FEATURE ARTICLE: REVIEW

Framework for understanding marine ecosystem health


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ABSTRACT: Although the terms ‘health’ and ‘healthy’ are often applied to marine ecosystems and communicate information about holistic condition (e.g. as required by the Ecosystem Approach), their meaning is unclear. Ecosystems have been understood in various ways, from non-interacting populations of species to complex integrated systems. Health has been seen as a metaphor, an indicator that aggregates over system components, or a non-localized emergent system property. After a review, we define good ecosystem health as: ‘the condition of a system that is self-maintaining, vigorous, resilient to externally imposed pressures, and able to sustain services to humans. It contains healthy organisms and populations, and adequate functional diversity and functional response diversity. All expected trophic levels are present and well interconnected, and there is good spatial connectivity amongst subsystems.’ We equate this condition with good ecological or environmental status, e.g. as referred to by recent EU Directives. Resilience is central to health, but difficult to measure directly. Ecosystems under anthropogenic pressure are at risk of losing resilience, and thus of suffering regime shifts and loss of services. For monitoring whole ecosystems, we propose an approach based on ‘trajectories in ecosystem state space’, illustrated with time-series from the northwestern North Sea. Change is visualized as Euclidian distance from an arbitrary reference state. Variability about a trend in distance is used as a proxy for inverse resilience. We identify the need for institutional support for long time-series to underpin this approach, and for research to establish state space co-ordinates for systems in good health.

Changes in the northern North Sea, 1958–2008, plotted in a state space defined by the breeding success of kittiwakes, abundance of copepods Calanus spp., and simulated annual primary production.

Image: P. Tett,
Photos: R. Gowen (kittiwakes), D. Altin, BioTrix (Calanus spp.)

KEY WORDS: Ecosystem approach · Functional and response biodiversity · Resilience · State space · Regime shift · EU Marine strategy Framework Directive

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INTRODUCTION

Marine ecosystems provide services that lead to societal benefits (Atkins et al. 2011, Barbier et al. 2012). Three common strategies aim to protect these services against anthropogenic pressures. One conserves key organisms and habitats, one protects against disturbance, and one manages resource exploitation. These strategies, however, often neglect the interactions amongst ecosystem components, which in some cases ameliorate pressure effects. In other cases the effects multiply, leading to major disturbances to marine food webs and their use by humans (Fogarty & Murawski 1998, Jackson et al. 2001, Thurstan & Roberts 2010).

The need for integrated environmental management is recognized in the ‘Ecosystem Approach’, and the idea that ecosystems can have an optimal, or healthy, state is beginning to be used as a goal for holistic management, appearing in the marine environmental protection laws of several nations (Table S1 in Supplement 1; all supplements at www.int-res.com/articles supp/ m494p001_supp.pdf). However, it has been argued that this idea ‘is based on controversial, value-based assumptions that masquerade as science’ (Lackey 2001, p. 437). Crucial aspects of health have been presented in terms of metaphors such as those of stability landscapes (Holling 1973, Scheffer et al. 2001) rather than quantitative theories. Proposed holistic methodologies have focussed on the diagnosis of ecosystem pathologies (McLusky & Elliott 2004) or of undesirable disturbance (Tett et al. 2007), rather than the identification of healthy states or the good status required, for example, by recent EU Directives (Borja et al. 2010). Thus, there is a need to clarify what is meant by ‘good ecosystem health’ and to develop methods for assessing it.

The present article originated in workshops that had been tasked with developing a sound scientific basis for evaluating the overall status of UK marine waters as required by EU Directives. The article includes a selective review of the literature relevant to ecosystem health, a glossary of key terms (Box 1), and a proposed method for tracking change in the state of marine ecosystems by means of plots in state space. We exemplify the method with data from the North Sea, but believe it to be of general applicability.

REVIEW OF ECOSYSTEM HEALTH

Nature of ecosystems and aims of management

Building on Tansley (1935), Lindeman (1942) defined an ecosystem as a ‘system composed of physico-chemical-chemical-biological processes active within a space-time unit of any magnitude, i.e. the biotic community plus its abiotic environment’. This definition is widely accepted, but a continuing debate concerns the processes. Do they belong to the system or to its components? Is there such a thing as an ecosystem, or is the word merely a label for a human-delineated collection of species and habitats?

