#### AQUATIC CONSERVATION: MARINE AND FRESHWATER ECOSYSTEMS

Aquatic Conserv: Mar. Freshw. Ecosyst. 16: 469-482 (2006)

Published online in Wiley InterScience (www.interscience.wiley.com). DOI: 10.1002/aqc.769

# VIEWPOINT

# Alternatives to taxonomic-based approaches to assess changes in transitional water communities

D. MOUILLOT<sup>a,\*</sup>, S. SPATHARIS<sup>b</sup>, S. REIZOPOULOU<sup>c</sup>, T. LAUGIER<sup>d</sup>, L. SABETTA<sup>e</sup>, A. BASSET<sup>e</sup> and T. DO CHI<sup>a</sup>

<sup>a</sup> UMR CNRS-UMII 5119 Ecosystèmes Lagunaires, Université Montpellier II, Montpellier, France

<sup>b</sup> Department of Marine Sciences, University of the Aegean, Mytilene, Greece

<sup>c</sup> Hellenic Centre for Marine Research (HCMR) Institute of Oceanography, Anavissos, Greece

<sup>d</sup> French Research Institute for Exploitation of the Sea (Ifremer), Sète, France

<sup>e</sup> Department of Biological and Environmental Sciences and Technologies, University of Lecce, Lecce, Italy

## ABSTRACT

- 1. Transitional waters, described as critical transition zones because of their position at terrestrial, freshwater and marine interfaces, provide essential goods and services to the biosphere including human populations. These ecotones face increasing human influence mainly due to population density increase in coastal areas.
- 2. Transitional water bodies have, to date, received little attention in the development of ecological status indicators; this is a critical deficiency when trying to meet the Water Framework Directive objective of all significant water bodies achieving good ecological status by the year 2015.
- 3. In order to assess changes in transitional water communities many taxonomic-based indicators have already been proposed but there are a number of concerns for their use such as taxonomic classification difficulties, their unsuitability for multi-site comparisons and their inconsistent relationships with disturbance or stress.
- 4. Alternative methods based on body size, abundance distribution among functional groups, functional diversity and productivity descriptors are proposed. These methods offer the opportunity to compare sites with different taxonomic compositions and allow derivation of indicators related to ecological status of communities under scrutiny.
- 5. Finally, the suitability of these taxonomic-free descriptors to provide relevant information for each of the four main biotic compartments in coastal lagoons is discussed. The use of biomass distribution among functional groups for fish, benthos and macrophyte and to use body-size distribution for benthos and plankton is proposed.

Copyright © 2006 John Wiley & Sons, Ltd.

KEY WORDS: Water Framework Directive WFD; body size; functional traits; functional diversity; productivity; biotic descriptors

<sup>\*</sup>Correspondence to: David Mouillot, UMR CNRS-UMII 5119 Ecosystèmes Lagunaires, Université Montpellier II cc 093, 34095 Montpellier Cedex 05, France. E-mail: mouillot@univ-montp2.fr

## **INTRODUCTION**

The Water Framework Directive (WFD, 2000) provides an exciting challenge to scientists and environmental managers: all significant water bodies must achieve a good ecological status based upon physicochemical, hydromorphological, chemical and biological criteria by the year 2015. Behind the vague notion of good ecological status many fundamental aspects remain unresolved (Basset and Abbiati, 2004). What are the relevant ecological indicators of a good status? Which spatio-temporal scale must be defined to carry out the sampling programme? How does one classify the different status? Thus, in order to address the WFD purpose, the question is not yet whether the water bodies are in a good ecological status, but how to define a good ecological status. Even more modestly, how does one assess changes in water bodies' ecological status?

Among water bodies, transitional waters have been defined as Critical Transition Zones (CTZ) owing to their position at terrestrial, freshwater and marine interfaces (Levin *et al.*, 2001). These zones provide essential ecosystem services such as shoreline protection, water quality improvement, fisheries resources, habitat and food for migratory and resident animals and recreational areas for human populations (Levin *et al.*, 2001). These CTZ are also very productive ecosystems (Knoppers, 1994), but are under severe stress owing to human activities and climatic change inducing sea-level rise. Human impacts are mainly due to permanent and seasonal population density increases, aquaculture, fisheries, agriculture and industry. The increasing pressure inevitably leads to the expansion of environmental crises such as anoxia, over-exploitation of resources, destruction of habitat, eutrophication and pollutant contamination from the land-use watershed (Crooks and Turner, 1999).

Despite their obvious ecological and economical interests, transitional water bodies have received little attention about development of ecological status indicators. This is partly due to the inherent and extreme spatio-temporal variability observed in such ecosystems. Basset *et al.* (2001) point out that the daily range of nutrient concentration can reach the level of annual range in some extreme cases. This is the consequence of low inertia to external events owing to shallowness of water masses coupled with an interface position between marine and watershed water bodies leading to continuous perturbations such as tidal variations, winds, storms and chaotic freshwater discharges. For instance wind events are able to cause significant suspension of bottom materials injecting dissolved and particulate benthic materials into the water column leading to large increases in nutrient levels (Lawrence *et al.*, 2004). Thus monitoring nutrient concentrations or abiotic elements becomes a difficult task in such systems in order to assess an ecological status (Fano *et al.*, 2003). As a consequence physico-chemical parameters are supposed to provide supplementary information to the WFD but are not supposed to define ecological status on their own.

As an alternative to abiotic indicators, biocriteria have been developed and successfully applied in aquatic surveys (e.g. Lougheed and Chow-Fraser, 2002). However, most of them were implemented for continental (Garcia-Criado *et al.*, 1999; Nichols *et al.*, 2000) or marine water bodies (Borja *et al.*, 2000; Mouillot *et al.*, 2002; Simboura and Zenetos, 2002). When biotic indicators were proposed for lagoon or estuarine ecosystems they were mainly based on indicator species, taxon richness and diversity indices (Gibson *et al.*, 2000; Fano *et al.*, 2003; Panayotidis *et al.*, 2004); but these taxonomic-based indicators raise several problems. First of all, in addition to being time-consuming, identification of individuals to the species level is not an easy task for many groups owing to the size of specimens or to the presence of sibling or cryptic species (Knowlton, 1993), and many taxonomic errors may occur in ecological assessment studies (e.g. Sheppard, 1998). Moreover, the observed taxonomic richness is an underestimation of the true taxonomic richness (real number of species living on the studied site) and the error involved will depend on sampling effort (Gaston, 1996), and is rarely consistent among sites. Finally, the specific diversity or richness is difficult to relate to an ecosystem disturbance level because the diversity–disturbance relationship remains controversial and its shape is often unimodal with the highest diversity being observed for medium levels of disturbance or stress (Huston, 1979; Mackey and Currie, 2001). For these reasons,

community richness does not seem to be a useful indicator of disturbance in applied ecology (Drobner et al., 1998) because a low richness can be related to a high stress level with very few species able to survive in such demanding conditions or related to low stress and high productivity levels with only the best competitors able to persist (Connell, 1978; Huston, 1979). Furthermore, monitoring ecological status of transitional waters not only at the regional but also at the continental scale (e.g. combining in the same evaluation lagoons or estuaries from Portugal to Greece) inevitably leads to a new problem associated with taxonomic-based indicators. Species and lineages inhabiting transitional water bodies are likely to be different among locations to be compared (Basset et al., this issue). As a consequence ecological indices based on an indicator taxon will be influenced by anthropogenic constraints but also by the distance between sites. For instance a species supposed to reveal a good ecological status in Greece may be absent in Spain or Portugal because this species' geographic range is likely to be limited and not because the ecological status of transitional water body masses is different in the latter two countries.

