, of which 31 are known in Argentina (Martínez 2005). However, few species capture and retain water in their leaf axils. Despite the fact that *Eryngium* plants and bromeliads have a similar physiognomy, their phytotelmata are simpler, lacking the semi-central and central cavities. Water in external cavities is ephemeral and makes favourable habitats for semi-aquatic macroinvertebrates. The real cavity containing water for a long time is limited to axils of internal leaves (Campos 2010).

The aquatic fauna occurring in axils of several species of *Eryngium* from Argentina (Vucetich and Rossi 1980; Campos 2010; Campos and Fernández 2011) and Brazil (Machado 1976) has been inventoried. The taxonomic composition of aquatic and semi-aquatic organisms at the ordinal level revealed that Diptera were the best represented in the *Eryngium* communities. Nine aquatic and six semi-aquatic families have been reported (Campos 2010 appendix). However, only Culicidae and Chironomidae have been studied; addressing issues such as selective oviposition of *Culex castroi* Casal and García, (Balseiro 1983); seasonality and population dynamics of three species of *Culex* in *Eryngium pandanifolium* Cham. and Schlecht and *Eryngium cabreræ* Pontiroli (Campos and Lounibos 1999); epizootiology of *Amblyospora camposi* Micieli, García and Becnel (Microsporidian) in *Culex renatoi* Lane and Ramalho (Micieli et al. 2007), mosquito species richness in *E. pandanifolium*, *Eryngium serra* Cham. and Schltld. and *Eryngium eburneum* Decne (Albicócco et al. 2011), and seasonal prevalence of Harpellales (Zygomycota: Trichomycetes) from chironomid larvae (Siri et al. 2008).

The *Eryngium* species are perennial plants with an annual cycle that wastes the water retained in their axils, when the plant goes into senescence after flowering at the end of the autumn. This situation led to speculation that the dipteran communities inhabiting them may differ in richness of species and abundance, mainly between the spring, when the plants begin to grow and retain more water in their axils, and autumn, when the plants become senescent.

The present study aims to compare the assembly of dipterans, the main taxon that develops, in four sympatric species of *Eryngium* during temperate seasons, coinciding with the time of flowering and senescence of the host plants. The sample is designed to find most dipteran species that inhabit the leaf axils during the wet spring and dry autumn.

## Material and methods

### *Study area and phytotelmata species*

Field studies were conducted in 'Ernesto Tornquist' Provincial Park, Sierra de la Ventana (38°10'S, 62°8' W), situated in the southeast of the Austral Pampean biogeographic district in Buenos Aires Province, Argentina (Cabrera and Willink 1980). This district is characterized by a plain with extensive pastures and a mountain range 170 km long with a maximum height of 1243 m above sea level, oriented from northwest to southeast (Harrington 1947). The climate of the mountain range is wet-subwet with a slight deficit of water (Burgos and Vidal 1951). The main rains occur during spring-summer and the drought period is autumn-winter; the winters are severe with occasional snow and a strong annual temperature variation (Kristensen and Frangi 1995). Annual mean temperature and precipitations are 14°C and 896 mm, respectively (SMN 1981, 1986).

Four *Eryngium* species out of seven recognized from Sierra de la Ventana (Frangi and Bottino, 1995) – *Eryngium horridum* Malme, *Eryngium stenophyllum* Urb., *Eryngium* aff. *serra* Cham. and Schltld. and *Eryngium elegans* Cham. and Schltld. – hold water and debris in their leaf axils (Figure 1). All species are terrestrial, *E. horridum* and *E. stenophyllum* occurred in dry and rocky soil, whereas *E. aff. serra*



**Figure 1.** *Eryngium horridum*: (A) flowering plant; (B) top view of a non-flowering plant showing arrangement of leaves; (C) detail of axils containing water.

and *E. elegans* grow near the edges of streams, in moist soils and in fields prone to flooding (Campos 2010).

### **Sampling design and sample extraction**

Forty plants of *E. horridum*, *E. stenophyllum*, *E. aff. serra* and *E. elegans* were randomly selected and sampled in spring and autumn. *Eryngium horridum* and *E. stenophyllum* were sampled from dry, open, sloping terrain, approximately > 20 m from streams, in 2006, and *E. aff. serra* and *E. elegans* were sampled from stream edges and ravines in 2007. All plants were exposed to sunlight because there are no trees in the study area. All samples were taken in the same area; about 500 m above sea level, in spring (flowering time) and autumn (senescence time). Two species of *Eryngium* were sampled from the edge of streams and two from 20 m away from the stream because they naturally grow in those sites. The two species growing on the bank (*E. aff. serra* and *E. elegans*) do not grow in areas away from the bank and vice versa. For this study, plants containing water in their axils were studied but no attempts were made to estimate the proportion of plants with or without water.

Fluid contents were extracted with a pipette attached to a lift pump, and subsequently all leaves except the most exterior were removed, from centre to periphery, and washed in a pan to release all adhering organisms; then the cavity formed by the remaining leaves was flushed twice with clean tap water, and the

aquatic contents were extracted with a pipette after each wash. All organisms were killed in the field and preserved in 80% commercial ethyl alcohol, except for some individuals of each morpho-species, which were carried alive to the laboratory in individual plastic tubes to be reared and identified. The fluid extracted (original siphoning plus washes) from the plant was observed under a stereo-microscope at 40 × magnification, and the immature dipterans were removed from the debris. Taxonomic resolution was attempted to species level; however, some aquatic larvae have not yet been matched to their adult stage, and unidentifiable species were recorded as morpho-species. Individuals of all taxa were sent to specialists for identification. Taxa for which no specialists were known in Argentina were classified to family following the keys of Stehr (1991).

