

## Community structure in mediterranean shallow lentic ecosystems: size-based vs. taxon-based approaches

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### ABSTRACT

The main environmental variables determining the community structure and the functioning of Mediterranean shallow lentic ecosystems are described. These ecosystems are characterized by the unpredictability of their water inputs and the high variability in their water level and physical and chemical composition. Variations in flooding, salinity, and water turnover are determinant in species composition and nutrient dynamics. Taxon-based and size-based approaches to the study of the community structure of aquatic organisms that colonise these ecosystems are also compared. The conventional taxonomic approach, based on the determination of species composition, has been used for the identification of patterns in species richness, distribution and temporal dynamics, and for ecological requirements of species and their potential use as ecological indicators. This taxon-based approach has been compared with a size-based approach, where individuals are classified by their size. Size-based approach gives complementary information about community structure and dynamics, especially when communities are dominated by a single species. The use of size diversity combined with species diversity is suggested for a more complete understanding of community structuring in this type of ecosystem. Detailed examples of two Mediterranean shallow lentic ecosystems, the salt marshes of the Empordà wetlands and the Espolla temporary karstic pond, which differ in hydrology and water origin, are used to discuss the suitability of these different approaches.

**Keywords:** Mediterranean wetlands, shallow lakes, size diversity, species diversity, biomass size spectra.

### RESUMEN

*Los ecosistemas leníticos someros mediterráneos se caracterizan por la impredecibilidad en las entradas de agua y por la elevada variabilidad en el nivel del agua y su composición física y química. Se describen aquí las principales variables ambientales que determinan la estructura de la comunidad y el funcionamiento de estos ecosistemas. Variaciones en la inundación, la salinidad y la tasa de renovación del agua son determinantes en la composición de especies y en la dinámica de nutrientes. Se comparan también aproximaciones al estudio de la comunidad basadas en la composición taxonómica con las basadas en la distribución de tamaños. La aproximación taxonómica convencional, basada en la determinación de la composición de especies, es adecuada para la identificación de patrones en la distribución y en la dinámica temporal de la especie, así como para el análisis de los requerimientos ambientales de las diferentes especies y su uso potencial como indicadores ecológicos. Esta aproximación taxonómica se ha comparado con otra aproximación basada en el tamaño corporal, donde los individuos se clasifican por su tamaño. La aproximación basada en el tamaño proporciona información complementaria de la estructura y dinámica de las comunidades, especialmente cuando en estas comunidades hay una especie dominante. Se sugiere el uso combinado de una diversidad de tamaños y una diversidad de especies para una comprensión más completa de la estructuración de las comunidades en este tipo de ecosistemas. Para discutir si estas diferentes aproximaciones son o no adecuadas, se presentan datos de dos ecosistemas mediterráneos leníticos y someros: las marismas de Aiguamolls de l'Empordà y la laguna de Espolla, una surgencia cárstica de inundación temporal.*

**Palabras clave:** Humedales mediterráneos, lagos someros, diversidad de tamaños, diversidad de especies, espectros de biomasa.

### INTRODUCTION

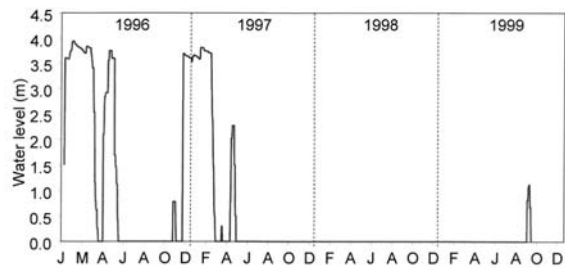
Mediterranean shallow lentic ecosystems include shallow lakes, wetlands, coastal marshes and ponds, which present a hydrology strongly con-

ditioned by the Mediterranean climate. Irregularity and unpredictability of water inputs and flooding periods have a strong influence on their hydrology, causing not only sharp water level fluctuations but also wide fluctuations in

their physical, chemical, and biological composition (Fernández-Aláez, *et al.*, 1999; Beklioglu *et al.*, 2003; Romo *et al.*, 2004; Alvarez-Cobelas *et al.*, 2005). Spatial and temporal variability in water composition favours large environmental gradients over short time and spatial scales. As a consequence, species richness is very high, although this is frequently not well known and often neglected (Boix *et al.*, 2001b; Boix & Sala 2002; Martinoy *et al.*, in press).

Most of these ecosystems are characterized by sudden inputs of water, coinciding with flooding events, which are frequently followed by long periods of confinement. Decreases in water levels during confinement may lead to desiccation or, at least, to very low levels. Therefore, these ecosystems may become temporary and suffer one or two dry periods throughout the year. This characteristic hydrology strongly determines the ecological functioning of these ecosystems, where the duration of the hydroperiod and water turnover rate have been determined to affect water composition, nutrient dynamics, contaminant contents and aquatic community structure (Quintana *et al.*, 1998a and 1998b; López-Flores *et al.*, 2003; Boix *et al.*, 2004). This sudden flooding – confinement hydrology is less determinant for the nutrient cycle in those Mediterranean salt marshes or coastal lagoons where other factors such as human regulation, riverine inputs or a permanent connexion with the sea are more relevant (Heurteaux *et al.*, 1992; Comín & Valiela, 1993; Frascari *et al.*, 2002; Pérez-Ruzafa *et al.*, 2002; Villena & Romo, 2003).

Temporary habitats have no fish unless they are filled with water coming from permanently flooded ecosystems. In the absence of fish, invertebrate predation and competitive interactions among invertebrates become more relevant in the trophic structure of the aquatic community (Wellborn *et al.*, 1996). Situations in which a single invertebrate species dominates are frequent in Mediterranean shallow water bodies (Quintana *et al.*, 1998b; Brucet *et al.*, 2005b) due to competitive exclusion. When the invertebrate community is dominated by one single species, intraspecific competitive interactions will take place and the separation of niches among diffe-



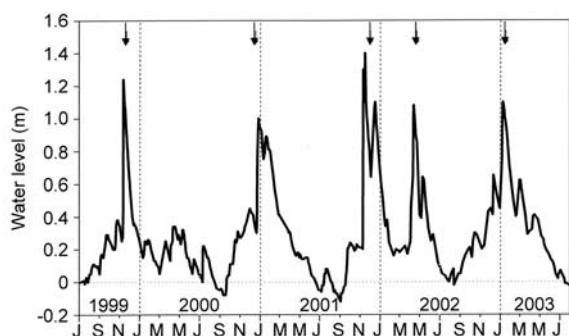
**Figure 1.** Water level over 7 consecutive hydroperiods at the Espolla pond (1996-1999). *Nivel del agua a lo largo de 7 hidroperíodos consecutivos en la laguna de Espolla (1996-1999).*

rent stages may allow the dominant species to reduce intraspecific competition (Werner & Gilliam 1984). In this case, a size-based approach of the study of the community structure would probably show the niche partitioning better than the traditional taxon-based approach.