Clements (1916, 1936) viewed the climax of terrestrial vegetation and its associated animals as ‘a complex organism inseparably connected with its climate and often continental in extent’ (Clements 1936, p. 253) and possessing ontogeny and phylogeny. Odum (1969) considered that ecological succession culminated in a stabilized ecosystem that maximised biomass, information content, and symbiotic interactions amongst organisms, for a given energy flux. Moss (2008) argued that what he called ecosystems (but which might be better seen as biomes) have evolved, through the natural selection of component species, to become optimal systems for the use of natural resources. In contrast, Gleason (1926) held that terrestrial floras were no more than contingent associations of species that had been selected by the species’ environmental requirements and their abilities to disperse. Davis & Slobodkin (2004) denied that ecosystems existed as ‘some integrated entity … that grows, lives, reproduces and dies, or can be injured or healed’ (p. 1). However, Winterhalder et al. (2004) argued that ‘biotic communities and ecological systems … do show a very high degree of integration, or “coherency” in their responses to perturbations of various kinds’ (p. 5). If integration exists, a likely explanation is that the biota contribute to the ‘ecological theatre’ in which they act out the ‘evolutionary play’ (Hutchinson 1965, Post & Palkovacs 2009) and so co-evolve (Urban & Skelly 2006, Johnson & Stinchcombe 2007).

A related debate concerns stability in ecosystems (Holling 1973, Botkin 1990, Cuddington 2001, Gowen et al. 2012). Are natural systems in balance, and do they tend to return to that balance if perturbed? Or are they intrinsically, and perhaps unpredictably, variable? If pristine ecosystems tend towards a balance of organisms, the goal of environmental protection seems clear: to ensure that (commensurate with human aspirations) communities are assisted towards, or maintained at, their ecological climaxes. If, instead, ecosystems are ‘open, complex and dynamic systems that are characteristically transient and unstable’ (Spieles 2010, referring to Botkin 1990), as evidence increasingly suggests, then the goal of ecosystem management (as opposed to the
Box 1. Glossary of key terms relating to ecosystem health. The definitions are those used in this article, unless otherwise qualified. The terms are italicized here, on the first substantive use in the text, and sometimes thereafter