### Data analyses

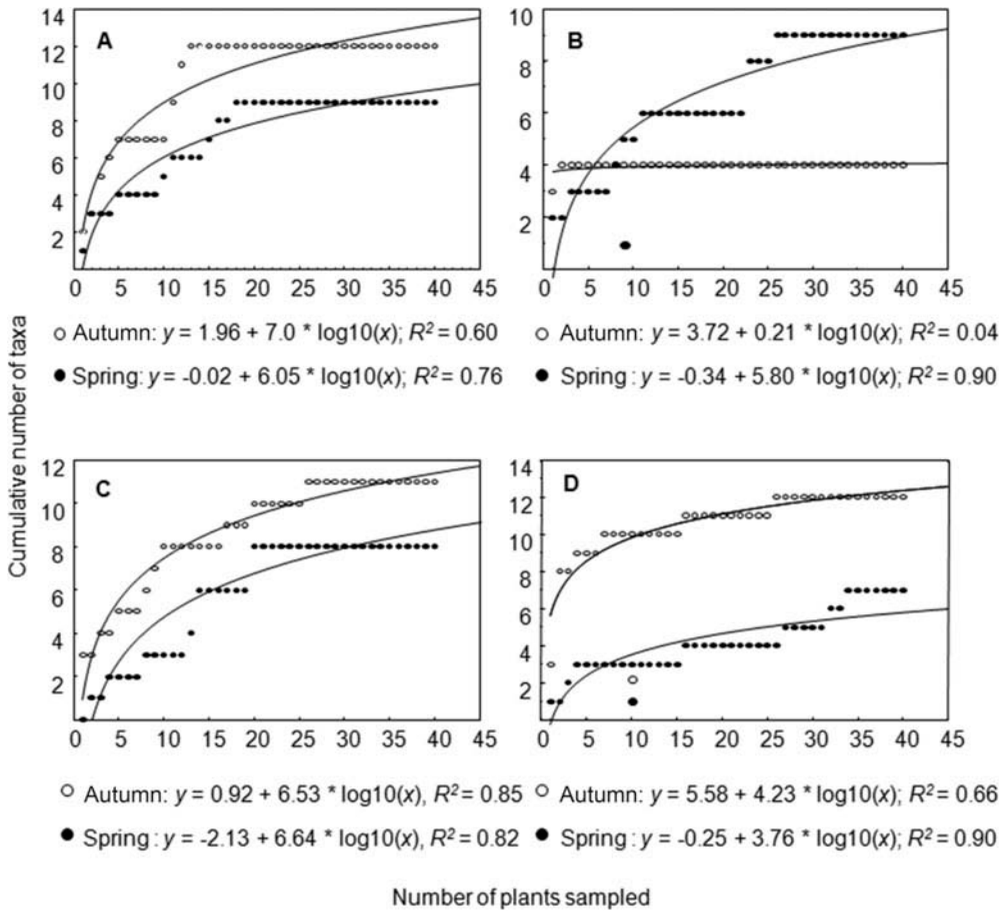
To estimate the sample size (number of plants) that quantified the number of taxa inhabiting each phytotelm, cumulative dipterans were plotted and fitted to the best curves. Comparisons between the proportion of *Eryngium* containing immatures of dipteran families in autumn and spring were tested by G-test ( $G_{adj}$ ). As the data obtained from samples violated the assumption of normality and homogeneity of variances, the non-parametric two-way analysis of variance with Scheirer–Ray–Hare extension (Scheirer et al. 1976; as described in Sokal and Rohlf 1995) was used to determine the main and interactive effects of each variable (season and *Eryngium* species) on the abundance of chironomid species. Multiple comparisons were performed by Mann-Whitney *U*-tests. In order to compromise between a large overall Type I error, which would arise with an  $\alpha$  of 0.05 for each separate test, and a large overall Type II error, which would arise with a strict application of Bonferroni correction (i.e. with an  $\alpha$  of  $0.05 \times 15^{-1} = 0.003$ ), an  $\alpha$  of 0.01 in each separate post hoc test was used (Nugues and Bak 2006).

Similarity between the four *Eryngium* species related to dipteran communities was analysed by cluster analysis, using arithmetic averages and the un-weighted pair-group method. For cluster analysis, the Morisita–Horn index was used because it is not dependent on sample size, and is the index least influenced by species richness (Wolda 1981). The analyses were performed using Bio-Dap software (Magurran 1988).

## Results

### Adequacy of sampling

The evaluation carried out to estimate the total taxa inhabiting each plant species indicated that relationships between number of plants sampled and taxa identified, fitted to logarithmic functions, which demonstrated tendencies to reach a plateau (Figure 2).  $R^2$  values were  $> 0.66$  and significant ( $p < 0.001$ ) for all species of plants in autumn and spring, except for *E. stenophyllum* in autumn ( $R^2 = 0.04$ ;  $p = 0.09$ ) where two plants yielded all the Diptera taxa recorded from this phytotelm.



**Figure 2.** Cumulative total of Diptera observed in successive plants ( $n = 40$  plants for each species in autumn and in spring). (A) *Eryngium horridum*; (B) *Eryngium stenophyllum*; (C) *Eryngium aff. serra*; (D) *Eryngium elegans*. Logarithmic equations with  $R^2$  values;  $p < 0.001$  for all  $R^2$  except for *E. stenophyllum* in autumn ( $p = 0.09$ ).

### Community composition and seasonal occurrence of dipterans inhabiting *Eryngium phytotelmata*

The community of dipterans that live in the axils of *Eryngium* consisted of aquatic and semi-aquatic larvae, belonging to six and seven families, respectively (Table 1). Of the 13 families in the community, Chironomidae accounted for 87.6%, followed by Culicidae (3.9%), Perisclididae (3.4%) and Ceratopogonidae (3.1%). The remaining families accounted for 2%.