Thus, even if the WFD does not require the use of similar indicators in all the countries, some intercalibration studies are necessary among sites with different taxonomic compositions. The severe flaws related to the use of richness or indicator species in order to assess the ecological status of transitional water body masses highlight the need for more integrated community-based indicators. Simpson (1949) suggested that the number of species only represents one aspect of species diversity, noting that the diversity of a given community also depends on the quantitative equilibrium between species, termed evenness. To take into account the community structure, numerous diversity or evenness indices have been proposed such as the Shannon-Weaver (1949) index and its associated evenness or the Simpson (1949) index. These diversity indices are classically used in ecology to assess the environmental impacts on ecosystems (e.g. Pires et al., 2000); however, some authors criticized the limited ability of these indices to discriminate between communities or to detect changes within a given community (Cao et al., 1996; Pires et al., 2000; Orfanidis et al., 2001). For instance, the Shannon diversity index appeared inadequate for impact assessment since it cannot discriminate natural stress from anthropogenic impact (Reizopoulou et al., 1996; Reizopoulou and Nicolaidou, 2004). Moreover, these classical species diversity indices do not take into account functional differences between species, though some authors pointed out the necessity of including these differences between species to estimate a diversity related to changes in environmental conditions or influencing ecosystem processes (Diaz and Cabido, 2001; Mouillot et al., 2005). Also, the inefficiency of the available multivariate techniques applied in transitional waters has been attributed to the fact that they were making use only of the existing information at the species level without taking into account their phylogenetic/ taxonomic relationships (Arvanitidis et al., 2005a,b). Thus, the most important question is not whether a proposed statistic summarizing a community satisfies some theoretical criterion, but whether it allows useful distinctions, with a functional meaning, between ecosystems to be made with field data.

Regarding the limitations listed above and induced by the taxonomic-based approach, some alternative CTZ systemic indicators have to be developed in order to address the main Water Framework Directive issues. The aim of this viewpoint is to propose an overview of alternative methodologies thus overcoming problems related to taxonomic ecological indicators for four main biotic components in coastal lagoons: fish, benthos, macrophyte and phytoplankton.

#### **BODY SIZE OR SIZE SPECTRA**

Body size is one of the most important traits of an organism determining, to a large extent, the type and the strength of ecological interactions to which individuals are subjected (De Roos *et al.*, 2003). More precisely, the physical structure of aquatic habitats has often been cited as a driver of body-size distributions in communities (Robson *et al.*, 2005) such as discontinuities in body-size distributions explained by habitat architecture (Holling, 1992). In addition, individual development could be density-dependent and

influenced by changes in environmental conditions (De Roos *et al.*, 2003). Body size is generally easy to measure and amenable to intercalibration procedures, it is comparable across taxa, guilds and sites, and, as a community feature, is expected to vary along disturbance gradients, according to energetic and ecological constraints (Basset *et al.*, 2004).

This taxonomic-free indicator was successfully used for some biotic components of transitional water body masses but is still under scrutiny for others. The most promising application comes from the benthos; the biomass profile of benthic communities has already been used to investigate community structure in coastal lagoons (Reizopoulou *et al.*, 1996; Lardicci and Rossi, 1998; Basset *et al.*, 2004); changes of benthic community biomass under disturbed conditions are well documented in benthic ecology (Pearson and Rosenberg, 1978; Warwick, 1986), since small-bodied invertebrates may characterize environments with high instability, and small body size could be a consequence of environmental/anthropogenic pressures imposed on organisms. As an illustrative example, a biometric index (Index of Size Distribution, ISD) was recently developed and applied as a new tool for quantifying the degree of disturbance in coastal lagoons (Reizopoulou and Nicolaidou, 2005). This kind of index may highlight alterations in benthic ecosystems since size variability of benthic organisms may decrease along pollution gradients (Reizopoulou and Nicolaidou, 2005).

Robust indicators describing the community structure such as size spectra in fish communities are also promising but are not yet well developed (Rochet and Trenkel, 2003), but size spectra in fish communities could be a suitable indicator allowing discrimination in coastal lagoon environments. In a different context, it is shown that the slope of the size spectrum decreases quasi-linearly with fishing mortality and that the curvature could help to detect ecosystem overexploitation (Shin and Cury, 2004).

Plankton size structure is also a common taxon-independent tool for the study of community and ecosystem structure of aquatic environments (Rodríguez et al., 1987; Chisholm, 1992; Quinones, 1994; Cavender-Bares et al., 2001). Size fractions (Sieburth, 1979) and size spectra, by grouping cells into logarithmic size classes (Rodríguez et al., 1998; 2002), are conventional descriptors of phytoplankton size structure. Various structural abiotic environmental factors were found to explain phytoplankton size structure; they include growth factors (Helbling et al., 1991), vertical water dynamics (Rodríguez et al., 2001; Serra et al., 2003), water depth (Gaedke, 1992) and trophic state (Glover et al., 1985; Tamigneaux et al., 1999; Yew-Hoong Gin et al., 2000; Maranon et al., 2001; Iriarte and Gonzalez, 2004).

Although there is evidence of changing phytoplankton size structure with respect to shifts in trophic conditions (Maranon *et al.*, 2001; Iriarte and Gonzalez, 2004), no methodology exists using phytoplankton biometric measures as a means for discriminating among trophic levels. In coastal marine areas, body-size spectra showed common patterns of variation according to major environmental forcing factors, independently of the taxonomic composition of phytoplankton (Sabetta *et al.*, 2005). Furthermore, for the purposes of the Water Framework Directive, new techniques such as flow cytometry allow rapid, reliable and reproducible data acquisition on phytoplankton size structure (Vaquer *et al.*, 1996; Rutten *et al.*, 2005). The regularity of phytoplankton size-abundance distributions can be translated into useful size-based descriptors of community or ecosystem structure and incorporated into environmental monitoring and conservation programmes for the purposes of the WFD, even though at this stage a methodological standardization of descriptors for ecological status classifications is lacking.