<table>
<thead>
<tr>
<th>Term</th>
<th>Definition</th>
</tr>
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<tbody>
<tr>
<td>Attractor</td>
<td>A point, or repeating trajectory, in state space to which a system tends</td>
</tr>
<tr>
<td>Autopoïetic</td>
<td>A self-making or self-maintaining system</td>
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<td>(Varela &amp; Maturana 1980)</td>
<td></td>
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<tr>
<td>Basin of attraction</td>
<td>A region in state space in which the system tends to remain (Walker et al. 2004, p. 3)</td>
</tr>
<tr>
<td>Biodiversity</td>
<td>The phenotypic variability amongst organisms within an ecosystem, and the genetic basis of that variability; cf. 'includes diversity within species, between species and of ecosystems' (Article 2, Convention on Biological Diversity 1992: www cbd int/convention/text/default.shtml)</td>
</tr>
<tr>
<td>Biome</td>
<td>A distinctive combination of plants and animals in a fully developed or climax community, characterized by a uniform life form of vegetation, development stages (Smith 1992), extended to refer to the combination expected under particular ecological conditions</td>
</tr>
<tr>
<td>Community</td>
<td>The biota in an ecosystem as distinct from the abiotic environment: i.e. the organisms or species and the trophic links amongst them</td>
</tr>
<tr>
<td>Compliance</td>
<td>The ratio of ecosystem state change to external pressure change (Fig. 3); the inverse of resistance (to pressure) in the elasticity analogue for resilience (Supplement 6)</td>
</tr>
<tr>
<td>Complex</td>
<td>A system with sufficient components and interactions to exhibit emergent properties or behaviour, such as non-linearity, homeostasis or autopoïesis, often as a result of nested sub-systems (hierarchy) and feedback loops</td>
</tr>
<tr>
<td>Connectance</td>
<td>The fraction of all possible links that are realized in a [trophic] network (Dunne et al. 2002)</td>
</tr>
<tr>
<td>Connectivity</td>
<td>Refers to spatial links allowing exchanges of individuals or genetic materials amongst meta-populations (Steneck &amp; Wilson 2010) and of materials and energy amongst habitats or sub-regions within an ecosystem (Dakos et al. 2010)</td>
</tr>
<tr>
<td>CPR</td>
<td>Continuous Plankton Recorder, towed by ships of opportunity, to sample plankton (Richardson et al. 2006)</td>
</tr>
<tr>
<td>Domain</td>
<td>A defined region in an ecosystem</td>
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<tr>
<td>DPSIR</td>
<td>Acronym for Driver (within society)-Pressure-State-Impact-Response (within society) paradigm, originally by Luiten (1999), updated by Atkins et al. (2011)</td>
</tr>
<tr>
<td>Ecobhydrodynamic</td>
<td>The physical conditions that select for species and communities in the sea, including water depth, stirring and stratifying tendencies, light penetration, and sediment type (Tett et al. 2007)</td>
</tr>
<tr>
<td>Ecological quality ratio</td>
<td>The ratio (between 0 and 1) of the value of an ecological indicator to the value under good conditions</td>
</tr>
<tr>
<td>Ecological status</td>
<td>An expression of the quality of the structure and functioning of aquatic ecosystems associated with surface waters’ (WFD article 2.21 &amp; Annex V)</td>
</tr>
<tr>
<td>Ecosystem</td>
<td>The system composed of the physical-chemical-biological processes active within a space-time unit of any magnitude, i.e. the biotic community plus its abiotic environment’ (Lindeman 1942, p. 400)</td>
</tr>
<tr>
<td>Emergent property</td>
<td>(System) behaviors that cannot be identified through functional decomposition (Johnson 2006) and that are more than the sum of the system’s parts, even if explicable in terms of within-system processes (O’Connor &amp; Wong 2009)</td>
</tr>
<tr>
<td>Empirical</td>
<td>Conceptualization of ecosystem health: a view of ecosystems as the sum of their parts and hence able to be simulated by mechanistic models; cf. systemic</td>
</tr>
<tr>
<td>Empirical model</td>
<td>A model that, even if theory-based, has its parameter values adjusted to best fit simulations to observations; cf. mechanistic model</td>
</tr>
<tr>
<td>Endogenous</td>
<td>Pressure generated within the local social-ecological system and hence susceptible to local management; in contrast, exogenous pressures are externally generated, and local management can only deal with the consequences (Elliott 2011)</td>
</tr>
<tr>
<td>Environment</td>
<td>(1) The abiotic component of an ecosystem (cf. community); (2) What lies outside a given system</td>
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<tr>
<td>ERSEM</td>
<td>European Regional Seas Ecosystem Model, a mechanistic model for marine ecosystems including pelagic and benthic components (Baretta et al. 1995, Baretta-Bekker &amp; Baretta 1997)</td>
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<tr>
<td>Euclidian distance</td>
<td>The scalar distance between 2 points in state space, calculated from the square root of the sum of squares of the distance along each axis</td>
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<td>External description</td>
<td>Of a system: properties of the system that characterize its holistic behavior when seen from outside</td>
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<tr>
<td>Functional-group diversity</td>
<td>The sets of species (or components of biodiversity) responsible for ecosystem functions; the sets correspond to benthic guilds, or pelagic life forms</td>
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<tr>
<td>Functional-response diversity</td>
<td>Biodiversity contributing to different responses to environmental change within a functional group</td>
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<td>GES</td>
<td>Good Environmental Status as defined in Article 3.5 of the MSFD, which includes the requirement that 'the structure, functions and processes of the constituent marine ecosystems...allow those ecosystems to function fully and to maintain their resilience to human-induced environmental change'</td>
</tr>
<tr>
<td>GETM</td>
<td>General Estuarine Transport Model (www getm.eu), a hydrodynamic model designed for use in shelf seas including those with significant intertidal areas</td>
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<tr>
<td>Granularity</td>
<td>The existence, nature and scale of spatial patchiness within an ecosystem</td>
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<tr>
<td>Hierarchy</td>
<td>Arrangements whereby systems contain subsystems; the emergent properties of the latter contribute to the functioning of the main system, which provides boundary conditions for the subsystems; see also panarchy</td>
</tr>
<tr>
<td>Impact</td>
<td>The consequences of ecosystem state change on services to human societies</td>
</tr>
<tr>
<td>Integrity</td>
<td>Of a system with open boundaries: what maintains the distinctiveness between a system and what is outside it</td>
</tr>
<tr>
<td>Internal description</td>
<td>Of a system: description in terms of system components and connections, e.g. in terms of a set of state variables</td>
</tr>
<tr>
<td>k-adapted (species)</td>
<td>Slow in growth but efficient at resource acquisition; cf. r-adapted</td>
</tr>
</tbody>
</table>

(Box continues on next page)
Box 1 (continued)