All *E. horridum* and *E. aff. serra*, 37 *E. stenophyllum* and 39 *E. elegans* contained larvae of dipterans in autumn, and 38 *E. horridum*, 40 *E. stenophyllum*, 37 *E. aff. serra* and 34 *E. elegans* contained larvae of dipterans in spring. Ten families of dipterans were recorded from *E. horridum*, all present in autumn, eight of which were also present in spring, except Asilidae and Dolichopodidae. Nine families were present in *E. stenophyllum* in spring, and only Scatopsidae and Chironomidae in autumn; this was

**Table 1.** Taxonomic classification and number of individuals of the aquatic dipterans inhabiting four *Eryngium* species censused in autumn and spring at Sierra de la Ventana, Argentina.

Taxa	<i>Eryngium horridum</i>		<i>Eryngium stenophyllum</i>		<i>Eryngium aff. serra</i>		<i>Eryngium elegans</i>	
	Autumn	Spring	Autumn	Spring	Autumn	Spring	Autumn	Spring
Tipulidae								1
Cecidomyiidae*	1	5		9	2	7	10	
Scatopsidae*	87	39	47	75	13	1	8	7
Culicidae								
<i>Culex castroi</i>	87	29						
<i>Culex renatoi</i>	247	514		3	22		3	
Ceratopogonidae								
<i>Dasyhelea</i> sp.					181	48	501	14
<i>Forcipomyia</i> sp.						1		
Chironomidae								
<i>Metriocnemus eryngiotelmatus</i>	1230	712	2455	1393	352	188	860	63
<i>Polypedilum parthenogeneticum</i>	4990	3814	64	1124	1830	74	1331	62
<i>Cricotopus</i> sp.								1
<i>Larsia</i> sp.					4	1	28	1
Asilidae*	3							
Dolichopodidae	4			22				1
Phoridae*					1			
Syrphidae								
Morpho-species 1	17	4						
Morpho-species 2				3				
Tanypezidae*	3	1		12	1		2	
Sphaeroceridae*	25	14		3		1	3	1
Perisclididae: <i>Stenomicro</i> sp.	460	319		4	1		3	
Unidentified Diptera*	4		24		1	4	3	

\*Semi-aquatic.

the only *Eryngium* where the greatest diversity occurred in spring. In *E. aff. serra* nine families of dipterans occurred, eight in autumn and five in spring. Of the families present in spring, four were also present in autumn, except Sphaeroceridae. Whereas for *E. elegans*, 10 families were recorded, nine in autumn and five in spring. Of the families present in spring, four were also present in autumn, except Dolichopodidae (Table 1).

The presence of Chironomid larvae in 70% and 97.5% of all the *E. horridum*, and in 75% and 97.5% of all the *E. elegans* in spring and autumn, respectively, indicates a high and significant presence in autumn ( $p < 0.01$ ). Furthermore, the occurrence of ceratopogonid larvae in *E. aff. serra* and in *E. elegans* in spring (10% and 15%) and in autumn (45% and 65%), indicates a significant preference for the latter season in both *Eryngium* phytotelmata ( $p < 0.001$ ). Sphaeroceridae larvae occurred in more *E. horridum* plants in autumn (50%) than in spring (25%;  $p < 0.05$ ). G tests for the other families of dipterans did not significantly support seasonal independence of phytotelmata use (Table 2).

## Relevant families of dipterans of the communities inhabiting *Eryngium phytotelmata*

### Chironomidae

Four species of Chironomidae were found breeding in the leaf axils of *Eryngium*, one belonging to the subfamily Chironominae (*Polypedilum parthenogeneticum* Donato and Paggi) two to Orthocladiinae (*Metriocnemus eryngiotelmatus* Donato and Paggi, and

**Table 2.** G-test results comparing the proportion of *Eryngium* containing immatures of dipteran families in autumn and spring.

Families	<i>E. horridum</i>		<i>E. stenophyllum</i>		<i>E. aff. serra</i>		<i>E. elegans</i>	
	G <sub>adj</sub>	P	G <sub>adj</sub>	P	G <sub>adj</sub>	P	G <sub>adj</sub>	P
Cecidomyiidae	0.26	>0.70			0.18	>0.70		
Scatopsidae					1.69	>0.20	0	>0.99
Culicidae	0.05	>0.80						
Ceratopogonidae					11.14	<0.001	19.79	<0.001
Chironomidae	10.22	<0.01	1.28	>0.30	2.65	>0.20	7.42	<0.01
Sphaeroceridae	4.37	<0.05					0.27	>0.70
Perisclididae	0.27	>0.70						
Unident. Diptera					0.88	>0.50		