In this paper the main physical and chemical characteristics of several shallow lentic water bodies located in Catalonia (NE Spain) are described, and studies of the composition and size distribution of their species are compared. We mainly discuss data from two ecosystems with different hydrology and water origin: the salt marshes of the Empordà wetlands (Alt Empordà and Baix Ter wetlands) and the temporary karstic pond of Espolla. The former are mainly flooded by sudden seawater inputs during sea storms, whilst the latter is flooded by karstic groundwater springs. Our aim is to show the main limnological characteristics of these shallow lentic ecosystems, and to compare the conclusions obtained in the study of the community structure by means of two different approaches, the taxon-based and the size-based approaches.

## HYDROLOGY AND NUTRIENT DYNAMICS

Hydroperiods in the karstic Espolla pond (Fig. 1) are characterized by rapid flooding and drying and by a high water turnover rate throughout the hydroperiod (Boix *et al.*, 2004). Conductivity values do not increase significantly during the drying out period, indicating no substantial



**Figure 2.** Variation in the water level (m above the average sea level) in Fra Ramon lagoon (Baix Ter wetlands) during four hydrological cycles (from 1999 to 2003, separated by dotted lines). Arrows indicates sea storm events. The zero water level corresponds to the average sea level over the last 15 years. *Variación del nivel del agua (cota en m sobre el nivel medio del mar) en la laguna Fra Ramon ("Aiguamolls del Baix Ter") durante cuatro ciclos hidrológicos (desde 1999 a 2003, separados por líneas discontinuas). Las flechas indican temporales de mar. El nivel cero corresponde al nivel medio del mar en la zona durante los últimos 15 años.*

changes in water ionic composition. The water is poor in nutrients, especially in phosphorus, due to its subterranean origin. High water turnover during flooding and a relatively short drying period lead to fast changes in nutrient composition only at the end of the hydroperiod.

The hydrology in the Empordà salt marshes is determined by sudden flooding, mostly caused by marine intrusions during sea storms, when waves cross the sand bar. At least one sea storm occurs every year, but in some unusual years (e.g. during the 2001–2002 cycle) several sea storms have been registered. Although sea storms are more frequent between autumn and spring, their appearance is very irregular and no seasonal pattern can be observed (Fig. 2). Flooding periods are followed by long periods of confinement, when the water level gradually decreases and salinity increases due to evaporation (Quintana, 2002b; Quintana *et al.*, 2004; Badosa *et al.*, in press). Neither the Espolla pond nor the Empordà salt marshes have any regular seasonal pattern in hydrology and ecological processes.

Nutrient dynamics is strongly affected by this sudden flooding–long confinement hydrology in the Empordà salt marshes. Nitrate content increases during flooding periods, whilst

confinement favours the accumulation of phosphorus and the loss of the inorganic forms of nitrogen, most likely caused by denitrification (Badosa *et al.*, in press). The simultaneous accumulation of phosphorus and loss of nitrogen lead to especially low N/P ratios, which is called differential confinement of nutrients (Quintana *et al.*, 1998a). Water turnover and differential confinement also affects the accumulation of other compounds. Thus, some persistent pesticides and heavy metals tend to concentrate in some lagoons during confinement, whilst some more degradable pesticides appear only when the water turnover rate is high due to runoff (López-Flores *et al.*, 2003; Salvadó *et al.*, in press).

Moreover, a study of salinity and nutrient composition and their variations throughout the year carried out in several water bodies located in the area of the Empordà wetlands, reveals five types of shallow water ecosystems (Trobajo *et al.*, 2002): 1) hyperhaline wetlands, 2) brackish wetlands, 3) hypereutrophic freshwater wetlands; 4) meso-eutrophic freshwater wetlands; and 5) freshwater springs. Martinoy *et al.*, (in press) added more water body types, such as rice fields, estuarine lotic habitats, and ephemeral wetlands (Table 1). Salinity, water permanence, and nutrient composition appear to be the main environmental factors determining wetland classification and the shift between alternative stable states of macrophyte-periphyton-phytoplankton predominance. This is a common feature of most Mediterranean wetlands and shallow lakes (Britton & Podlejski, 1981; Alonso, 1998; Romo *et al.*, 2005; Fernández-Aláez *et al.*, 2004).

## COMMUNITY STRUCTURE OF AQUATIC ORGANISMS: TAXON-BASED APPROACHES

Classification of shallow lakes according to physical and chemical characteristics may be verified by their faunal composition. A study developed in Catalonia, within the framework of the implementation of the Water Framework

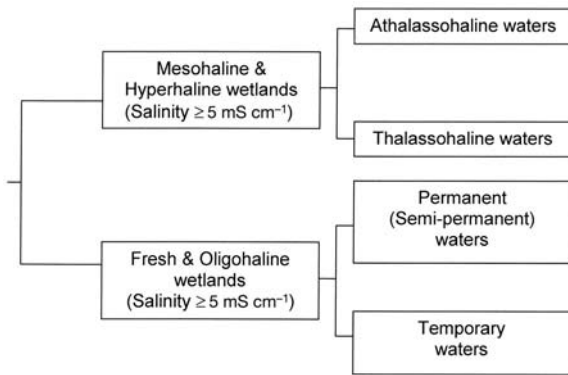
**Table 1.** Physical, chemical and biological characteristics of each water body type of the Empordà wetlands. Mean values in bold and coefficients of variation (%) in italics. Legend: HHW: hyperhaline wetlands; BW: brackish wetlands; HTW: hypereutrophic freshwater wetlands; MFW: meso-eutrophic freshwater wetlands; FS: freshwater springs; LW: lotic waters; EBW: ephemeral brackish wetlands; EFW: ephemeral freshwater wetlands; RF: rice fields; —: no data available. *Características físicas, químicas y biológicas de las diferentes tipologías de masas de agua de los humedales del Empordà. Valores medios en negrita y coeficientes de variación (%) en cursiva. Leyenda: HHW: humedales hiperhalinos; BW: humedales salobres; HTW: humedales hipertróficos de agua dulce; MFW: humedales meso-eutróficos de agua dulce; FS: surgencias; LW: sistemas lóticos; EBW: humedales salobres efímeros; EFW: humedales de agua dulce efímeros; RF: arrozales; —: datos no disponibles.*