| **Latitude** | the greatest amount that a system can change before losing its ability to regain or remain within its original regime (Walker et al. 2004) |
| **Lifeform** | a set of species (not necessarily taxonomically related) that play similar roles in ecosystem function; e.g. ‘algal silicon users’ such as diatoms and silicoflagellates |
| **Mechanistic model** | equations and parameters assembled from hypotheses validated in controlled experiments; cf. **empirical model** |
| **MVA (MultiVariate Analysis)** | statistical methods for analysing relationships amongst multiple variables; including **PCA** |
| **Net primary production** | formation of organic material by photosynthesis, net of respiratory losses by the same organisms over the time-period considered; in the case of the model-simulated production reported here, photosynthesis and respiration have been integrated over the water column and the day-night cycle |
| **Open** | of a system whose boundaries allow inputs and/or outputs |
| **Organization** | the types and arrangements or interconnections of the components of a system; cf. **vigour** |
| **Panarchy** | a view of dynamic systems (including ecosystems) as made up of nested or linked subsystems cycling adaptively through development and collapse (Holling 2004) |
| **PCA: Principal Component Analysis** | which converts a set of observations of possibly correlated variables into a smaller set of values of linearly uncorrelated variables called ‘principal components’ |
| **Precariousness** | the closeness of a system to: the state at which resilience collapses and a regime shift occurs (Walker et al. 2004); the edge of the ‘basin of attraction’ (Holling 1973); the edge of the ‘cliff’ or the ‘elastic limit’ |
| **Pressure** | (1) a link in the **DPSIR** chain (Luijten 1999), referring to external (anthropogenic) pressure on an ecosystem; (2) the human-altered influxes, outflows and disturbances acting on an ecosystem; in either case, dimensionally undefined |
| **Principal axis (or component)** | a line drawn through a set of points in a multivariable space so as to minimize the scatter of points about it (typically by minimizing the sum-of-squares of deviations); see **PCA** |
| **Production** | the formation of new organic matter at a given trophic level |
| **Qualitative descriptor** | 1 of 11 components listed in Annex I of the **MSFD** as determining the characteristics of **GES** |
| **r-adapted (species)** | populations capable of rapid increase but inefficient at resource acquisition and liable to high predation; cf. **k-adapted** |
| **Recovery** | return towards undisturbed system state as pressure is relaxed; as a component of resilience, the capability of a system to recover |
| **Regime** | a bundle of trajectories in system **state space** |
| **Regime shift** | a substantial and persistent change in ecosystem state, condition, or regime that involves many ecosystem components, impacts substantially on services, and in systems theory is explained by a shift to a new attractor |
| **Resilience** | ‘the capacity of a system to absorb disturbance and reorganize while undergoing change so as to maintain essentially the same functions, structure, identity and feedbacks’ (Folke et al. 2004) |
| **Resistance** | one of the components of resilience—a measure of difficulty in moving a system within a **basin of attraction** (Walker et al. 2004); ‘the ability of an ecosystem to resist displacement from its reference state during a perturbation stress’ (Vallina & Le Quéré 2011) |
| **Scalar variability** | describes the variability of system state about a long-term trend when both are expressed as **Euclidian distance** using self-standardized variables |
| **SEM (Structural Equation Modelling)** | a statistical method aimed at eliciting a group of factors, and connections or relations, that best fits a data set (Hox & Bechger 1998) |
| **SMP: (UK) Sea-bird monitoring programme** (jncc.defra.gov.uk/page-1550) |
| **Social-ecological system** | a linked system of people and nature (Berkes & Folke 1998); a spatially-bounded region containing an ecosystem and a social system interacting with each other (Tett et al. 2013) |
| **Stability** | the tendency for system state to remain near an attractor in **state space** |
| **Stability landscape** | metaphor in which the state of an ecosystem is represented by the position of a ball in an undulating landscape |
| **State** | (1) of a system a single set of values of a set of **state variables** sufficient to specify the system’s condition uniquely, and plotting to a point in the corresponding **state space**; (2) ‘the state of the environment (external to human society) as affected by pressure’ (Luijten 1999) |
| **State space** | ‘the n-dimensional space of possible locations of [state] variables’ (von Bertalanffy 1972, p. 417) |
| **State variable** | a quantification of a system property |
| **Status** | the condition of (all or part of) an ecosystem assessed relative to a norm |
| **System** | ‘a set of elements standing in interrelation among themselves and with [their] environment’ (von Bertalanffy 1972) |
| **Systemic (conceptualization of ecosystem health)** | based on a view of ecosystems in accordance with General Systems Theory (von Bertalanffy 1972) and allowing for emergent properties such as resilience; cf. **empirical model** |
| **Trajectory** | a sequence of system states plotted in **state space** |
| **Type-specific reference conditions**: in the **WFD** (Annexes II & V), the conditions ‘normally associated with that [water body] type under undisturbed conditions’ |
| **Variability** | in **state space** has 3 components: semi-cyclical (e.g. associated with seasonal cycles and classed as part of organization); medium-term (about a trend); and long-term (i.e. a trend) (Fig. 1) |
| **Vigour** | the ability of a system to maintain or renew its organization by drawing on production (Costanza 1992, vigour) |
more restricted aims of species and habitat conserva-
tion) is harder to define.

A further complication arises if sustainability is
seen as a property, or goal, of ‘social-ecological sys-
tems’ (Berkes & Folke 1998) which have psychologi-
cal and social dimensions as well as physical exist-
ence (Tett et al. 2013). Although both ecological
knowledge and preferences for certain states of eco-
systems lie in the mental and social worlds, it is
important to distinguish social values from ecological
facts and theories because they relate to different
sorts of ‘validity claims’ (Habermas 1984). This does
not mean that ecological knowledge should only be
used instrumentally, to guide actions decided by
political processes (Lackey 2001, Davis & Slobodkin
2004). Our view is that science must inform debate as
well as help implement its outcome. Exploring the
meaning of ‘the health’ of ecosystems in terms of
their structure and function can show why aiming at
‘good ecosystem health’ might be good for societies
using services from these ecosystems (Atkins et al.
2011).