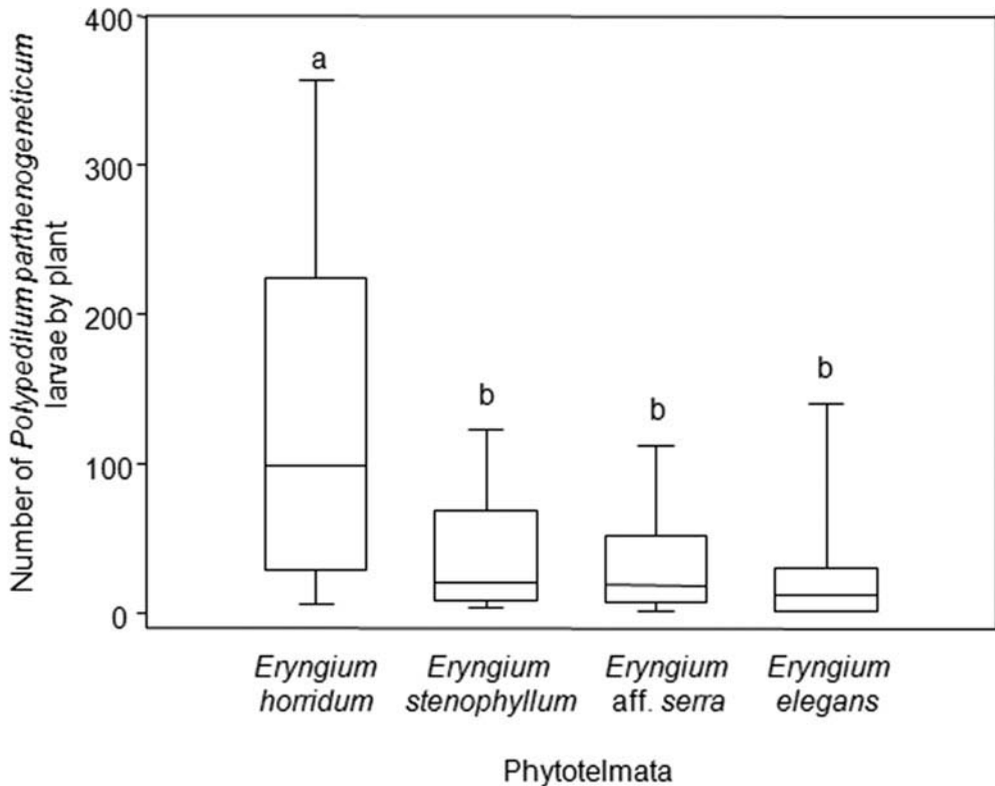
Families represented by less than 30 individuals and/or lacking in a station, were not tested.

*Cricotopus* sp.), and a Tanypodinae (*Larsia* sp.). *Metriocnemus eryngiotelmatus* and *P. parthenogeneticum* were found in the four species of *Eryngium*; *Larsia* sp. in *E. aff. serra* and *E. elegans*; and *Cricotopus* sp. in *E. elegans*. *Metriocnemus eryngiotelmatus* and *P. parthenogeneticum* were the dominant Chironomidae, *Larsia* sp. was scarce, and *Cricotopus* was rare, a single individual being found (Table 1).

Because *M. eryngiotelmatus* and *P. parthenogeneticum* occurred in the four *Eryngium* species, it was of interest to compare the abundance of both chironomids within phytotelmata and the seasons in which they occurred. The number of larvae of *P. parthenogeneticum* per plant was significantly higher in *E. horridum* than in the other three phytotelmata ( $H = 39.37$ ,  $df = 3$ ,  $p < 0.001$ , Figure 3). Whereas for *M. eryngiotelmatus* it was significantly higher in *E. horridum* and *E. stenophyllum* than in *E. aff. serra* and *E. elegans* ( $H = 63.39$ ,  $df = 3$ ,  $p < 0.001$ , Figure 4).

The abundance of *P. parthenogeneticum* was affected by seasons, *Eryngium* phytotelmata and their interaction (Table 3). All *Eryngium* phytotelmata showed significant differences in the number of individuals of *P. parthenogeneticum* ( $U_{E. horridum} = 1827$ ;  $p < 0.05$ ;  $U_{E. stenophyllum} = 1264.5$ ;  $U_{E. aff. serra} = 2339$ ;  $U_{E. elegans} = 2007$ ;  $n = 40$ ;  $p < 0.001$ ) being higher in autumn than in spring, except for *E. stenophyllum* where it was the inverse (Table 1). The number of *P. parthenogeneticum* larvae by *Eryngium* phytotelmata in autumn and spring was significant ( $H_{Autumn} = 73.90$ ;  $H_{Spring} = 28.80$ ;  $df = 3$ ;  $p < 0.001$ ). Paired comparisons for autumn detected significant differences in the number of *P. parthenogeneticum* larvae for all *Eryngium* phytotelmata. Whereas for spring, four of six comparisons were significantly different (Table 3).

The abundance of *M. eryngiotelmatus* was affected by season, *Eryngium* phytotelmata and their interaction (Table 4). All *Eryngium* phytotelmata showed significant differences in the number of individuals of *M. eryngiotelmatus* between autumn and spring ( $U_{E. horridum} = 1965$ ;  $U_{E. aff. serra} = 2002.5$ ;  $U_{E. elegans} = 2274$ ;  $n = 40$ ;  $p < 0.001$ ), except for *E. stenophyllum* ( $U = 1808$ ;  $n = 40$ ;  $p > 0.05$ ). As with *P. parthenogeneticum*, the highest abundance of *M. eryngiotelmatus* occurred in autumn (Table 1). The number of *M. eryngiotelmatus* larvae by *Eryngium* phytotelmata in autumn and in spring was significant ( $H_{Autumn} = 23.58$ ;  $H_{Spring} = 32.21$ ;  $df = 3$ ;  $p < 0.001$ ). Significant differences were found in the number of *M. eryngiotelmatus* larvae for four *Eryngium* phytotelmata comparisons and they were nonsignificant for two, in both, autumn and spring (Table 4).