Macrophyte size spectra descriptors seem much less promising than other biotic components, as most of lagoon macrophytes consist of seagrasses and/or free-living macroalgae, which are not size-bounded. As a consequence, macrophyte size structure at the community level reflects more physical ecosystem attributes such as depth or mechanistic constraints (fragmentation) than disturbances. However, at the population level Hackney and Durako (2004) demonstrate the sensitivity of size-frequency distributions of the seagrass *Thalassia testudinum* morphometric characteristics to differences in environment factors in coastal areas. The challenge would be to determine a macrophyte species with a high morphometric plasticity to allow discrimination of coastal lagoon environments at the Mediterranean level.

In order to apply body size related descriptors to monitoring programmes it is important to clarify whether they are just phenomenological descriptors, indirectly related to ecosystem ecological status, or whether they are emergent properties of communities. Several authors have expressed concerns about this issue (Leaper et al., 2001; Robson et al., 2005), even though the question is no longer whether habitat architecture and environmental variables matter for body size distribution within communities, but how they matter and what are underlying mechanisms and assembly rules. The recent development of metabolic theory (Brown et al., 2004) confers the theoretical background to body size related descriptors of ecological status of ecosystems, i.e. ecosystem health, by relating body size constraints on patch selection and giving up behaviour to interspecific coexistence into a metabolic theory of coexistence (Basset, 1995). Accordingly, a tendency of body size to be positively related to population energy use has been observed (e.g. Maurer and Brown, 1988) and a relative invariance of size spectra compared to taxonomic composition was also detected in many different guilds of both macrobenthos (Bourassa and Morin, 1995; Basset et al., 2004) and plankton (Sprules and Munawar, 1986; Sabetta et al., 2005). Therefore, the ecological relevance of body size has already received strong experimental support; however, further research on transitional aquatic ecosystems is needed to test the relevance of the metabolic theory of coexistence (Basset, 1995) on the assembly rules responsible of the body size spectra shape at different spatial and temporal scales in such complex ecosystems.

## BIOMASS DISTRIBUTION AMONG FUNCTIONAL GROUPS

Many studies have highlighted the complexity of ecological systems and their fundamental unpredictability owing to multiple interactions (Huisman and Weissing, 2001). One way to overcome this problem is a simplification of communities through partitioning of species into a variety of guilds, functional groups or functional types (Simberloff and Dayan, 1991; Mathieson *et al.*, 2000; Blondel, 2003; Jauffret and Lavorel, 2003). Until now the guild concept has been more widely used than that of functional groups for animals. However, the guild concept refers exclusively to the mechanisms of resource-sharing by species in a competitive context. Within the framework of biodiversity and ecosystem functioning, a wide range of functions and responses are required from organisms and the 'functional groups' partitioning seems more appropriate than the 'guilds' one.

This approach seems particularly relevant for macroorganisms such as fish, benthos and macrophytes because these organisms can provide functional or life history traits allowing a classification of species into homogeneous functional units as a preliminary step. Species classification into functional groups is already well tested for fish (Dumay *et al.*, 2004; Nagelkerken and van der Velde, 2004) and benthos (Pearson, 2001; Gerino *et al.*, 2003) but not so commonly for macrophytes (Steneck and Dethier, 1994) and plankton. Based on functional form typology, Littler and Littler (1980) showed that macrophyte species within a functional group 'behave' similarly along stress-disturbance gradients regardless of their taxonomic or biogeographic features. Thus, algal-dominated communities, when examined at the functional group level, appear to be much more temporally stable and predictable than when examined at the species level. For plankton, a promising methodology is based on phytoplankton photopigments (Paerl *et al.*, 2003) that can be a surrogate for phytoplankton functional groups such as chlorophytes, cryptophytes, cyanobacteria, diatoms and dinoflagellates.

Shannon and Cury (2004) introduced an indicator of interaction strength (IS) which quantifies the effect that a change in biomass of one group has on abundance of other groups in pelagic fish communities. Nevertheless, these indices are more related to interactions than to environmental influences and do not correspond to the aim of the Water Framework Directive. A second example deals with the distribution of fish biomass along a gradient of trophic level which can be considered as a criterion for a functional classification because the 'trophic level', as a simple descriptor of the position of fish species in a food web,

is certainly a key component to study the functional role of the fish compartment in an ecosystem (Cruz-Escalona et al., 2000; Hajisamae et al., 2004). Based on these considerations, Sosa Lopez et al. (2005) developed a useful indicator (BDAtroph) that reflects changes in the trophic structure beyond simple changes of the mean trophic level. This indicator depends on information taken from FishBase 'ecology table' (Pauly and Christensen, 1995; Froese and Pauly, 2005) consisting of trophic level data for each fish species based on both the diet composition (trophDC) and food items (trophFI). The fish biomass was then arranged along trophDC intervals in order to build the general shape of the cumulated relative biomass (cumulated relative BDAtroph) smoothed by weighted least squares (McLain, 1972). The Kolmogorov-Smirnov test (Zar, 1997) was performed between two samples in order to assess differences in the general shape of the cumulated relative biomass along the trophic levels (BDAtroph). The apparent BDAtroph consistency across short-term periods suggests that this indicator could be considered as a useful quantitative ecological indicator when long-term comparisons between fish communities are performed (Sosa Lopez et al., 2005). The several shapes of the cumulated relative BDAtroph deserve to be evaluated more deeply in order to link them to specific processes such as top-down regulation of community dynamics contrasted with bottom-up factors affecting plant or animal prey availability. However, the proposed indicator could provide ecologists with the opportunity to address a variety of questions on changes and differences in fish community structure, and their relationships with environmental and human influences.

The recently developed Ecological Evaluation Index EEI (Orfanidis et al., 2001; Panayotidis et al., 2004) is based on the classification of macrophyte species in two Ecological State Groups, based on morphological and life strategy traits (Littler and Littler, 1980). The ESG I, characterizing pristine conditions, comprised the thick leathery, the articulate upright calcareous and the crustose calcareous species, most of them being K-selected species. ESG II, characterizing disturbed conditions, grouped the foliose, the filamentous and the coarsely branched upright species together, most of them being R-selected species. Sampling sites are classified in one of five Ecological Status Classes (bad, low, moderate, good, high) after a cross-comparison of the mean abundance value of the ESG I and II, on a matrix (Figure 1). Various applications of the index on macrophyte communities of the Aegean, as well as comparisons with other indices, showed that it is an efficient tool for revealing disturbed conditions (Spatharis et al., 2003; Panayotidis et al., 2004). However, its efficiency remains to be assessed for other European coastal areas as well.

Thus biomass distribution among functional groups can lead to the development of useful indicators within the Water Framework Directive context to help assess differences in coastal lagoon status. These indicators would themselves be taxonomic-free but would require considerable biological knowledge about the constituent species and their functional attributes in order to classify them into functional groups. Such

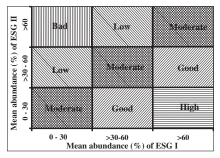


Figure 1. Matrix for the comparison of the mean abundances of macrophytes (percentage cover) classified in Ecological State Groups I and II.