**Ecosystem health as a metaphor**

Metaphors are important in human speech and
thought (Pinker 2007). They can help to explain com-
plex things, such as ecosystems, that cannot be ade-
quately described in terms drawn from direct human
experience. But they can be ambiguous, misleading
when the metaphor is confused with reality, and dan-
gerous when used to guide action.

Human health is an attractive metaphor for ecosys-
tem condition because there is a universal under-
standing of what it means to be well: to be free from
illness, to function well, to be vigorous, to resist and
recover from disease, to maintain physical and men-
tal integrity in the face of stress. That is to say, to be
in a socially and individually desired condition,
defined by a certain range of physiological and psy-
chological states, and to be able to maintain that con-
dition or to recover it after disturbance.

The metaphor has aided ecologists in conceptualizing
ecosystem functioning as well as explaining ecosystem
condition to the general public and suggesting regula-
tory goals to government. But it may be dangerous
(Lackey 2001, Davis & Slobodkin 2004) if it allows per-
sonal or sectoral values to be intruded into what are
claimed as objective assessments. Thus there is a need
to consider whether, and to what extent, the compo-
nents of human health map to those of ecosystem
health.

**Ecosystem health as an aggregate property**

One approach to a definition is to see health not as
a single property of an ecosystem, but as an aggreg-
gate of contributions from organisms, species and
processes within a defined area. We will refer to this
approach as *empirical*, because, although not with-
out theoretical content, it is based largely on expert
observations of ecosystems.

Elliott (2011) listed 6 levels of biological organiza-
tion, each of which could be termed healthy or un-
healthy: cell, tissue, organism, population, *community*
and ecosystem. At the level of organisms, the
meaning of ‘health’ seems unambiguous and identical
with that of human physical well-being. At the next
level, health concerns the viability of populations or
species. At the level of communities and ecosystems
the argument becomes more complex. Elliott (2011)
described ‘community health’ as that of an assemblage
of organisms that can continue to function in terms of
inter-species relationships; and ‘ecosystem health’ as
providing protection against the ‘ecosystem patholo-
(Table S2 in Supplement 2). Monitoring at this level
allows ‘detection of things going wrong’ against a
background of system variability (Elliott 2011). Earlier,
Odum (1985) had listed trends expected in ‘stressed
ecosystems’ (Table S3 in Supplement 2), basing these
on a conceptual model of ecosystem succession under
undisturbed conditions (Odum 1969). We have drawn
on these 2 sets of ill-health diagnostics as the basis for
the empirical and aggregatable criteria for marine
ecosystem health in Table 1.

The first column in Table 1 provides generic criteria.
However, there are several types of marine ecosys-
tems, each with their characteristic *lifeforms* of pri-
mary producers, and each linked to particular *ecohy-
drodynamic* (Tett et al. 2007) and climatic conditions.
Criteria that are applicable across all these types
might provide little guidance in managing pressures
on a particular type. For example, biodiversity is natu-
raly low in the physically stressed environment of
estuaries (Elliott & Quintino 2007), where biomass can
be high as a result of inputs of allochthonous organic
matter (Elliott & Whitfield 2011). In contrast, the oligo-
trophic waters of the eastern Mediterranean support a
high diversity of pelagic micro-algae (Ignatiades et al.
2009). Thus, there is need for the ecological norms of
column 2, which link the general criteria to what
might be expected in a particular ecosystem under
undisturbed conditions.

The European Union Water Framework Directive
(*WFD*) is based on such a norm-based approach
Table 1. Components of (good) ecosystem health, according to the empirical approach, and as interpreted by EU Directives. Column 3 gives corresponding specifications from Annex V of the EU Water Framework Directive (WFD) for ‘high quality status’ (which we equate with good health) in ‘transitional’ and ‘coastal’ waters. Column 4 refers to the relevant ‘qualitative descriptors’ for determining good environmental status’ in Annex I of the Marine Strategy Framework Directive (MSFD), which are expanded by COM (2010).