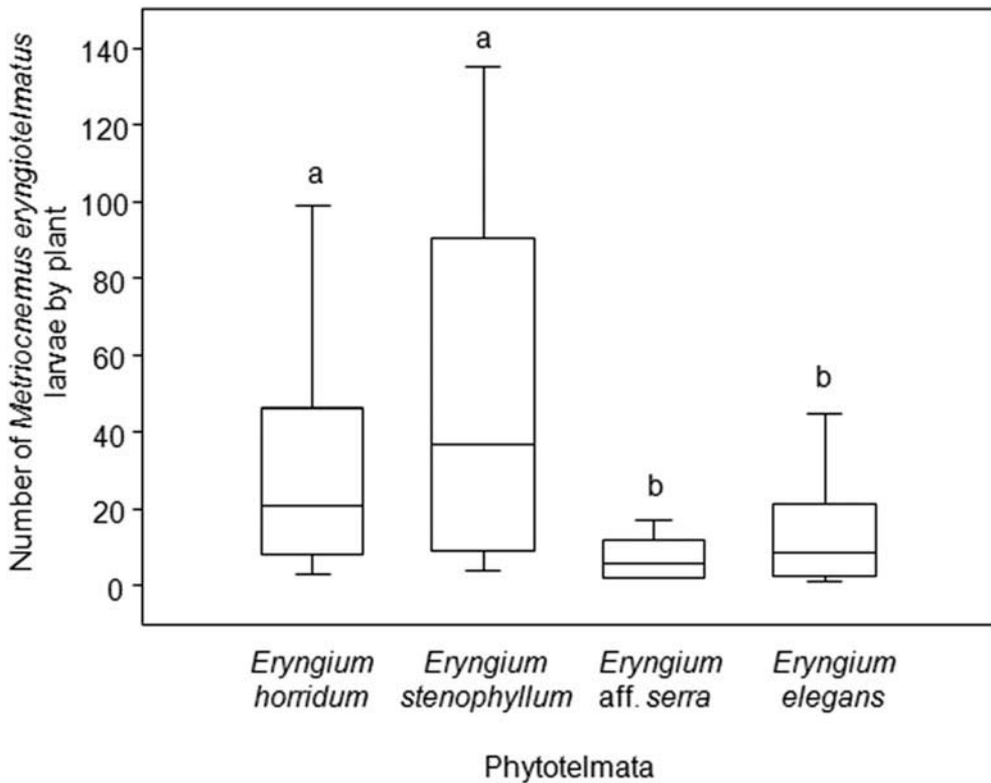


**Figure 3.** Median number of *Polypedilum parthenogeneticum* larvae per plant in four *Eryngium* phytotelmata. The box indicates the 25th and 75th centiles, the line within the box indicates the median and the whiskers indicate the 90th and 10th centiles. Different letters indicate significant differences detected by Dunn's test ( $p < 0.05$ ).

Because one plant can be occupied by *P. parthenogeneticum* and *M. eryngiotelmatus*, their co-occurrence in each phytotelmata was analysed. Of all plants containing both species of chironomids, *E. horridum* was the most relevant. Over 80% were occupied by both species and *P. parthenogeneticum* was the most abundant species in 49 plants. The occurrence of both species of chironomids was lower in the other three *Eryngium* species, and *P. parthenogeneticum* was dominant in most plants. In only four plants, both species of chironomids occurred in equal numbers (Table 5). The average number of individuals of *P. parthenogeneticum* per plant, where it was the dominant species, was greater than *M. eryngiotelmatus* in plants where this was the dominant species (except for *E. stenophyllum*). Conversely, the average number of individuals of *M. eryngiotelmatus* per plant, where it was not the dominant species, was higher than that of *P. parthenogeneticum* in plants where this was not the dominant species (Table 5).

### **Culicidae**

Two species of mosquitoes were found breeding in *Eryngium* phytotelmata, *Culex castroi* Casal and García and *Culex renatoi* Lane and Ramalho. *Culex renatoi* occurred



**Figure 4.** Median number of *Metriocnemus eryngiotelmatus* larvae per plant in four *Eryngium* phytotelmata. The box indicates the 25th and 75th centiles, the line within the box indicates the median and the whiskers indicate the 90th and 10th centiles. Different letters indicate significant differences detected by Dunn's test ( $p < 0.05$ ).

**Table 3.** Results of two-way analysis of variance with Scheirer–Ray–Hare extension on the ranks of *Polypedilum parthenogeneticum* larval abundance by seasons and *Eryngium* phytotelmata, and post-hoc multiple comparisons (pairwise comparisons by Mann–Whitney *U*-test).

Source	df	SS	SS/MS <sub>Total</sub>	<i>p</i> -value
Season	1	156,777.778	18.315	<0.001
<i>Eryngium</i>	3	421,309.856	49.218	<0.001
S × E	3	411,527.978	175.476	<0.001

Comparisons	Autumn		Spring	
	<i>U</i>	*	<i>U</i>	*
<i>E. horridum</i> – <i>E. stenophyllum</i>	2312	*	1781.5	ns
<i>E. horridum</i> – <i>E. aff. serra</i>	1874	*	2007.5	*
<i>E. horridum</i> – <i>E. elegans</i>	2060	*	2011	*
<i>E. stenophyllum</i> – <i>E. aff. serra</i>	892.5	*	1921.5	*
<i>E. stenophyllum</i> – <i>E. elegans</i>	1187.5	*	1927.5	*
<i>E. aff. serra</i> – <i>E. elegans</i>	1951.5	*	1622.5	ns

Kruskal–Wallis test showed significant differences among phytotelmata ( $H_{Autumn} = 73.90$ ;  $H_{Spring} = 28.80$ ;  $df = 3$ ;  $p < 0.001$ ). \* =  $p < 0.01$ , ns = not significant ( $p > 0.01$ ).

in the four species of *Eryngium*, but was abundant only in *E. horridum* where it coexisted with *C. castroi*. The abundance of *C. renatoi* was higher in spring, in contrast to *C. castroi* (Table 1).

**Table 4.** Results of two-way analysis of variance with Scheirer–Ray–Hare extension on the ranks of *Metriocnemus eryngiotelmatus* larval abundance by seasons and *Eryngium* phytotelmata, and post-hoc multiple comparisons (pairwise comparisons by Mann–Whitney *U*-test).