D

D

Descriptor Descriptor	Biotic compartment			
	Fish	Benthos	Macrophyte	Phytoplankton
Body-size distributions	D	S	×	S
Biomass distribution among functional groups	S	S	S	D

S

D

D

D

Table 1. Suitability of four taxonomic free descriptors of four biotic compartments to assess changes in coastal lagoons (S: suitability with tools already implemented and tested, D: suitable but with more development needed, ×: not suitable)

an approach is already tried and tested for fish, benthos and macrophytes, and deserves to be implemented for phytoplankton (Table 1).

## **FUNCTIONAL DIVERSITY**

'Functional diversity', which was initially defined as the value and range of functional traits of the organisms present in a given ecosystem (Diaz and Cabido, 2001), has been measured in different ways, capturing different facets of this diversity such as the variation in the functional attributes of species (Walker *et al.*, 1999; Petchey and Gaston, 2002; Mason *et al.*, 2003), the complexity of food webs (Hulot *et al.*, 2000), the number of plant functional groups present (Diaz and Cabido, 2001) and the regularity of abundance distribution in functional trait space (Mouillot *et al.*, 2005). Thus a more general definition of functional diversity is clearly needed embracing these different components. Mason *et al.* (2005) recently redefined functional diversity as a measure (or group of measures) of the distribution of the species and abundance of a community in functional attribute space that represents: (a) the amount of functional attribute space filled by species in the community (functional richness), (b) the evenness of abundance distribution in filled niche space (functional evenness) and (c) the degree to which abundance distribution in niche space maximizes divergence in functional attributes within the community (functional divergence).

It is now generally accepted that functional diversity, rather than species diversity per se, is the key for ecosystem processes such as productivity, stability, resilience or invadability (Hulot *et al.*, 2000; Naeem and Wright, 2003; Petchey *et al.*, 2004; Roscher *et al.*, 2004; Waldbusser *et al.*, 2004). Most of experiments relating functional diversity to ecosystem processes were carried out on terrestrial plants, but two recent studies underline the positive effect of benthic functional diversity on organic matter processes in sediments (Waldbusser *et al.*, 2004; Mermillod-Blondin *et al.*, 2005), but has yet to address other biotic components such as fish, macrophytes and phytoplankton. Within the context of the Water Framework Directive, further investigation of the influence of environmental factors on functional diversity is needed in order to evaluate its ability to be a relevant indicator of ecological status.

Very few studies have dealt with this issue but Bremner *et al.* (2003) using benthic invertebrates, identified some consistent patterns of functional diversity across scales and Bady *et al.* (2005) demonstrated that functional diversity had a greater accuracy than species richness with less sampling effort. As an application Fano *et al.* (2003) developed the ecofunctional quality index (EQI) for the evaluation of environmental quality in lagoons using a set of data embracing primary productivity, seaweed and seagrass biomasses, structure and productivity of the benthic community, taxonomic diversity of macrozoobenthos, and finally, trophic complexity, expressed as macrozoobenthic functional diversity. However, this index requires a lot of information on different biotic components and must be tested on a large set of coastal lagoons with various taxonomic compositions.

Functional diversity

Productivity

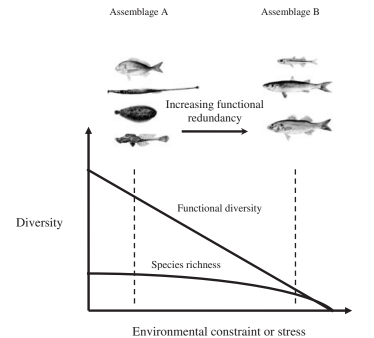


Figure 2. Theoretical relation between functional diversity, species richness and environmental constraints in fish assemblages. When environmental constraints increase functional redundancy, or similarity increases in assemblages through niche filtering (between A and B), functional diversity decreases at a higher rate than species richness.

From a theoretical point of view, we can suspect that functional diversity decreases with increasing environmental constraints or stress (Figure 2). When environmental constraints increase, coexisting species are more likely to be similar to one another because environmental conditions (i.e. abiotic properties of the habitat) act as a filter, allowing only a narrow spectrum of species to survive. The species that make it through the environmental filters are likely to share many biological/ecological characteristics through the niche filtering concept (Franzen, 2004; Statzner *et al.*, 2004). More precisely, environmental factors could limit the presence of certain functional traits at certain sites and thus decrease functional diversity of local communities in sites under environmental pressure such as confined parts in lagoons (Figure 2). For instance, increasing organic disturbance in coastal lagoons has been shown to lead to a decrease of macrobenthic size in three Mediterranean lagoons (Reizopoulou and Nicolaidou, 2005): small-bodied specimens were mainly represented by tolerant and opportunistic deposit feeders, while the larger-bodied specimens comprised mostly filter feeding bivalves, carnivorous polychaetes, etc.

The hypothesis that there is decreasing functional diversity along a disturbance gradient could be tested on the four biotic components. The issue that functional traits allow functional diversity estimation, however, must be resolved first, although this appears the case for benthic invertebrates (Bremner *et al.*, 2003; Fano *et al.*, 2003; Waldbusser *et al.*, 2004; Bady *et al.*, 2005; Mermillod-Blondin *et al.*, 2005) and for fishes (Sibbing and Nagelkerke, 2001; Dumay *et al.*, 2004; Nagelkerken and van der Velde, 2004) but still unresolved for macrophyte and plankton. There has been a recent effort to produce a functional classification of freshwater phytoplankton associations based on commonly shared adaptive features (Reynolds *et al.*, 2002) and to associate them with trophic states (Padisak *et al.*, 2003). Additionally, phytoplankton functional attributes such as cell volume, growth rate, and the ratio between minimum quotas of total *N* and total *P* have also been investigated (Seip and Reynolds, 1995).

Once the functional attributes are estimated for each species, many functional diversity indices can be applied, providing either an overall diversity index including all functional traits (Petchey and Gaston, 2002) or a univariate diversity index for each trait (Mason *et al.*, 2003, 2005; Mouillot *et al.*, 2005). These indices are not taxonomic-based since species identity and its taxonomy is never included in the analysis, allowing comparison between distant sites. Nevertheless more theoretical and empirical studies are awaited before considering functional diversity as a tool to assess changes in coastal lagoon ecological status.

## BIOMASS OR PRODUCTIVITY MEASURES

Biomass has often been used as a surrogate for productivity of primary producers, which is the most commonly cited ecosystem property in ecological studies (e.g. Loreau, 2000). Some statistical methodologies, which do not involve species identification but are based on chlorophyll *a* and nutrient concentrations in the water column, have been proposed for water quality assessment (Kitsiou and Karydis, 1998). As an application, the Joint Research Centre of the European Union (Druon *et al.*, 2004) is developing a risk index (EUTRISK) as an indicator of European coastal areas sensitive to eutrophication. EUTRISK is based on observations of phytoplankton biomass in the top layer of water column and determined as chlorophyll abundance by remote sensing. Nevertheless these methods contribute to the assessment of eutrophication levels but not to the assessment of biological status of water body masses as stated in the WFD. This is more the task of the 'Eutrophication activity' WFD working group that has not yet been completed.