<table>
<thead>
<tr>
<th>1. Generic component of ecosystem health</th>
<th>2. Ecological norm</th>
<th>3. WFD ‘high quality status’</th>
<th>4. MSFD ‘qualitative descriptors’</th>
</tr>
</thead>
<tbody>
<tr>
<td>Autochthonous primary photosynthetic production plus import of organic matter is roughly in balance with consumption, so that there is no large excess of respiration that might lead to deoxygenation nor substantial export of unconserved material</td>
<td>Life-form of primary producer is typical of ecohydrodynamic type and production is within characteristic range for undisturbed example of this type</td>
<td>Phytoplankton biomass to be ‘consistent with the type-specific physico-chemical condition’, macro-algal cover, and angiosperm abundance to be ‘consistent with undisturbed conditions’. [O]xygen balance ... remain[s] within the range ... normally associated with undisturbed conditions.’</td>
<td>'(5) Human-induced eutrophication is minimised, especially adverse effects thereof, such as losses in biodiversity, ecosystem degradation, harmful algae blooms and oxygen deficiency in bottom waters.'</td>
</tr>
<tr>
<td>Nutrient supply, cycling rates and elemental ratios are adequate to support community functioning and structure; communities make efficient use of these resources</td>
<td>Nutrient seasonal cycles, amounts, and elemental ratios are similar to those under undisturbed conditions</td>
<td>‘Nutrient concentrations remain within the range normally associated with undisturbed conditions.’</td>
<td>Not explicitly mentioned</td>
</tr>
<tr>
<td>Sufficient biodiversity to fulfill all the necessary bio-geochemical roles, to support species at higher trophic levels, and to provide a reserve in case of loss of species; keystone species flourishing where essential for community functioning, there is a mixture of r- and k-adapted species, and a mixture of reproductive and young individuals within populations</td>
<td>All aspects of diversity are appropriate for undisturbed example of ecosystem type as determined by climate and local (eco) hydrodynamic conditions</td>
<td>‘The composition and abundance of phytoplanktonic taxa are consistent with undisturbed conditions.’ ‘All disturbance-sensitive macroalgal and angiosperm taxa associated with undisturbed conditions are present.’ ‘The level of diversity and abundance of invertebrate taxa is within the range normally associated with undisturbed conditions.’</td>
<td>'(1) Biological diversity is maintained. The quality and occurrence of habitats and the distribution and abundance of species are in line with prevailing physiographic, geographic and climatic conditions. (2) Non-indigenous species introduced by human activities are at levels that do not adversely alter the ecosystems.'</td>
</tr>
<tr>
<td>Community structure includes multiple trophic levels and a variety of trophic links between levels; in cases where autogenic or responsive successions are important, then either a substantial proportion of the ecosystem is in the mature state, or there are no impediments to reaching such a state</td>
<td>Structure is that characteristic of this ecosystem type under undisturbed conditions</td>
<td>Not explicitly dealt with by this Directive</td>
<td>'(4) All elements of the marine food webs, to the extent that they are known, occur at normal abundance and diversity and levels capable of ensuring the long-term abundance of the species and the retention of their full reproductive capacity.'</td>
</tr>
<tr>
<td>Individual organisms are healthy and reproductively fit, not showing widespread pathologies, nor substantially contaminated with pollutants, nor exhibiting reduced resistance to disease or stress or reduced ability to detoxify</td>
<td>Body burden of contaminants below defined threshold, no substantial differences in performance compared with individuals at unpolluted station</td>
<td>Pollutant concentrations ‘remain within the range normally associated with undisturbed conditions’.</td>
<td>'(8) Concentrations of contaminants are at levels not giving rise to pollution effects.'</td>
</tr>
</tbody>
</table>

to assessing ecological status. Values for physico-chemical and biological quality elements are compared with those ‘normally associated with that [water body] type under undisturbed conditions’, i.e. with those of the type-specific reference conditions.

The third column in Table 1 quotes from the Directive’s specifications for high quality status, which is that of the reference condition, and which it seems logical to equate with good ecosystem health. However, finding current examples of the reference conditions in European coastal seas and estuaries has proven difficult (Hering et al. 2010, Borja et al. 2012).

Column 4 maps the criteria to the ‘qualitative descriptors’ of ‘good environmental status’ (GES) in the European Marine Strategy Framework Directive (MSFD), which takes a functional approach to good condition, in contrast (Borja et al. 2012) to the WFD’s norm-referencing.

Finally, there is another problem to be solved before using aggregation as an integrative method. It
concerns how to combine indicators whilst (1) avoiding double-counting of primary variables and (2) recognising each indicator’s importance in relation to ecosystem function. The problem is especially marked when aggregating over levels. Do community and ecosystem effects arise linearly from aggregation of organism-level and population-level effects, or can species replacement sustain ecosystem function even if some populations are damaged (for example, by pollution)? One combinatorial strategy has been to use weightings: for example, Aubry & Elliott (2006) multiplied indicators for estuarine disturbance by weights based on expert judgements, and Borja et al. (2011) used weights based on relative importance and indicator reliability to combine ecological quality ratios, for the MSFD’s qualitative descriptors of good condition, into a single integrated assessment. Another strategy is the precautionary ‘one-out, all-out’ principle used by the WFD (Borja & Rodríguez 2010), where a water body is demoted from ‘good’ status if a single quality element is scored below ‘good’. A general objection to all these procedures is that their outcomes may depend as much on decisions made about the combinatorial rules as on the actual condition of the ecosystem (Borja & Rodríguez 2010, Caroni et al. 2013).