	Waterbody types								
	HHW	BW	HTW	MFW	FS	LW	EBW	EFW	RF
Approximated hydroperiod length (months·year <sup>-1</sup> )	12	7-12	12	7-12	12	12	<2	<2	7
Conductivity (mS·cm <sup>-1</sup> )	<b>49.6</b> <i>48</i>	<b>18.9</b> <i>66</i>	<b>1.3</b> <i>136</i>	<b>3.9</b> <i>179</i>	<b>1.1</b> <i>99</i>	<b>0.8</b> <i>76</i>	<b>6.9</b> <i>96</i>	<b>1.4</b> <i>39</i>	<b>1.2</b> <i>39</i>
Chlorophyll- <i>a</i> (µg·L <sup>-1</sup> )	<b>21.5</b> <i>186</i>	<b>20.9</b> <i>183</i>	<b>12.1</b> <i>107</i>	<b>14.3</b> <i>150</i>	<b>18.1</b> <i>143</i>	<b>6.5</b> <i>103</i>	—	—	—
Ammonium (µM)	<b>9.1</b> <i>196</i>	<b>4.3</b> <i>260</i>	<b>34.1</b> <i>109</i>	<b>8.6</b> <i>245</i>	<b>2.6</b> <i>351</i>	<b>29.8</b> <i>214</i>	—	—	—
Nitrite (µM)	<b>0.7</b> <i>569</i>	<b>0.5</b> <i>234</i>	<b>6.9</b> <i>156</i>	<b>2.0</b> <i>289</i>	<b>5.5</b> <i>148</i>	<b>10.9</b> <i>193</i>	—	—	—
Nitrate (µM)	<b>2.1</b> <i>253</i>	<b>11.1</b> <i>312</i>	<b>109.9</b> <i>58</i>	<b>53.7</b> <i>151</i>	<b>705.9</b> <i>25</i>	<b>205.6</b> <i>100</i>	—	—	—
Phosphate (µM)	<b>4.3</b> <i>298</i>	<b>3.7</b> <i>253</i>	<b>5.8</b> <i>173</i>	<b>2.9</b> <i>174</i>	<b>0.93</b> <i>368</i>	<b>8.5</b> <i>159</i>	—	—	—
Total N (µM)	<b>219.3</b> <i>160</i>	<b>164.8</b> <i>94</i>	<b>212.7</b> <i>49</i>	<b>125.0</b> <i>75</i>	<b>851.9</b> <i>27</i>	—	—	—	—
Total P (µM)	<b>10.5</b> <i>248</i>	<b>9.2</b> <i>213</i>	<b>11.0</b> <i>83</i>	<b>6.1</b> <i>121</i>	<b>2.0</b> <i>197</i>	—	—	—	—

Directive and using CA multivariate analysis of invertebrate species composition, revealed four types of shallow lentic ecosystems, each one with a characteristic faunal composition (Boix *et al.*, 2005): 1) permanent (or semi-permanent) freshwater wetlands, including fresh or oligohaline waters (conductivity < 5 mS·cm<sup>-1</sup>); 2) temporary freshwater wetlands, with the same conductivities, which dry out every year; 3) thalassohaline wetlands, including temporary and permanent wetlands with average conductivities higher than 5 mS·cm<sup>-1</sup> and marine influence; and 4) athalassohaline wetlands, with conductivities higher than 5 mS·cm<sup>-1</sup>, not caused by marine intrusion, but by the endorheic concentration of salts (Fig. 3). The salt marshes of the Empordà

wetlands belong to thalassohaline ecosystem type, whilst the Espolla pond is an example of freshwater temporary ecosystem. It is worth pointing out that temporary and permanent thalassohaline water bodies have a similar faunal composition, but different community structures (Brucet *et al.*, 2005b; Gascón *et al.*, 2005).

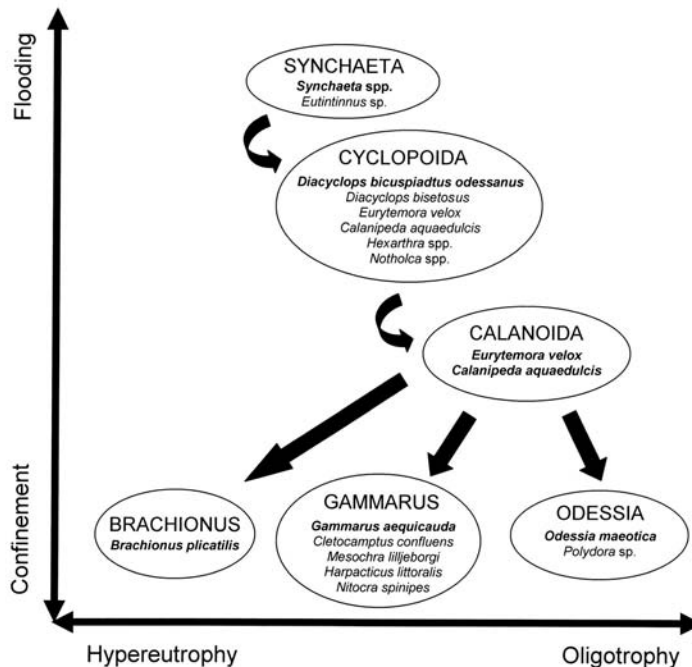
Successional patterns of aquatic invertebrates are related to hydrology. Although some changes are associated with seasonality, no regular seasonal pattern is observable in aquatic invertebrate dynamics, and changes in species composition are irregular throughout the year, as a consequence of the irregular hydrology (Quintana *et al.*, 1998b; Boix *et al.*, 2001a). In the Empordà salt marshes, several situations or phases have



**Figure 3.** Dichotomic diagram that identifies the main wetland types of Catalonia, modified from Boix *et al.* (2005). *Diagrama dicotómico que identifica los principales tipos de humedales de Catalunya, modificado de Boix et al. (2005).*

been identified in the zooplankton temporal pattern (Fig. 4), each one dominated by a characteristic species (Quintana *et al.*, 1998b; Brucet *et al.*, 2005b; Badosa *et al.*, in press): The rotifer *Synchaeta* spp. dominates at the beginning of the cycle just after an intense input of water, and is replaced soon after by copepod populations, for-

med by the cyclopoid *Diacyclops bicuspidatus odessanus* and nauplii of the calanoid *Eurytemora velox*. Later, when the water level decreases, late stages of *E. velox* may reach close to 100 % of the zooplankton total biomass. In spring the population of *E. velox* is substituted by another calanoid, *Calanipeda aquaedulcis*. During drought periods, benthic species such as the harpacticoid *Cletocamptus confluens* or the amphipod *Gammarus aequicauda* colonize the water column. Species composition during drought periods may differ depending on the degree of eutrophy. Under oligotrophic conditions, with high salinity and low N/P ratios, dense populations of the medusa *Odessia maeotica* and polychaete larvae (*Polydora* sp.) may appear. On the other hand, a drastic reduction of the water level in high eutrophic waters frequently causes incidents of hypereutrophy, where oxygen oversaturations during midday alternate with nocturnal anoxias. The rotifer *Brachionus plicatilis* and several species of ciliates dominate in these hypereutrophic situations.