Biomass density per se does not appear to be a reliable parameter for distinguishing between different ecological status of macrophyte communities. Indeed, along a disturbance gradient such as eutrophication, the amount of total primary production is roughly maintained (Schramm, 1999), and the shift concerns progressive regression of slow-growing perennials and final replacement by fast-growing ephemeral forms; suggesting that an indicator of biomass turnover rates, combining standing stocks and productivity, is a promising taxonomic-free approach.

In the classic model of benthic ecology of Pearson and Rosenberg (1978), benthic variables such as macrofaunal abundance or biomass along an organic enrichment gradient are expected initially to increase to a certain level and then to decrease. Thus, this non-monotonic indicator is not suitable to assess changes in benthic communities. Under disturbed conditions, the larger, long-lived species are the first to be eliminated and the communities are dominated by smaller, short-lived opportunistic species. A useful indicator could estimate the biomass turnover between these two sets of species.

A measure of fish biomass or density per surface area is still very problematic to obtain in coastal lagoons because the sampling methodology is far from homogeneous among regions. An active capture method, such as a drawnet, or a passive method using 'capetchade' or trammel net, can be used to give fish biomass or density. With the former methodology it is possible to sample the entire water column from the bottom to the surface and a certain area but the largest individuals and some species (Mugilidae) are almost impossible to catch, introducing a bias in the results. The passive method causes selectivity problems, i.e. some species are more easily caught than others, and thus the samples can be biased. In addition, there is the difficulty of relating an extremely variable indicator such as the biomass/density of fish to a given ecological status. A most promising tool is certainly to carry out experiments based on cages to measure mortality and growth of juveniles. This method may give insights into the ability of coastal lagoon environments to provide suitable conditions for juvenile fish to survive and grow, a fundamental service provided by coastal ecosystems (Baldo and Drake, 2002).

## **CONCLUSION**

Transitional waters, which provide essential goods and services to the biosphere, are facing increasing human influence mainly due to population density increases on coastal areas. Thus, within the European Water Framework Directive context, indicators of ecological status in order to classify different sites are required. Monitoring nutrient concentrations or abiotic elements is a very difficult task in such an environment because, owing to the shallowness of lagoons combined with the interface between marine and watershed water bodies that leads to continuous perturbations, extreme spatio-temporal variability was observed causing inconsistencies in impact assessment studies (Basset et al., 2001). As an alternative to abiotic indicators, biocriteria have been developed, mainly related to indicator species or taxonomic description of the communities (Gibson et al., 2000; Panayotidis et al., 2004). However, the current study highlights some concerns about the extensive use of such taxonomic-based indicators to classify water body masses. Individual classification into taxonomic units remains time-consuming and problematic for some groups such as plankton. Moreover, taxonomic richness is often assumed to represent a second order (unimodal) relationship with ecosystem productivity or disturbance negating its usefulness as a relevant indicator (Drobner et al., 1998). In addition, most species and taxa are not present in all the sites being compared. As a consequence we cannot relate the presence-absence of some taxon to a given ecological status in multi-site comparisons. Alternative descriptors of communities, based on taxonomic-free attributes: body size, abundance distribution among functional groups, functional diversity and productivity are proposed. Such methods overcome many problems related to taxonomic indicators and deserve to be applied in extensive comparative studies: for example, biomass distribution among functional groups for fish, benthos and macrophyte and the use of body size distribution for benthos and plankton. Functional diversity has the potential to define the ecological status of communities for every kind of organism, but more theoretical and conceptual work is needed.

## **ACKNOWLEDGEMENTS**

This paper results from the contributions of the participants to the Naples Lagunet Forum on 'Major challenges to bridge basic ecology to applications' (Naples, June 17–19, 2004). We want to thank all of them for their enthusiasm and the effort brought to the discussion of these topics. We also feel grateful to Anna Fauci and the staff of the Ecology group of the Naples Federico II University for their invaluable contribution to the organization of the Forum and to the Presidency of the Campania Region which provided the financial resources. A. Basset, L. Sabetta and S. Reizopoulou were supported by the EU-project TWReferenceNET; T. Do Chi and D. Mouillot were supported by the EU-project DITTY (EVK3 CT 2002 00084). Two anonymous referees and the Editor (J. Baxter) greatly improved the manuscript.

#### REFERENCES

- Arvanitidis C, Chatzigeorgiou G, Koutsoubas D, Dounas C, Eleftheriou A, Koulouri P. 2005a. Mediterranean lagoons revisited: weakness and efficiency of the rapid biodiversity assessment techniques in a severely fluctuating environment. *Biodiversity and Conservation* 14: 2347–2359.
- Arvanitidis C, Chatzigeorgiou G, Koutsoubas D, Kevrekidis T, Dounas C, Eleftheriou A, Koulouri P, Mogias A. 2005b. Estimating lagoonal biodiversity in Greece: comparison of rapid assessment techniques. *Helgoland Marine Research* **59**: 177–186.
- Bady P, Doledec S, Fesl C, Gayraud S, Bacchi M, Scholl F. 2005. Use of invertebrate traits for the biomonitoring of European large rivers: the effects of sampling effort on genus richness and functional diversity. *Freshwater Biology* **50**: 159–173.
- Baldo F, Drake P. 2002. A multivariate approach to the feeding habits of small fishes in the Guadalquivir Estuary. *Journal of Fish Biology* **61**: 21–32.