Health as an emergent property of complex systems

An alternative to the empirical approach is what we call systemic, because its perspective is that of health as an emergent property of the whole ecological system rather than of any of its components. According to General Systems Theory (von Bertalanffy 1968, 1972), a system is ‘a set of elements standing in interrelation among themselves and with [their] environment’. The theory’s principles include a subset that apply to the open, complex, hierarchical and autopoietic systems that are exemplified by organisms and ecosystems. Living systems must be open (to their external environment) because they need a throughput of matter and energy to keep their components in a thermodynamically unlikely state (Schrödinger 1944, Boulding 1956, Lovelock & Margulis 1976, Moss 2008). They can persist only if they can maintain their internal organization and functions against these fluxes, or are able to restore internal states after perturbation. Health, in the systemic view, is this ability: a healthy system is one that maintains its integrity and is resilient under pressure. Thus, the concept of ecosystem health is more than a metaphor carried over from human health, and it is more than the sum of properties of components. It refers to patterns of system behaviour that are common to both organisms and ecosystems, and ill-health is recognized by a breakdown of this pattern.

Systems can be characterized in 2 ways (von Bertalanffy 1968, 1972). Internal description uses state variables, and can be exemplified for biological systems by the Lotka-Volterra equations (Slobodkin 1962, Hastings 1996). In a simple case, the state variables might be the population sizes of a predator and its prey. The behaviour of even small and species-poor ecosystems is more complex than such binary systems, and many variables may be needed to represent their internal state. Nevertheless, whether a system is simple or complex, change can be expressed ‘geometrically ... by the trajectories that the state variables traverse in the state space, that is, the n-dimensional space of possible location of these variables’ (von Bertalanffy 1972, p. 417). Fig. 1 de-

![Diagram of system state variable space](image-url)

Fig. 1. Definitions relating to system state variable space, exemplified for 2 axes, but generalizable to any number of dimensions. A state is represented by a point in state space; a trajectory is a (temporal) sequence of states; a regime is a coherent bundle of trajectories, such as those arising from seasonal cycles; a domain is a region in state space. The Euclidian distance gives the (shortest) scalar distance between 2 points in a state space of any dimensionality. The inset box shows the 3 types of variability discussed in this paper.
fines the terms used in discussing such state space diagrams.

In the case of external description, the behaviour of the system is described in terms of interactions with what is outside the system, often by specifying the relationship between inputs to the system and the resulting outputs. Fig. 2 combines a systemic view of an ecosystem with the Driver-Pressure-State-Impact-Response (DPSIR) paradigm of Luiten (1999) as updated by Atkins et al. (2011). Anthropogenic disturbance to trans-boundary fluxes constitutes a pressure, resulting in changes in the (internal) state of the system with consequent impacts on society. Resilience is the system property that determines the response of state to a pressure change; it is an emergent property because it cannot be localized in any particular component of the system.

Following a review of system-based definitions of health (Table S4 in Supplement 3), Costanza (1992) proposed that ecosystem health was best seen as a comprehensive, multiscale, dynamic, hierarchical measure of system resilience, organization, and vigour, [concepts that] are embodied in the term ‘sustainability’ which implies the system’s ability to maintain its structure (organization) and function (vigour) over time in the face of external stress (resilience).

All 3 components are necessary (Mageau et al. 1995): (1) a system lacking in vigour would tend towards abiotic thermodynamic equilibrium; (2) systems with excess vigour but ‘little or no organization, such as nutrient enriched lakes, …, or early successional ecosystems dominated … by ‘r’ selected species’ (p. 204) tend towards excessive blooms; and, (3) ‘certain highly managed systems, such as agriculture, aquaculture, and plantations’ (p. 204), lack resilience and require continuous human intervention for their maintenance. Costanza & Mageau (1999) thought that organization might be quantified through the analysis of trophic networks (e.g. Ulano - wicz & Kay 1991, Christensen & Pauly 1992) and vigour quantified by measurements of ‘ecosystem metabolism’, including its primary production.

Resilience is presently seen as the key component of system health. Holling (1973) defined it as ‘a measure of the ability of [eco]systems to absorb changes of state variables, driving variables, and parameters’ (p. 17) without ceasing to exist. Folke et al. (2004) saw resilience as ‘the capacity of a system to absorb disturbance and reorganize while undergoing change so as to maintain essentially the same functions, structure, identity and feedbacks’ (p. 558). Loss of resilience is now seen as leading to regime shift: a
change to a new internal organization, rather than extinction. Holling (1973) visualized continued existence as requiring trajectories in state space to remain within a *basin of attraction*, and distinguished resilience from ‘stability’—‘the ability of a system to return to equilibrium after a temporary disturbance’ (p. 17). Much subsequent literature, however, has equated ‘resilience’ with Holling’s stability and the ability to recover after perturbation. This included Elliott et al. (2007) and Tett et al. (2007). However, it now seems best to see resilience as having several components, one of which is recovery to ‘equilibrium’, bearing in mind that the restored equilibrium may be dynamic or correspond to a complex attractor in state space (Gowen et al. 2012). Other components of resilience (Walker et al. 2004) are resistance to pressure, and latitude: the greatest amount that a system can change before losing its ability to regain (or remain in) its original regime, and corresponding to the size of Holling’s ‘basin of attraction’.