**Figure 4.** Conceptual model of plankton succession in the salt marshes of the Empordà wetlands, adapted from Quintana *et al.* (1998b), Brucet *et al.* (2005c) and Badosa *et al.* (2006). Each group corresponds to different situations of the succession, characteristic taxa are shown, dominant taxa in bold. *Modelo conceptual de sucesión planctónica en la marisma de los “Aiguamolls de l’Empordà”, adaptado de Quintana et al. (1998b), Brucet et al. (2005b) y Badosa et al. (2006). Cada grupo corresponde a diferentes situaciones de la sucesión, se muestran los taxones característicos, y en negrita los dominantes.*

Similarly, heleoplankton succession in the temporary karstic pond of Espolla is related to the changes in the water turnover rate. Thus, just after flooding, when the water turnover is maximal, the ostracods *Heterocypris incongruens* and *Eucypris virens* are the dominant species of heleoplankton. If the high water turnover rate is maintained for approximately three weeks, the dominant species are substituted by the copepod *Megacyclops vireidis* and the cladoceran *Simocephalus vetulus*. Finally, when the water turnover rate is minimal (at the end of the hydroperiod), the more abundant species are the cladocerans *Daphnia pulicaria* and *Moina brachiata* (only in spring hydroperiods), and the copepod *Cyclops* sp.

Changes in hydrology also determine the physical and chemical composition of sediment and, hence, the spatial distribution of benthic species composition in the Empordà salt marshes: 1) non-disturbed permanent waters are characterized by a high density of individuals, total biomass and diversity; 2) non-disturbed temporary waters have lower richness and diversity values with little temporal variability; 3) disturbed temporary waters present a high taxonomic singularity and a high temporal variability of species richness and diversity (Gascón *et al.*, 2005). Moreover, changes in hydrology have different effects on planktonic and benthic communities, the former responding to temporal changes due to hydrological disturbances (Quintana, 2002) and the latter being more related to spatial patchiness in water permanence (Gascón *et al.*, 2005).

### Species' ecological requirements

The wide range of physical and chemical characteristics found in the different water bodies of the Empordà wetlands, as well as the existence of spatial and temporal gradients, allows for the ecological preferences of the different species found therein to be analysed, thus contributing to the knowledge of their autoecology. Similar patterns are found in organisms with very different characteristics, such as ostracods and periphytic diatoms (Gifre *et al.*, 2002; Trobajo *et al.*, 2004a). In both cases, the range of variation of water characteristics where the

different species are captured is very high for most species, especially for those species found in salt marshes. This seems to be common in other aquatic organisms that colonize these fluctuant environments (Boronat *et al.*, 2001). Fish such as *Aphanius iberus* tolerate wide ranges in salinity variations (Fernández-Delgado *et al.*, 1988; Vargas & Sostoa, 1997; Oltra & Todolí, 2000) and macrophytes such as *Ruppia cirrhosa* adapt their reproductive patterns to changes in water level (Gesti *et al.*, 2005). Some diatoms may also change their shape as a response to changing environmental conditions (Trobajo *et al.*, 2004b). Salinity, water turnover, water permanence, and productivity are described as the main factors determining species distribution in these ecosystems. In the case of the Empordà wetlands, the aquatic systems under salinity or trophic constraint (hyperhaline and hypereutrophic water bodies) are characterized by low values of species richness per visit and singularity, and by a higher proportion of species of crustaceans than of insects (Table 2).

Moreover, the interannual variability of these environmental factors is high in Mediterranean aquatic systems (Álvarez-Cobelas *et al.*, 2005; Britton & Crivelli, 1993). For example, hydroperiod length may vary considerably among years causing drastic interannual changes in the population abundance of some species that do not tolerate desiccation, such as the amphipod *Corophium orientale* (Gascón *et al.*, in press). Even population dynamics and reproduction of species that exclusively live in temporary waters are highly dependent on the flooding duration. This is the case of *Triops cancriformis* in the Espolla pond (Boix *et al.*, 2002).

### Biological indicators

The knowledge of species composition and its relation with trophic state variables, allow its use as an ecological indicator. To this effect, new tools for the assessment of ecological status have been successfully developed at a regional scale in Catalonia as required by the European Water Framework Directive (Directive 2000/60/EC). This includes the use of

**Table 2.** Species richness per visit, of crustaceans and insects for each water body type of the Empordà wetlands. Singularity, cumulative species richness, and crustaceans-insects ratio are also showed. Singularity was measured as the proportion of the number of species found only in one water body type in relation with total number of species of this water body type. Legend: HHW: hyperhaline wetlands; BW: brackish wetlands; HTW: hypertrophic freshwater wetlands; MFW: meso-eutrophic freshwater wetlands; FS: freshwater springs; LW: lotic waters. *Riqueza de especies por visita de crustáceos e insectos en cada tipología de masa de agua de los humedales del Empordà. También se muestra la singularidad, la riqueza acumulada de especies y la relación entre crustáceos e insectos. La singularidad se ha calculado como la proporción del número de especies encontradas únicamente en una tipología de masa de agua en relación con el número total de especies para esa tipología de masa de agua. Leyenda: HHW: humedales hiperhalinos; BW: humedales salobres; HTW: humedales hipertróficos de agua dulce; MFW: humedales meso-eutróficos de agua dulce; FS: surgencias; LW: sistemas lóticos.*

	Waterbody types					
	HHW	BW	HTW	MFW	FS	LW
<b>Species richness per visit</b>						
Mean	5	8	6	8	10	7
Maximum	10	18	11	19	19	19
Minimum	1	1	3	1	3	2
<b>Cumulative species richness</b>						
Crustacea	53	109	72	171	70	128
Insecta	35	62	48	76	24	47
	18	47	24	95	47	81
<b>Singularity</b>						
Within Crustacea	0.04	0.19	0.04	0.20	0.13	0.29
Within Insecta	0.00	0.21	0.06	0.11	0.04	0.06
	0.11	0.35	0.09	0.34	0.17	0.51
<b>Ratio</b>						
Crustacea/Insecta	1.9	1.3	2.0	0.8	0.5	0.6

aquatic crustaceans and insects as water quality indicators (Boix *et al.*, 2005) and the rapid assessment of the conservation status in terms of morphology, human use, water characteristics, and wetland vegetation (Sala *et al.*, 2004).