- Basset A. 1995. Body size-related coexistence: an approach through allometric constraints on home range use. *Ecology* **76**: 1027–1035.
- Basset A, Abbiati M. 2004. Challenges to transitional water monitoring: ecological descriptors and scales. *Aquatic Conservation: Marine and Freshwater Ecosystems* 14: S1–S3.
- Basset A, Carlucci D, Fiocca A, Vignes F. 2001. Water transparency and health of coastal salt marshes: simple enclosure experiments of nutrient dynamics. *Aquatic Conservation: Marine and Freshwater Ecosystems* 11: 273–279.
- Basset A, Sangiorgio F, Pinna M. 2004. Monitoring with benthic macroinvertebrate: advantage and disadvantages of body size descriptors. *Aquatic Conservation: Marine and Freshwater Ecosystems* 14: S43–S58.
- Blondel J. 2003. Guilds or functional groups: does it matter? Oikos 100: 223-231.
- Borja A, Franco J, Perez V. 2000. A marine Biotic Index to establish the ecological quality of soft-bottom benthos within European estuarine and coastal environments. *Marine Pollution Bulletin* 40: 1100–1114.
- Bourassa N, Morin A. 1995. Relationships between size structure of invertebrate assemblages and trophy and substrate composition in streams. *Journal of the North American Benthological Society* **14**: 393–403.
- Bremner J, Rogers SI, Frid CLJ. 2003. Assessing functional diversity in marine benthic ecosystems: a comparison of approaches. *Marine Ecology Progress Series* **254**: 11–25.
- Brown JH, Gillooly JF, Allen AP, Savage VM, West GB. 2004. Towards a metabolic theory of ecology. *Ecology* **85**: 1771–1789.
- Cao Y, Bark AW, Williams WP. 1996. Measuring the responses of macroinvertebrate communities to water pollution: a comparison of multivariate approaches, biotic and diversity indices. *Hydrobiologia* **341**: 1–19.
- Cavender-Bares KK, Rinaldo A, Chisholm SW. 2001. Microbial size spectra from natural and nutrient enriched ecosystems. *Limnology and Oceanography* **46**: 778–789.
- Chisholm SW. 1992. Phytoplankton size. In *Primary Productivity and Biogeochemical Cycles in the Sea*, Falcowski PG, Woodhead AD (eds). Plenum Press: New York.
- Connell JH. 1978. Diversity in tropical rain forests and coral reefs. Science 199: 1302–1310.
- Crooks S, Turner RK. 1999. Integrated coastal management: sustaining estuarine natural resources. *Advances in Ecological Research* 29: 241–289.
- Cruz-Escalona VH, Abitia-Cardenes LA, Campos-Davila L, Galvan-Magaña F. 2000. Trophic interrelations of the three most abundant fish species from Laguna San Ignacio, Baja California Sur, Mexico. *Bulletin of Marine Science* **66**: 361–373.
- De Roos AM, Persson L, McCauley E. 2003. The influence of size-dependent life-history traits on the structure and dynamics of populations and communities. *Ecology Letters* **6**: 473–487.
- Diaz S, Cabido M. 2001. Vive la difference: plant functional diversity matters to ecosystem processes. *Trends in Ecology and Evolution* **16**: 646–655.
- Drobner U, Bibby J, Smith B, Wilson JB. 1998. The relation between community biomass and evenness: what does community theory predict, and can these predictions be tested? *Oikos* 82: 295–302.
- Druon JN, Schrimpf W, Dobricic S, Stips A. 2004. Comparative assessment of large-scale marine eutrophication: North Sea area and Adriatic Sea as case studies. *Marine Ecology Progress Series* 272: 1–23.
- Dumay O, Tari PS, Tomasini JA, Mouillot D. 2004. Functional groups of lagoon fish species in Languedoc Roussillon, southern France. *Journal of Fish Biology* **64**: 970–983.
- Fano EA, Mistri M, Rossi R. 2003. The ecofunctional quality index (EQI): a new tool for assessing lagoonal ecosystem impairment. *Estuarine Coastal and Shelf Science* **56**: 709–716.
- Franzen D. 2004. Plant species coexistence and dispersion of seed traits in a grassland. Ecography 27: 218–224.
- Froese R, Pauly D. 2005. Fishbase world wide web electronic publication. www.fishbase.org [version 16 March 2004]. Gaedke U. 1992. The size distribution of plankton biomass in a large lake and its seasonal variability. *Limnology and Oceanography* 37: 1202–1220.
- Garcia-Criado F, Tome A, Vega FJ, Antolin C. 1999. Performance of some diversity and biotic indices in rivers affected by coal mining in northwestern Spain. *Hydrobiologia* **394**: 209–217.
- Gaston KJ. 1996. Biodiversity. Blackwell: Oxford.
- Gerino M, Stora G, Francois-Carcaillet F, Gilbert F, Poggiale JC, Mermillod-Blondin F, Desrosiers G, Vervier P. 2003. Macro-invertebrate functional groups in freshwater and marine sediments: a common mechanistic classification. *Vie et Milieu–Life and Environment* 53: 221.
- Gibson GR, Bowman ML, Gerritsen J, Snyder BD. 2000. Estuarine and Coastal Marine Waters: Bioassessment and Biocriteria Technical Guidance. EPA 822-B-00-024. U.S. Environmental Protection Agency, Office of Water, Washington, DC.
- Glover HE, Smith AE, Shapiro L. 1985. Diurnal variations in photosynthetic rates: comparisons of ultraphytoplankton size fraction. *Journal of Plankton Research* 7: 519–535.