Source of variation	df	SS	SS/MS <sub>Total</sub>	<i>p</i> -value
Season	1	405,341.628	47.353	<0.001
<i>Eryngium</i>	3	312,537.206	36.511	<0.001
S × E	3	93,628.153	10.938	<0.02
Comparisons	Autumn		Spring	
	<i>U</i>		<i>U</i>	
<i>E. horridum</i> – <i>E. stenophyllum</i>	1429	ns	1335.5	*
<i>E. horridum</i> – <i>E. aff. serra</i>	2005.5	*	1660.5	ns
<i>E. horridum</i> – <i>E. elegans</i>	1732.5	ns	1802	ns
<i>E. stenophyllum</i> – <i>E. aff. serra</i>	2010	*	2041	*
<i>E. stenophyllum</i> – <i>E. elegans</i>	1899.5	*	2184	*
<i>E. aff. serra</i> – <i>E. elegans</i>	1308.5	*	1872	*

Kruskal–Wallis test showed significant differences among phytotelmata ( $H_{Autumn} = 23.58$ ;  $H_{Spring} = 32.21$ ;  $df = 3$ ;  $p < 0.001$ ). \* =  $p < 0.01$ , ns = not significant ( $p > 0.01$ ).

**Table 5.** Biological parameters of *Eryngium* plants where *Polypedilum parthenogeneticum* and *Metriocnemus eryngiotelmatus* (Chironomidae) co-occur.

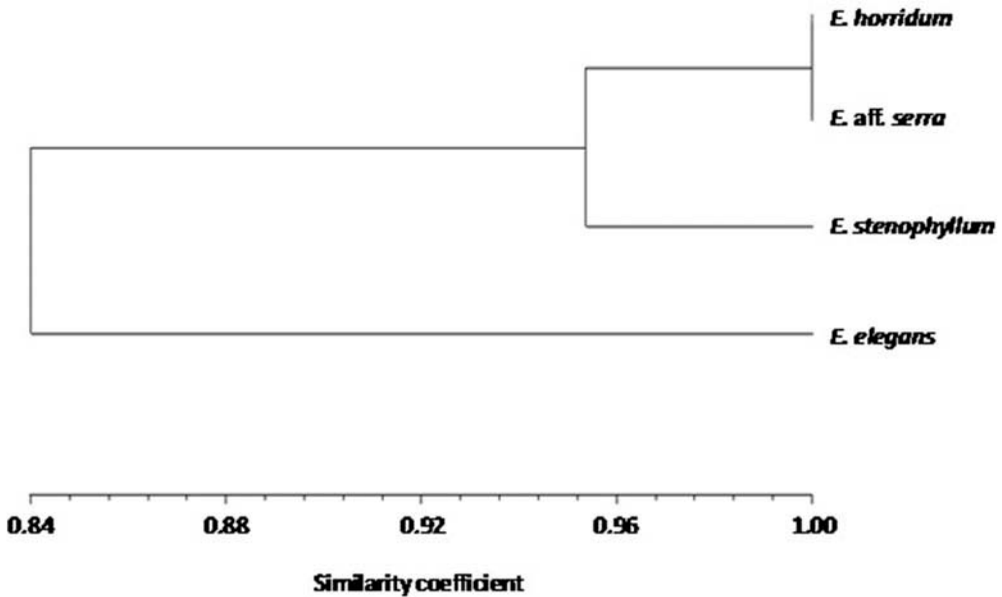
Parameters	<i>Eryngium horridum</i>	<i>Eryngium stenophyllum</i>	<i>Eryngium aff. serra</i>	<i>Eryngium elegans</i>
No. of plants where chironomid species co-occur/no. of plants with chironomids (%)	53/66 (80.3)	24/72 (33.3)	41/76 (53.9)	29/68 (42.6)
No. of plants where <i>P. parthenogeneticum</i> is dominant.	49	10	36	15
No. of plants where <i>M. eryngiotelmatus</i> is dominant.	4	13	2	14
No. of plants with similar no. of individuals of both chironomid species.	0	1	3	0
Mean no. ± standard deviation of individuals of <i>P. parthenogeneticum</i> when it is dominant.	171.6 ± 150.2	70.1 ± 35.1	51.1 ± 47.2	79.5 ± 92.6
Mean no. ± standard deviation of individuals of <i>P. parthenogeneticum</i> when it is not dominant.	15.5 ± 10.5	9.6 ± 6.0	2.0 ± 1.4	8.9 ± 8.4
Mean no. ± standard deviation of individuals of <i>M. eryngiotelmatus</i> when it is dominant.	44.2 ± 39.8	95.4 ± 48.3	9.5 ± 10.6	25.6 ± 26.5
Mean no. ± standard deviation of individuals of <i>M. eryngiotelmatus</i> when it is not dominant.	33.2 ± 32.4	12.0 ± 9.1	8.5 ± 5.9	18.6 ± 16.9

### Perisclididae

An unidentified species of Perisclididae of the genus *Stenomicroa* Coquillett was found in the axils of the four species of *Eryngium*. However, its presence was strongly associated with *E. horridum* where 99% of individuals (779/787 immature collected) occurred. No seasonal difference in the individuals' abundance was observed (Table 1), but occurrence of pupae was limited to spring.

### Ceratopogonidae

Two unidentified species of Ceratopogonidae of the genera *Dasyhelea* and *Forcipomyia* were collected from *Eryngium* axils. *Forcipomyia* had a single individual present in *E. aff. serra*, whereas *Dasyhelea* sp. occurred in *E. aff. serra* and *E. elegans*, and was more abundant in autumn (Table 1).



**Figure 5.** Cluster analysis of the *Eryngium* phytotelmata in autumn, using the Morisita–Horn index and UPGMA linkage.

### Similarity among dipteran communities

Morisita–Horn indexes of similarity were high for all combinations of species in both autumn and spring. Cluster analysis for the autumn (Figure 5), shows that most differences involved dipteran communities inhabiting *E. elegans*. Whereas in spring (Figure 6), two groups were defined, one composed of *E. horridum* and *E. aff. serra*, and the second group composed of *E. stenophyllum* and *E. elegans*.