Species composition may also show other ecological aspects not related to water quality. Short term changes in zooplankton species composition have been used as a measure of disturbance intensity in the Empordà salt marshes, indicating a strong dependence of the zooplankton community structure on hydrological changes (Quintana, 2002a). This relationship between community structure and disturbance intensity allows the testing of Connell's (1978) intermediate disturbance hypothesis, which postulates that maximum diversities are found at intermediate disturbances. Considering the results obtained in the Empordà salt marshes, this hypothesis may be reformulated as follows: at intermediate severity, the maximum links between species are found (Quintana, 2002a). The

influence of hydrological disturbances to species composition in the Empordà salt marshes is not only restricted to planktonic communities. The Maturity index (MI) developed by Bongers (1990), applied to assess the disturbance effects on meiobenthic nematode assemblages, indicates that hydrological disturbances are also the main cause of variation in benthic communities (Gascón *et al.*, in press).

## SIZE BASED APPROACHES

During the last decades several authors emphasized the importance of body size as a community-organising unit in marine and freshwater ecosystems (Platt & Denman, 1977; Strayer, 1991; Rodríguez, 1994; Kerr & Dickie, 2001). It is based on the allometric relationship between body size and variables describing the main ecological processes, such as respiration, production, growth rate, and ingestion rate. The

representation (on a logarithmic scale) of abundance as a function of the size of individuals, what is called the biomass-size spectrum, gives rise to a straight line with a slope close to -1. Such a slope is achieved in many different environments, such as pelagic marine ecosystems, lakes, or shallow fluctuating ecosystems (Rodríguez & Mullin, 1986; Gaedke 1992; Gilabert, 2001; Quintana *et al.*, 2002). Besides the ecological relevance of this regularity, these authors also emphasized the interest in studying irregularities of the size-spectrum, rather than its regularity, for a more dynamic interpretation of the changes in size distribution relating to changes in environmental conditions.

Several reasons reveal the interest in studying the community structure using a size-based approach in aquatic communities of shallow lentic water bodies:

1. The different developmental stages of a single species may have different trophic roles. Changes in diet between stages are known for many copepods (Paffenhöfer, 1971, Poulet, 1977; Fernández, 1979). Differences in diet are found in the most abundant calanoids of the Empordà salt marshes (Compte unpublished data), and changes in amino acid composition during ontogeny, which may be related to changes in the diet (Guisande *et al.*, 2003), have been described for these species (Brucet *et al.*, 2005a).
2. Individuals of different species but of the same size may feed on the same preys, thus occupying the same trophic niche. Zooplankters found in the Empordà salt marshes (Quintana *et al.*, 1998b; Brucet *et al.*, 2005b, Badosa *et al.*, 2006) often belong to species with a relatively similar feeding ecology.
3. Dominances of a single species of zooplankter are very frequent in shallow lentic ecosystems, like the Empordà salt marshes (Quintana *et al.*, 1998b and 2004; Brucet *et al.*, 2005b). Under conditions of dominance, intraspecific competition acquires more relevance, favouring trophic niche partitioning among different stages of the same species.

### The biomass-size spectrum

Studies on biomass-size spectrum consist in the representation of abundance (either the number of individuals or biomass) as a function of body size (Rodríguez, 1994). This size distribution may be adjusted to a linear or non-linear model. Thereafter, changes in the model parameters, such as the fit ( $r^2$ ) or the slope are described as having ecological meaning. The biomass-size spectrum most commonly used is the normalised biomass-size spectrum, where abundance is represented as a function of the size of the individuals, classified in size classes of regular amplitude. The inconvenience of this procedure is that it produces a discrete model, which is dependent on the choice of the size scale and the interval amplitude (Han & Straskraba 1998). Vidondo *et al.* (1997) proposed the use of the underlying Pareto distribution, which is a probability density function and has the advantage of being a continuous function, where individuals should not be classified in size classes. The Pareto distribution has a cumulative distribution of probability defined as,

$$\text{prob}(s \geq S) = K^c s^{-c} \quad (1)$$

where  $\text{prob}(s \geq S)$  is the probability that a size ( $s$ ) of an individual taken at random will be greater than a threshold size  $S$ , expressed as a function of  $s$ . In practice, the term  $\text{prob}(s \geq S)$  is calculated for each individual as the fraction of all individuals larger than or equal to itself ( $N_{s \geq S}/N_t$ ). This model is linear at the logarithmic scale. Furthermore, the same authors define a more generalized non linear model, the Pareto type II model, which differs from the original Pareto model by the additive constant  $D$ ,

$$\text{prob}(s \geq S) = (K+D)^c (s+D)^{-c} \quad (2)$$

The ordinary Pareto or Pareto type I is a special case of the second type when  $D=0$ .

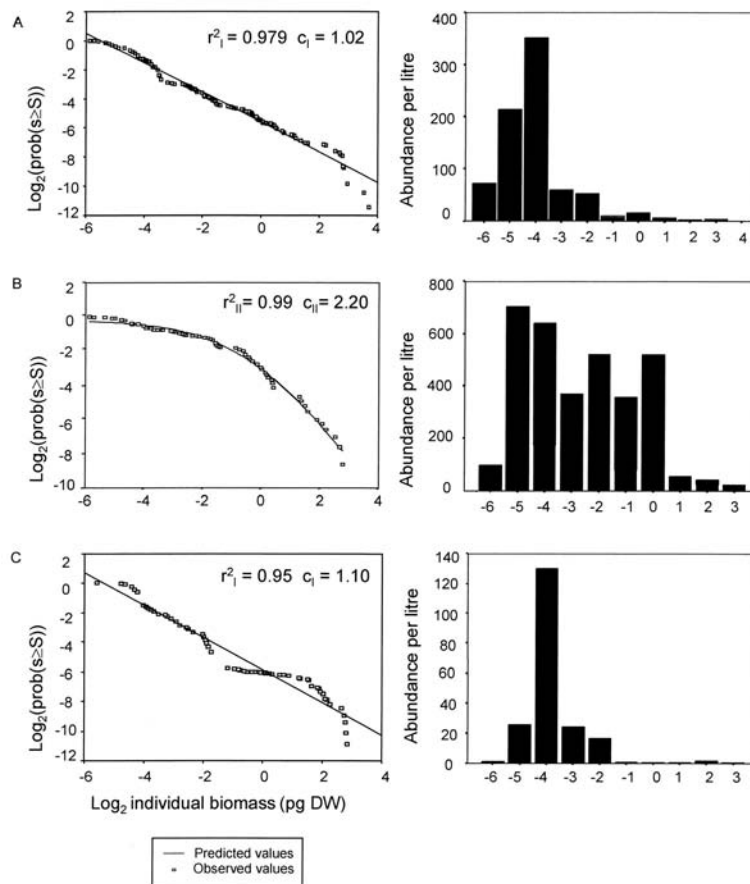
Several scalings in biomass-size spectra were described (Dickie *et al.* 1987). The primary or physiological scaling includes the size spectrum of all aquatic organisms (from bacteria to fish).