- Hackney JW, Durako MJ. 2004. Size-frequency patterns in morphometric characteristics of the seagrass *Thalassia testudinum* reflect environmental variability. *Ecological Indicators* **4**: 55–71.
- Hajisamae S, Chou LM, Ibrahim S. 2004. Feeding habitats and trophic relationships of fishes utilizing an impacted coastal habitat, Singapore. *Hydrobiologia* **520**: 61–71.
- Helbling EW, Villafane V, Holm-Hansen O. 1991. Effect of iron on productivity and size distribution of Antarctic phytoplankton. *Limnology and Oceanography* **36**: 1879–1885.
- Holling CS. 1992. Cross-scale morphology, geometry, and dynamics of ecosystems. *Ecological Monographs* **62**: 447–502.
- Huisman J, Weissing FJ. 2001. Biological conditions for oscillations and chaos generated by multispecies competition. *Ecology* **82**: 2682–2695.
- Hulot FD, Lacroix G, Lescher Moutoue FO, Loreau M. 2000. Functional diversity governs ecosystem response to nutrient enrichment. *Nature* **405**: 340–344.
- Huston MA. 1979. A general hypothesis of species diversity. American Naturalist 113: 81-101.
- Iriarte JL, Gonzalez HE. 2004. Phytoplankton size structure during and after the 1997/98 El Nino in a coastal upwelling area of the northern Humboldt Current System. *Marine Ecology Progress Series* **269**: 83–90.
- Jauffret S, Lavorel S. 2003. Are plant functional types relevant to describe degradation in arid, southern Tunisian steppes? *Journal of Vegetation Science* **14**: 399–408.
- Kitsiou D, Karydis M. 1998. Development of categorical mapping for quantitative assessment of eutrophication. Journal of Coastal Management 4: 35-44.
- Knoppers B. 1994. Aquatic primary production in coastal lagoons. In *Coastal Lagoon Processes*, Kjerfve B (ed.). Elsevier Science Publishers: Amsterdam: 243–286.
- Knowlton N. 1993. Sibling species in the sea. Annual Review of Ecology and Systematics 24: 189–216.
- Lardicci C, Rossi F. 1998. Detection of stress on macrozoobenthos: evaluation of some methods in a coastal Mediterranean lagoon. *Marine Environmental Research* **45**: 367–386.
- Lawrence D, Dagg MJ, Liu HB, Cummings SR, Ortner PB, Kelble C. 2004. Wind events and benthic–pelagic coupling in a shallow subtropical bay in Florida. *Marine Ecology Progress Series* **266**: 1–13.
- Leaper R, Raffaelli D, Emes C, Manly B. 2001. Constraints on body-size distributions: an experimental test of the habitat architecture hypothesis. *Journal of Animal Ecology* 70: 248–259.
- Levin LA, Boesch DF, Covich A, Dahm C, Erseus C, Ewel KC, Kneib RT, Moldenke A, Palmer MA, Snelgrove P, Strayer D, Weslawski JM. 2001. The function of marine critical transition zones and the importance of sediment biodiversity. *Ecosystems* 4: 430–451.
- Littler MM, Littler DS. 1980. The evolution of thallus form and survival strategies in benthic marine macroalgae: field and laboratory tests of a functional form model. *American Naturalist* **116**: 25–44.
- Loreau M. 2000. Biodiversity and ecosystem functioning: recent theoretical advances. Oikos 91: 3-17.
- Lougheed VL, Chow-Fraser P. 2002. Development and use of a zooplankton index of wetland quality in the Laurentian Great Lakes basin. *Ecological Applications* 12: 474–486.
- Mackey RL, Currie DJ. 2001. The diversity-disturbance relationship: is it generally strong and peaked? *Ecology* **82**: 3479–3492.
- Maranon E, Holligan PM, Barciela R, Gonzalez N, Mourino B, Pazo MJ, Varela M. 2001. Patterns of phytoplankton size structure and productivity in contrasting open-ocean environments. *Marine Ecology Progress Series* **216**: 43–56.
- Mason NWH, MacGillivray K, Steel JB, Wilson JB. 2003. An index of functional diversity. *Journal of Vegetation Science* 14: 571–578.
- Mason NWH, Mouillot D, Lee WG, Wilson JB. 2005. Functional richness, functional evenness and functional divergence: proposed primary components of functional diversity. *Oikos* 111: 112–118.
- Mathieson S, Cattrijsse A, Costa MJ, Drake P, Elliott M, Gardner J, Marchand J. 2000. Fish assemblages of European tidal marshes: a comparison based on species, families and functional guilds. *Marine Ecology Progress Series* **204**: 225–242.
- Maurer BA, Brown JH. 1988. Distribution of energy use and biomass among species of North American terrestrial birds. *Ecology* **69**: 1923–1932.
- McLain DH. 1972. Drawing contours from arbitrary data points. The Computer Journal 14: 318–324.
- Mermillod-Blondin F, Francois-Carcaillet F, Rosenberg R. 2005. Biodiversity of benthic invertebrates and organic matter processing in shallow marine sediments: an experimental study. *Journal of Experimental Marine Biology and Ecology* **315**: 187–209.
- Mouillot D, Culioli JM, Do-Chi T. 2002. Indicator species analysis as a test of non-random distribution of species in the context of marine protected areas. *Environmental Conservation* **29**: 385–390.
- Mouillot D, Mason WHN, Dumay O, Wilson JB. 2005. Functional regularity: a neglected aspect of functional diversity. *Oecologia* **142**: 353–359.

- Naeem S, Wright JP. 2003. Disentangling biodiversity effects on ecosystem functioning: deriving solutions to a seemingly insurmountable problem. *Ecology Letters* **6**: 567–579.
- Nagelkerken I, van der Velde G. 2004. A comparison of fish communities of subtidal seagrass beds and sandy seabeds in 13 marine embayments of a Caribbean island, based on species, families, size distribution and functional groups. *Journal of Sea Research* **52**: 127–147.
- Nichols S, Weber S, Shaw B. 2000. A proposed aquatic plant community biotic index for Wisconsin lakes. *Environmental Management* 26: 491–502.
- Orfanidis S, Panayotidis P, Stamatis N. 2001. Ecological evaluation of transitional and coastal waters: a marine benthic macrophytes-based model. *Mediterranean Marine Science* 2: 45–46.
- Padisak J, Borics G, Feher G, Grigorszky I, Oldal I, Schmidt A, Zambone-Doma Z. 2003. Dominant species, functional assemblages and frequency of equilibrium phases in late summer phytoplankton assemblages in Hungarian small shallow lakes. *Hydrobiologia* **502**: 157–168.
- Paerl HW, Valdes LM, Pinckney JL, Piehler MF, Dyble J, Moisander PH. 2003. Phytoplankton photopigments as indicators of estuarine and coastal eutrophication. *Bioscience* **53**: 953–964.
- Panayotidis P, Montesanto B, Orfanidis S. 2004. Use of low-budget monitoring of macroalgae to implement the European Water Framework Directive. *Journal of Applied Phycology* **16**: 49–59.
- Pauly D, Christensen V. 1995. Primary production required to sustain global fisheries. Nature 374: 255–257.
- Pearson TH. 2001. Functional group ecology in soft-sediment marine benthos: the role of bioturbation. In Oceanography and Marine Biology 39: 233.
- Pearson TH, Rosenberg R. 1978. Macrobenthic succession in relation to organic enrichment and pollution of the marine environment. *Oceanography and Marine Biology Annual Review* 5: 229–311.
- Petchey OL, Gaston KJ. 2002. Functional diversity (FD), species richness and community composition. *Ecology Letters* 5: 402–411.
- Petchey OL, Hector A, Gaston KJ. 2004. How do different measures of functional diversity perform? *Ecology* **85**: 847–857.
- Pires AM, Cowx IG, Coelho MM. 2000. Benthic macroinvertebrate communities of intermittent streams in the middle reaches of the Guadiana Basin (Portugal). *Hydrobiologia* **435**: 167–175.
- Quinones RA. 1994. A comment on the use of allometry in the study of pelagic ecosystem. In *The Size Structure and Metabolism of the Pelagic Ecosystem*, Rodríguez J, Li WKW (eds). *Scientia Marina* 58: 11–16.
- Reizopoulou S, Nicolaidou A. 2004. Benthic diversity of coastal brackish-water lagoons in western Greece. *Aquatic Conservation: Marine and Freshwater Ecosystems* 14: S93–S102.
- Reizopoulou S, Nicolaidou A. 2005. Index of Size Distribution (ISD): a method for assessing disturbance in lagoons. In *International Conference on Lagoons and Coastal Wetlands in Global Change Context: Impacts and Management Issues*, UNESCO Proceedings Series, Venice, in press.
- Reizopoulou S, Thessalou-Legaki M, Nicolaidou A. 1996. Assessment of disturbance in Mediterranean lagoons: an evaluation of methods. *Marine Biology* **125**: 189–197.
- Reynolds CS, Huszar V, Kruk C, Naselli-Flores L, Melo S. 2002. Towards a functional classification of the freshwater phytoplankton. *Journal of Plankton Research* 24: 417–428.
- Robson BJ, Barmuta LA, Fairweather PG. 2005. Methodological and conceptual issues in the search for a relationship between animal body-size distributions and benthic habitat architecture. *Marine and Freshwater Research* **56**: 1–11.
- Rochet MJ, Trenkel VM. 2003. Which community indicators can measure the impact of fishing? A review and proposals. *Canadian Journal of Fisheries and Aquatic Sciences* **60**: 86–99.
- Rodríguez J, Jiménez F, Bautista B, Rodríguez V. 1987. Planktonic biomass spectra dynamics during a winter production pulse in Mediterranean coastal waters. *Journal of Plankton Research* 9: 1183–1194.
- Rodríguez J, Blanco JM, Jiménez-Gomez F, Echevarría F, Gil J, Rodríguez V, Ruiz J, Bautista B, Guerrero F. 1998. Patterns in the size structure of the phytoplankton community in the deep fluorescence maximum of the Alborean Sea (southwestern Mediterranean). *Deep-Sea Research* **45**: 1577–1593.
- Rodríguez J, Tintoré J, Allen JT, Blanco JM, Gomis D, Reul A, Ruiz J, Rodríguez V, Echevarría F, Jiménez-Gomez F. 2001. Mesoscale vertical motion and size structure of phytoplankton in the ocean. *Nature* **410**: 360–363.
- Rodríguez J, Jiménez-Gomez F, Blanco JM, Figueroa FL. 2002. Physical gradients and spatial variability of the size structure and composition of phytoplankton in the Gerlache Strait (Antarctica). *Deep-Sea Research II* 49: 693–706.
- Roscher C, Schumacher J, Baade J, Wilcke W, Gleixner G, Weisser WW, Schmid B, Schulze ED. 2004. The role of biodiversity for element cycling and trophic interactions: an experimental approach in a grassland community. *Basic and Applied Ecology* **5**: 107–121.
- Rutten TPA, Sandee B, Hofman RT. 2005. Phytoplankton monitoring by high performance flow cytometry: a successful approach. *Cytometry* **64A**: 16–26.