## Discussion

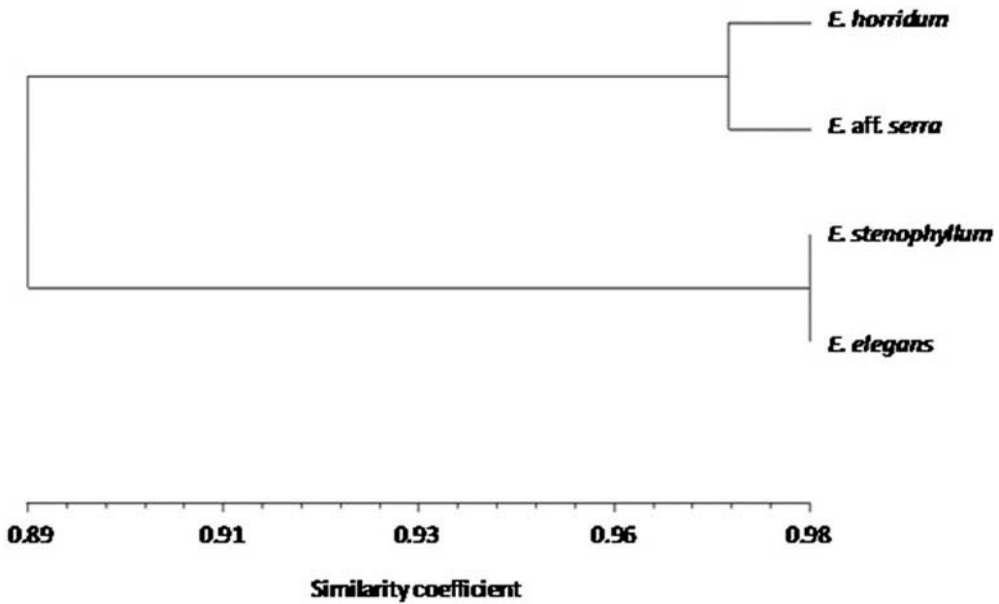
### Composition of dipterans communities inhabiting *Eryngium* phytotelmata

The communities of Diptera associated with the four species of *Eryngium* consist of saprophages that feed either on the suspended particulate matter, or on small particles that they scrape from package detritus (fine detritus feeders), including chironomids, *M. eryngiotelmatus* and *P. partenogeneticum* and Culicidae *Culex* (*phytotelmatomyia*).

Some syrphid larvae were found in the deepest parts of the axils and could feed by ‘scraping’ the substrate or the surface of plant leaves. Ceratopogonid larvae (*Dasyhelea* and *Forcipomyia*) that could have fed in the detritus (Kitching 2004) were less frequent.

The ‘fine detritus feeder’ dipterans were dominant in the communities inhabiting phytotelmata of *Eryngium*.

Among the dipterans inhabiting the axils of *Eryngium*, only free-swimming predators were recorded. Khoo (1984; cited in Kitching 2004) mentions that the larvae of periscelid *Stenomicro* are specialized predators in the axils of bromeliads, subsequently Fish (1976) reported that these larvae prey on mosquito larvae *Wyomyia* spp. Larvae of *Stenomicro*



**Figure 6.** Cluster analysis of the *Eryngium* phytotelmata in spring, using the Morisita–Horn index and UPGMA linkage.

were abundant in the axils of *E. horridum*, both in spring and in autumn. Although it was not possible to establish which species were their prey, it could be speculated that early Chironomidae larvae, given their abundance, could form part of their diet.

In the dipteran communities of the other three species of *Eryngium* studied here, no significant predators were detected. Although *Larsia* sp. chironomid larvae were found, their predatory action would be limited by their small size, and their scarce abundance, exerting a limited effect on the community.

### **Relevant families of dipterans of the communities inhabiting *Eryngium* phytotelmata**

#### ***Chironomidae***

Larvae of *M. eryngiotelmatus* and *P. parthenogeneticum*, representing over 87% of the number of individuals prevailed in the community of dipterans that lived in the axils of *Eryngium*. Kitching (1971) recorded up to three generations of the chironomid *Metriocnemus martinii* Thienermann in tree holes, with two population peaks, one in late summer and the other, greater, in autumn. This author considers that hibernation occurs in the larval stage, and the reason for the decrease in the density of the population during that season is mortality, presumably as a result of environmental factors. Although populations of *M. eryngiotelmatus* and *P. parthenogeneticum* inhabiting *Eryngium* were not sampled in all seasons, it was observed that the population peak also occurred in autumn, doubling the abundance of spring larvae. Furthermore, overlapping generations evidenced by

overlapping stages, and the presence of pupae in both spring and autumn, suggests that both species are multivoltine in the temperate region of Argentina.

The number of chironomid larvae per plant was highly variable between species and phytotelmata. This variation was more noticeable for *P. parthenogeneticum* than for *M. eryngiotelmatus*, and although this pattern was observed in the four species of *Eryngium*, it was more evident in *E. horridum*.

Kitching (1972), in a study on oviposition of *M. martinii* reported an average of 51.3 eggs per clutch with a wide range around the mean, without observing seasonal variation. So far, the biology of *M. eryngiotelmatus* oviposition is not known, but according to observations in the laboratory, *P. parthenogeneticum* reproduces by parthenogenesis and females oviposit between 12 and 24 hours after the emergence of the imago, each female producing more than three batches containing between 37 and 123 eggs per clutch (Donato and Paggi 2008). That reproductive potential, coupled with several females ovipositing simultaneously, explain the high number of larvae observed in some plants. Larvae observed in plants with an abundant number of individuals, mostly corresponded to early stages, while in plants with fewer individuals late larvae and pupae predominated. This variation in the number of larvae, could show that the higher incidence of mortality factors could be acting on the early stages of both chironomids.