The secondary or ecological scaling identifies regular sub distributions of the biomass-size spectrum for the whole community, covering well-defined size ranges, which generally correspond to ecological functional groups. A tertiary or population scaling is also identifiable when the spectrum shows a step pattern on the ecological scaling, generally associated with scant overlap in population size ranges.

Studies on biomass-size spectrum at the primary or physiological scaling in the Empordà salt

marshes show that size distribution mainly depends on hydrology. Hydrological disturbances cause a decrease in the fit ( $r^2$ ) of the biomass-size spectrum, and an increase in the slope. The slope also changes depending on the degree of eutrophy (Quintana *et al.*, 2002). When only one functional group is considered (secondary or ecological scaling), for example the zooplankton community, three different types of size distribution are shown (Fig. 5) depending on environmental conditions (Brucet *et al.*, 2005c). During

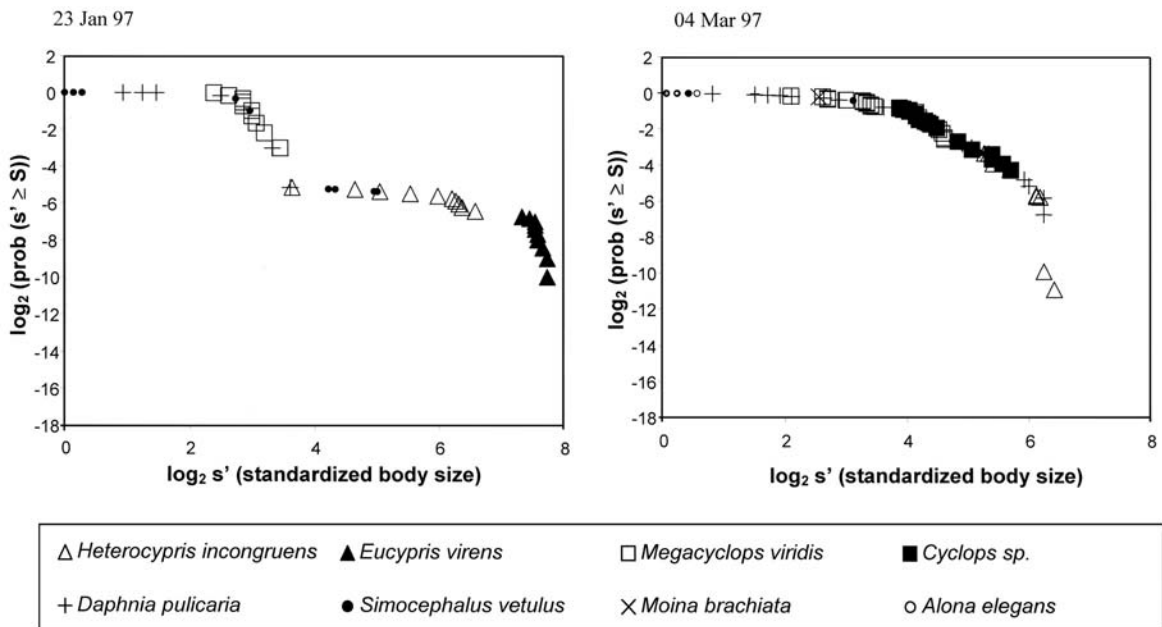


**Figure 5.** Types of biomass size spectra found in zooplankton samples and their fit to the Pareto model (left hand);  $r^2_{I}$  and  $r^2_{II}$  are the fit to the Pareto type I and the Pareto type II, respectively;  $c_I$  and  $c_{II}$  are the constant  $c$  of the Pareto type I and the Pareto type II models, respectively. (A) Linear spectrum, with better fit to linear model (Pareto type I); (B) curved spectrum, with better fit to non-linear model (Pareto type II); (C) example of step-like substructures. On the right, the graph that shows individuals abundance of each size class. All spectra are from a coastal lagoon in the Alt Empordà wetlands, modified from Brucet *et al.* (2005c). *Tipos de espectros de biomasa encontrados en muestras de zooplancton y su ajuste al modelo de Pareto (izquierda);  $r^2_{I}$  y  $r^2_{II}$  son el ajuste al modelo de Pareto tipo I y de Pareto tipo II, respectivamente;  $c_I$  y  $c_{II}$  son la constante  $c$  del modelo de Pareto tipo I y tipo II, respectivamente. (A) Espectro lineal, con un mejor ajuste al modelo lineal de Pareto (Pareto tipo I), (B) espectro curvado, con un mejor ajuste al modelo no lineal (Pareto tipo II), (C) ejemplo de subestructuras escalonadas. A la derecha el gráfico correspondiente que muestra la abundancia de individuos por clase de tamaño. Todos los espectros son de una laguna costera de los "Aiguamolls del Alt Empordà", modificado de Brucet *et al.* (2005c).*

flooding conditions, and when competitive interactions are weak, size spectra are linear shaped, with a high relative abundance of small sizes. Curved shaped spectra, with a higher relative abundance of larger sizes, are found during confinement conditions, when there is a lack of nutrient inputs and competitive interactions are expected to be stronger. Finally, stepped shapes with more than one mode appear whenever an increase of biomass takes place in a particular size range and are attributable to a variation in the population dynamics of the species involved. This subscale makes up the tertiary or population scaling, according to Dickie *et al.* (1987).

These patterns in spectra scaling are also easily identifiable in the size distributions in the Espolla pond. Most of its Pareto size distributions show biomass distributed in two groups, which make up the secondary scaling on the biomass spectrum: microcrustaceans (cladocerans, copepods, and ostracods) and macrofauna (turbellarians, oligochaetes, macrocrustaceans,

insects, gastropods, and amphibians). The inter-group minimum or swerve point of these cumulative distributions corresponds in all cases to an overlap zone of larger microcrustaceans (ostracods) and smaller macrofauna (turbellarians). In some cases, the tertiary scaling is also observable, especially at the beginning of the hydroperiod, when the initial hatching of the pioneering species takes place. In these circumstances the populations are composed of individuals of similar size, with population sizes well differentiated by species (Fig. 6). This tertiary scale has been related to a fractal nature of the biomass-size spectrum (Margalef, 1991). At the same time, the posterior disappearance of this tertiary scale could be interpreted as a self-organizing behaviour of biomass-size spectrum (Camacho & Solé, 2001). Thus, both of the theoretical biomass-size spectrum structure characteristics, the fractal, and the self-organizing, may be identified in tertiary scaling of size distribution along succession.