- Sabetta L, Fiocca A, Margheriti L, Vignes F, Basset A, Mangoni O, Carrada GC, Ruggieri N, Ianni C. 2005. Body size-abundance distributions of nano- and micro-phytoplankton guilds in coastal marine ecosystems. *Estuarine Coastal and Shelf Science* **63**: 645–663.
- Schramm W. 1999. Factors influencing seaweed responses to eutrophication: some results from EU-project EUMAC. *Journal of Applied Phycology* 11: 69–78.
- Seip KL, Reynolds CS. 1995. Phytoplankton functional attributes along trophic gradient and season. *Limnology and Oceanography* **40**: 589–597.
- Serra T, Granata T, Colomer J, Stips A, Møhlenberg F, Casamitjana X. 2003. The role of advection and turbulent mixing in the vertical distribution of phytoplankton. *Estuarine*, *Coastal and Shelf Science* **56**: 53–62.
- Shannon CE, Weaver W. 1949. *The Mathematical Theory of Communication*. University of Illinois Press: Urbana, IL. Shannon LJ, Cury PM. 2004. Indicators quantifying small pelagic fish interactions: application using a trophic model of the southern Benguela ecosystem. *Ecological Indicators* 3: 305–321.
- Sheppard CRC. 1998. Biodiversity patterns in Indian Ocean corals, and effects of taxonomic error in data. *Biodiversity and Conservation* 7: 847–868.
- Shin YJ, Cury P. 2004. Using an individual-based model of fish assemblages to study the response of size spectra to changes in fishing. *Canadian Journal of Fisheries and Aquatic Sciences* **61**: 414–431.
- Sibbing FA, Nagelkerke LAJ. 2001. Resource partitioning by Lake Tana barbs predicted from fish morphometrics and prey characteristics. *Reviews in Fish Biology and Fisheries* **10**: 393–437.
- Sieburth JMcN. 1979. Sea Microbes. Oxford University Press: New York.
- Simberloff D, Dayan T. 1991. The guild concept and the structure of ecological communities. *Annual Review of Ecology and Systematics* 22: 115–143.
- Simboura N, Zenetos A. 2002. Benthic indicators to use in Ecological Quality classification of Mediterranean soft bottom marine ecosystems, including a new Biotic Index. *Mediterranean Marine Science* 3: 77–111.
- Simpson EH. 1949. Measurement of species diversity. Nature 163: 688.
- Sosa Lopez A, Mouillot D, Do Chi T, Ramos Miranda J. 2005. Ecological indicators based on trophic rank distributions of fish biomass: an application to the Terminos coastal lagoon, Mexico. *ICES Journal of Marine Science* **62**: 453–458.
- Spatharis S, Panayotidis P, Montesanto B, Danielides D. 2003. Application of ecological indices on phytobenthos data for the implementation of the Water Frame Directive (WFD, 2000/60/EC). 2nd Mediterranean Symposium on Marine Vegetation.
- Sprules WG, Munawar M. 1986. Plankton size spectra in relation to ecosystem productivity, size and perturbation. *Canadian Journal of Fisheries and Aquatic Sciences* **43**: 1789–1794.
- Statzner B, Doledec S, Hugueny B. 2004. Biological trait composition of European stream invertebrate communities: assessing the effects of various trait filter types. *Ecography* 27: 470–488.
- Steneck RS, Dethier MN. 1994. A functional group approach to the structure of algal-dominated communities. *Oikos* **69**: 478–498.
- Tamigneaux E, Legendre L, Klein B, Mingelbeir M. 1999. Seasonal dynamics and potential fate of size-fractionated phytoplankton in a temperate near shore environment (western Gulf of St. Lawrence, Canada). *Estuarine Coastal and Shelf Science* **48**: 253–269.
- Vaquer A, Troussellier M, Courties C, Bibent B. 1996. Standing stock and dynamics of picophytoplankton in the Thau lagoon (northwest Mediterranean coast). *Limnology and Oceanography* **41**: 1821–1828.
- Waldbusser GG, Marinelli RL, Whitlatch RB, Visscher PT. 2004. The effects of infaunal biodiversity on biogeochemistry of coastal marine sediments. *Limnology and Oceanography* 49: 1482–1492.
- Walker B, Kinzig A, Langridge J. 1999. Plant attribute diversity, resilience, and ecosystem function: the nature and significance of dominant and minor species. *Ecosystems* 2: 95–113.
- Warwick RM. 1986. A new method for detecting pollution effects on marine macrobenthic communities. *Marine Biology* 92: 557–562.
- WFD. 2000. European Commission Directive 2000/60/EC of the European Parliament and of the Council of 23 October 2000 establishing a framework for Community action in the field of water policy. *Official Journal of the European Communities*, L 327/1-72, Brussels.
- Yew-Hoong Gin K, Lin X, Zhang S. 2000. Dynamics and size structure of phytoplankton in the coastal waters of Singapore. *Journal of Plankton Research* 22: 1465–1484.
- Zar JH. 1997. Biostatistical Analysis, 4th edn. Prentice-Hall International: Englewood Cliffs, NJ.