In several studies, the coexistence of chironomid species in other than *Eryngium* phytotelmata is mentioned. Picado (1913) described *Metriocnemus abdominoflavatus* and *Chirocladius pedipalpus* living in axils of tank bromeliads in Costa Rica, and Miller (1971) in a study of two species of bromeliads in the U.S. Virgin Islands, mentions the co-existence of three species of the genera *Tanytarsus*, *Pentaneura* and *Metriocnemus*, the latter being the least abundant. Moreover, Beaver (1983) observed species of the genera *Metriocnemus* and *Polypedilum* breeding in the pitcher plant *Nepenthes villosa* in Borneo, and in plants of the same species introduced to Britain from Sabah (cited in Cranston and Judd 1987). The co-occurrence of *M. eryngiotelmatus* and *P. parthenogeneticum* in *Eryngium* was reported by Donato and Paggi (2008). The ecological association between these two species was studied using the association coefficient  $C_8$  species, estimated for four species of *Eryngium* but significant only for *E. horridum* (Campos 2010). The present study shows that the co-existence of both species occurs in the four species of *Eryngium*, being higher in *E. horridum*, where both chironomids were present in 80% of the plants examined. *Polypedilum parthenogeneticum* was dominant in three of the four phytotelmata (see Table 5), however the number of individuals in the plants where only this species was found, was lower compared with that of *M. eryngiotelmatus* when this species was in the same situation. These observations require specific laboratory studies to distinguish between the effects of intra- and inter-specific competition.

The high reproductive potential of *P. parthenogeneticum* and the similarity in the pattern of abundance in the four species of *Eryngium*, could indicate a better fitness for this species than for *M. eryngiotelmatus*, particularly when inhabiting the same plant.

### **Culicidae**

The family Culicidae has been cited as one of the main inhabitants of phytotelmata, both in species diversity and in numerical abundance (Fish 1983). Three species of *Culex*

(*Phytotelmatomyia*) and an undetermined species inhabiting the axils of *Eryngium* spp. are known in the temperate zone of Argentina (Campos and Lounibos 1999; Albicocco et al. 2011) above the parallel 34°51'33"S. South of this latitude, in Sierra de la Ventana, two species of mosquitoes, *C. castroi* and *C. renatoi* were found (Campos 2010), but their abundance was low, with *E. horridum* phytotelmata the best host to both species. In the present study, both species were found in *E. horridum*, but only *C. renatoi* was found in the other species of *Eryngium*. Its population was abundant in *E. horridum*, with a notable increase in the spring. In this study, *C. castroi* was more abundant in autumn than in spring, whereas *C. renatoi* showed an inverse pattern. However, in both seasons, *C. renatoi* was much more abundant than *C. castroi*. In a previous study (Campos and Lounibos 1999), where the occurrence of these two species of mosquitoes in *E. cabreræ* Pontiroli was analysed, a predominance of *C. renatoi* with respect to *C. castroi* was also observed.

### **Periscelididae**

Larvae of *Stenomicro* were recently reported in Argentina, breeding in *Eryngium* spp. with a marked seasonality of pupation, which occurs in October and is produced on the leaves out of the water (Campos et al. 2010). In the present study, the same seasonal pattern was observed, suggesting that in the temperate zone of Argentina, *Stenomicro* sp. is univoltine, with a long life cycle and hibernation in the larval stage.

### **Ceratopogonidae**

Three species of ceratopogonids are known to breed in *Eryngium* phytotelmata, *Culicoides charruus* Spinelli and Martínez, and two undetermined species of the genera *Dasyhelea* and *Forcipomyia* (Campos 2010, Appendix.). *Culicoides charruus*, whose larvae were recorded from axils of *E. serra* (Spinelli and Martínez 1991) and *E. pandanifolium* Cham. and Schlecht (Ronderos et al. 2008), was not found in the species of *Eryngium* of Sierra de la Ventana. More than 20 species of *Dasyhelea*, breeding in pitcher plants, tree holes, water axils and bamboo internodes, are known (Kitching 2004). An undetermined species of *Dasyhelea* was the dominant ceratopogonid in *E. aff. serra* and *E. elegans* and was more abundant in autumn than in spring, but it was not recorded in *E. horridum* or in *E. stenophyllum*.

### **Similarity among dipteran communities**

Both clusters show that dipteran communities inhabiting *E. horridum* and *E. aff. serra* are similar, whereas the communities of dipterans inhabiting *E. stenophyllum* and *E. elegans* differ from the first in both autumn and spring. High values of the similarity coefficients indicate that the four phytotelmata provide a suitable habitat for all taxa of the dipteran communities. However, the strong similarity between communities could be attributed to the high sensitivity of the coefficient to the dominant species, being influenced by the abundance of Chironomidae.

### **Conclusion**

It is concluded that the four sympatric *Eryngium* spp. are used by Diptera differently and their populations show seasonal changes, with peaks in autumn. The communities of dipterans that live in the axils of the four *Eryngium* species are composed of larvae

belonging to six aquatic and seven semi-aquatic families. The main families are Chironomidae, Culicidae, Perisclididae and Ceratopogonidae. Two species of Chironomids (*M. eryngiotelmatus* and *P. polypedilum*) were the most abundant in all *Eryngium* species in both autumn and spring. Species of Culicidae, Perisclididae and Ceratopogonidae were more selective than Chironomidae. The first two families were best represented in the community of *E. horridum*, whereas Ceratopogonidae were found in *E. aff. serra* and *E. elegans*. The seasonal presence of the families of dipterans was different between *Eryngium* species. Major diversity was recorded from all *Eryngium* in the autumn, except for *E. stenophyllum*. Saprophages were the most abundant in the communities, with the fine detritus feeders being the dominant group. The perisclid *Stenomicro* was the most important free-swimming predator (abundant in *E. horridum* and scarce in the others species of *Eryngium*). No top predators were recorded in the four *Eryngium* phytotelmata.

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No potential conflict of interest was reported by the author.

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