**Figure 6.** Heleoplankton size distribution on two sampling days, showing the species composition in the Espolla pond. On the first example (23 Jan 97) a tertiary-scale structure appears due to a lack of size overlap among species, which disappears in the second example (04 Mar 97). Body size has been standardized dividing each value by the minimum size value found. *Distribución de tamaños del heleoplankton en dos días de muestreo, mostrando la composición de especies en la laguna de Espolla. En el primer ejemplo (23 enero 97) se observa una estructura de escala terciaria debida a la falta de solapamiento en los tamaños de las diferentes especies, que desaparece en el segundo ejemplo (04 mayo 97). El tamaño corporal se ha estandarizado dividiendo su valor por el mínimo valor de tamaño encontrado.*

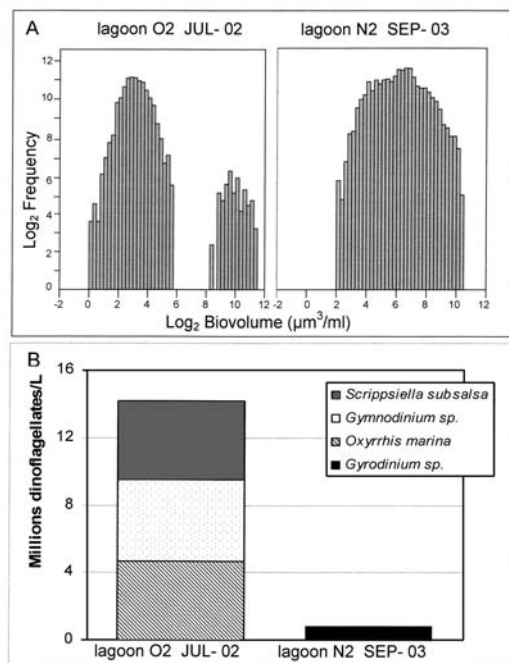
## COMBINATION OF SIZE-BASED AND TAXON-BASED APPROACHES

An approximation based on organism size is complementary to a taxonomic approach, not redundant (Strayer 1991; Rodríguez & Magnan 1993). Thus, the combination of taxon-based and size-based approaches in the study of community structure leads to a better understanding of the ecological functioning of aquatic ecosystems. A comparison of the zooplankton community structure in permanent and temporary water bodies of the Empordà salt marshes, show that they do not differ in their most abundant species, but they do differ in their temporal patterns and size structure (Brucet *et al.*, 2005b). These differences are a consequence of the ecological interactions of the species involved, such as competence and predation, where size acquires great relevance.

The combination of size-based and taxon-based approaches is especially suitable in the study of phytoplankton communities because of the different techniques currently used, such as observation by inverted microscope, identification of pigment contents by HPLC, or particle counting by flow cytometry, some of them more suitable for species (or functional groups) determination, others more suitable for counting. Studies on phytoplankton composition in the Empordà salt marshes show that dinoflagellates (and other potentially phagotrophic organisms such as haptophytes) play an important role in these ecosystems, dominating when nutrients are mainly found in organic form. Furthermore, phytoplankton biomass-size distributions frequently show a lack of intermediate sizes (2.5 - 4  $\mu\text{m}$  Equivalent Spherical Diameter), and its magnitude correlates with dinoflagellate biomass (Fig. 7). These results suggest that dinoflagellates take advantage of their mixotrophy by competing with, and grazing on, smaller phytoplankters simultaneously (López-Flores *et al.*, in press 1). Taxon-based approaches reveal that dinoflagellate species composition and dynamics of these coastal marshes differ from that observed in adjacent open coastal waters (López-Flores *et al.*, in press 2).

The study of the successional patterns found in the invertebrate community of the Espolla

pond is another example of the complementarity of size-based and taxon-based approaches. Boix *et al.* (2004) describe a 3-phase model of faunal succession in the Espolla pond by means of a taxon-based approach, which agrees with succession models described in several previous works for temporary ponds around the world (Kenk 1949; Barclay 1966; Lake *et al.*, 1989; Bazzanti *et al.*, 1996). The first and the last phases correspond to two allogenic succession processes. Flooding involves an available aquatic space with a trophic resource, low competition and low predation, while the disappearance of the flooded surface during drying periods involves changes in the environmental parameters (e.g. higher temperature values and oxygen fluctuations) and in the animal community (higher density values leading to higher competition



**Figure 7.** (A) Two kinds of phytoplankton biomass size distributions (sizes between 0 and 10  $\mu\text{m}$ ) in two lagoons of la Pletera salt marshes (Baix Ter wetlands). One gapped (left) and one unimodal distribution (right). (B) Abundances of the different dinoflagellate species present in the same samples, modified from López-Flores *et al.* (In press, 1). (A) *Dos tipos de distribuciones de biomasa (tamaños entre 0 y 10  $\mu\text{m}$ ) en dos lagunas de las marismas de la Pletera ("Aiguamolls del Baix Ter"). Una distribución discontinua (izquierda) y una unimodal (derecha).* (B) *Abundancias de las diferentes especies de dinoflagelados presentes en las mismas muestras, modificado de López-Flores et al. (en prensa, 1).*

and predation). Contrarily, the succession process observed between these two phases is autogenic. The size-based approach distinguishes five phases, which could be related to the three phases of the taxon-based approach, with the intermediate autogenic phase being divided into three additional phases, based on distinct community size structures, which appear throughout the autogenic succession. Observed changes in the middle phase are therefore more related to the size-based structure than to taxa abundance.

### Size diversity and species diversity

One of the main difficulties in the study of biomass-size spectra is the fact that the ecological meaning of the parameters of the size distribution models is normally difficult to interpret. The meaning of a change in the fit to a given model or a change in the slope is not trivial, and great effort is necessary to understand its ecological relevance. This is especially true when size distributions are adjusted to non-linear models, where several parameters are involved. The use of size diversity, that is the computation of a Shannon-Wiener index classifying individuals by their size instead of by their taxon, may solve these difficulties. A size diversity measure gives a unique value per size distribution and has the advantage of a more intuitive interpretation, since the concept of diversity and its ecological meaning is well established. However, a methodological problem emerges when measuring size diversity by clustering the different body sizes in size classes, because the division of a continuum variable (size) into an arbitrarily selected number of size classes is needed. As a result, different size diversity values for the same community were obtained depending on the number of size class intervals chosen (Ruiz, 1994). In order to overcome these problems, Lurie & Wagensberg (1984) proposed a measure of size diversity index appropriate for continuous variables ( $\mu$ ), based on the function of probability density of individuals with respect to size, where the summatory of the Shannon function is substituted by the integral of a continuous function of probability density  $p_i(s)$ ,

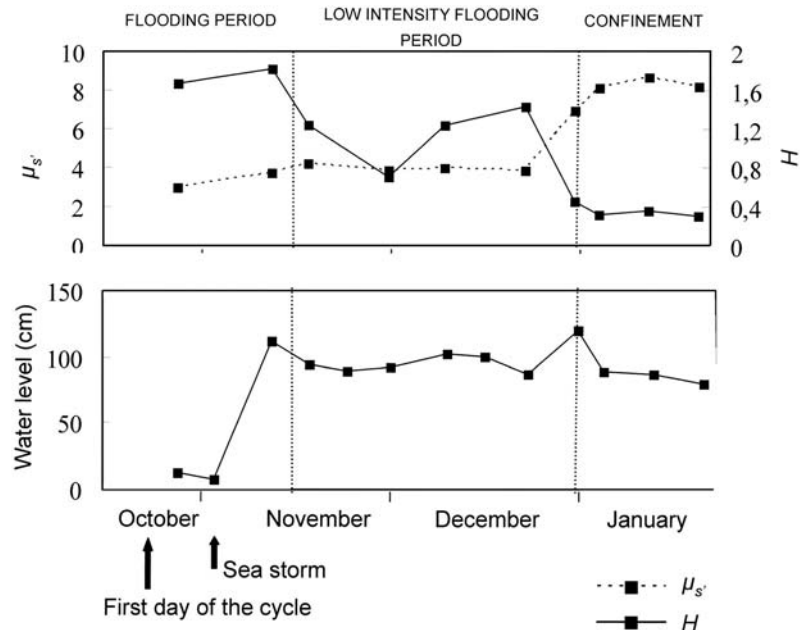
$$\mu = - \int p_i(s) \lg_2 p_i(s) ds \quad (3)$$

We have used this type of function and the results obtained are compared with those of the Shannon-Wiener index of species diversity (Brucet *et al.*, 2006).

Species diversity and size diversity give complementary information about zooplankton community structure. Relationships found between both diversities and several environmental variables (Table 3) suggest that the size diversity is mainly related to trophic interactions, such as fish predation when fish densities are high, or inter/intraspecific competition for food availability when fish are absent. On the other hand, species diversity appears to be more sensitive to abiotic factors such as trophic state. Differences among size and species diversity values are

**Table 3.** Pearson correlations between Shannon-Wiener diversity ( $H$ ), size diversity ( $\mu_s$ ) and environmental variables related to the trophic state (ammonium, soluble reactive phosphorus, total nitrogen, total phosphorus and total organic carbon), and variables related to species trophic interactions (fish density and food resource availability) of la Pletera (Baix Empordà wetlands). All these variables have previously been  $\log_{10}(x+1)$  except both diversity indexes. Significant correlation coefficients and non-significant relationships are indicated (\*  $p < 0.05$ , \*\*  $p < 0.01$ , n.s.  $p > 0.05$ ). Food resource availability was determined as a quotient between the [bacterial + phytoplankton] biovolume (in  $\mu\text{m}^3$ ) and the zooplankton biomass (in  $\mu\text{g}$  dry weight). *Correlaciones de Pearson entre la diversidad de Shannon-Wiener ( $H$ ), la diversidad de tamaños ( $\mu_s$ ) y variables ambientales relacionadas con el estado trófico (amonio, fósforo reactivo soluble, nitrógeno total, fósforo total y carbono orgánico total) y variables relacionadas con las interacciones tróficas (densidad de peces y disponibilidad de recurso) de la Pletera ("Aiguamolls del Baix Empordà"). Se indican los coeficientes de correlación significativos así como las relaciones no significativas (\*  $p < 0.05$ , \*\*  $p < 0.01$ , n.s.  $p > 0.05$ ). La variable disponibilidad de recurso se ha obtenido a partir del cociente entre biovolumen de [bacterio- + fitoplancton] (en  $\mu\text{m}^3$ ) y la biomasa de zooplancton (en  $\mu\text{g}$  de peso seco).*

	$H$	$\mu_s$
Ammonium (mg N·l <sup>-1</sup> )	- 0.299*	n.s.
Soluble reactive phosphate (mg P·l <sup>-1</sup> )	- 0.414**	n.s.
Total nitrogen (mg N·l <sup>-1</sup> )	- 0.395**	n.s.
Total phosphorus (mg P·l <sup>-1</sup> )	- 0.353**	n.s.
Total organic carbon (mg·l <sup>-1</sup> )	- 0.402**	n.s.
Fish density (ind·m <sup>-2</sup> )		
Low fish density lagoons	n.s.	n.s.
High fish density lagoons	n.s.	- 0.986**
Food resource availability	0.314*	- 0.427**



**Figure 8.** (A) Example of the evolution of size diversity ( $\mu_s$ ) and species diversity ( $H$ ) in a temporary lagoon. (B) Evolution of water level for the same cycle. Discontinuous lines separate three periods: the first correspond to a period of intense disturbances (arrows indicate flooding events with more than 100 % in water level increase); the second is a flooding period with low and irregular water inputs (water level increase always less than 40 %), and the third is a confinement period without water inputs, which causes a progressive decrease in water level. Modified from Brucet *et al.* (2006). (A) *Ejemplo de la evolución de la diversidad de tamaños ( $\mu_s$ ) y la diversidad de especies ( $H$ ) en una laguna temporal.* (B) *Evolución del nivel del agua durante el mismo ciclo. Las líneas discontinuas separan tres periodos: el primero corresponde a un periodo de perturbaciones intensas (las flechas indican inundaciones en las que el incremento del agua es mayor al 100 %), el segundo es un periodo de entradas de agua irregulares y de baja intensidad (incremento del agua siempre menor al 40 %), y el tercero es un periodo de confinamiento sin entrada de agua, que causa una disminución progresiva del nivel del agua. Modificado de Brucet *et al.* (2006).*

especially notable when they are compared along the succession gradient in fishless basins (Fig. 8). Species diversity increases after flooding, but decreases along the succession, when exploitative competition leads to the dominance of only one calanoid species. This agrees with the postulates of Connell's (1978) intermediate disturbance hypothesis. Besides, size diversity increases along the succession and the highest size diversity values coincide with the above-mentioned dominance of calanoids (Brucet *et al.*, 2006).

Comparing the behaviour of both diversity indexes lead to the conclusion that the combined use of size and species diversity may give a better understanding of ecological processes ruling the community structure. A wider interpretation of the concept of diversity, beyond the classical interpretation limited to species diversity was proposed earlier by Margalef (1980

and 1991), as discussed in Rodríguez (2005). Margalef pointed out the importance of size and turnover in diversity spectra and their relationship with some ecological aspects, such as those related to the transference of energy. Thus, all our work is only a small part of Margalef's heritage, being simply the development of Margalef previously proposed concepts.

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