Like Scheffer et al. (2001), Folke et al. (2004) argued that anthropogenic disturbance to ecosystems reduces resilience and increases the chances of regime shift. This leads to the metaphor of the ‘cliff’ in pressure-state diagrams (Fig. 3) (Elliott et al. 2007, Tett et al. 2007, van de Koppel et al. 2008). The paradigm in such diagrams is that natural ecosystems are resilient, and so resistant to anthropogenic pressures. Beyond a certain level of pressure, however, there is a danger of ecosystem collapse, from which it might be difficult to recover speedily (or at all) to the original conditions. An alternative metaphor (Fig. 4) is that of a *stability landscape*, in which a ball (representing system state) rolls to the lowest point in the valley, but can be displaced into other valleys either by change in the landscape or by increased movement of the ball (Scheffer et al. 2001, Walker et al. 2004). Duarte et al. (2009) pointed to the problem of ‘shifting baselines’ encountered when attempting to return an ecosystem to a prior state, such as that existing before eutrophication. They illustrated the problem with plots of state against pressure (corresponding to Fig. 3), but it could also be seen in terms of the changes in the topography of the landscape in Fig. 4. Finally, Scheffer et al. (2009) suggested that systems (such as ecosystems, but also economies) showed an increase in variability as they approached ‘critical transitions’ between regimes.

The idea of *panarchy* (Gunderson & Holling 2002, Holling 2004) is that on any particular spatial scale, systems naturally go through periods of collapse and recovery. During recovery, a system can adapt to changed circumstances, because post-collapse conditions can select for different species (in an ecosystem) or different institutions (in a society). The concept of panarchy also includes the interactions of

![Fig. 3. The cliff metaphor for change in ecosystem state (modified from Elliott et al. 2007 and Tett et al. 2007) used to illustrate resilience components (Walker et al. 2004) in terms of the effects of external pressure (P) on ecosystem condition or state (S). Latitude is shown as analogous to the ‘elastic limit’ of a mechanical system. Beyond this limit, the system deformation no longer changes linearly with pressure. Compliance is the ratio of state change to pressure change and thus is the inverse of resistance.](image1)

![Fig. 4. The landscape metaphor for stability and regime shift (Holling 1973). The ball, representing ecosystem state, moves between valleys denoting different regimes, following disturbance to the ball (Krebs 1988) or changes in the landscape (Scheffer et al. 2001). In the metaphor, stabilizing effects are likened to gravity. In mathematical terms, stability is the result of the system’s tendency to move towards an attractor state, shown at the bottom of each basin.](image2)
growth-collapse-recovery cycles across spatial scales or up and down hierarchies within the main system. Thus, heterogeneity within an ecosystem may contribute to vigour and resilience.

Although widely applied, a systemic approach to ecosystem health is currently less a refutable theory and more a point of view. Its cliff and landscape metaphors provide explanations that are useful in the public domain. The challenge, however, is to find ways to quantify ecosystem organization and vigour, and to understand how these quantifications relate to ecosystem resilience so that useful predictions of future ecosystem condition can be made from observations of present state and pressures.

Organisation and biodiversity

Costanza & Mageau (1999) explained 'high organization' as 'an efficient diversity of components and exchange pathways', and proposed its quantification through network analysis (Ulanowicz 1979, 2009). Despite some applications of the network approach (Christensen 1995, Christian et al. 2010), the difficulty in obtaining adequate data about exchanges has meant that most attempts to understand ecosystem function in terms of structure have continued to focus on the diversity of components rather than on the diversity of pathways in trophic or biogeochemical networks.

Biodiversity has a range of meanings. The 1992 international Convention for Biological Diversity (CBD) defined it as 'the variability among living organisms from all sources ... within species, between species, and of ecosystems' (Millennium Ecosystem Assessment 2005). Here, we define biodiversity as the phenotypic variability amongst organisms within an ecosystem, and the genetic basis of that variability. In most eukaryotes, that genetic basis is largely organized into reproductively isolated species, and it is at the level of species that biological diversity has mainly been studied. This focus derives from the standard paradigm of evolutionary ecology, which is that a species is a self-contained gene pool competing with other gene pools for territory in niche hyperspace, and thus fitness to survive in the physical world (Hutchinson 1957, 1965).

This paradigm has led some to propose equilibrium models of species-abundance distributions based on niche apportionment theory (MacArthur 1957, Tokesi 1993). However, others view species diversity as a non-equilibrium phenomenon: the result of high physical disturbance, or chaotic internal interactions, allowing multiple species to occupy 1 realized niche, and thus explaining (Scheffer et al. 2003) ‘the paradox of the plankton’ in aquatic ecosystems (Hutchinson 1961). Is high species diversity, then, more a symptom of instability than a cause of stability? Or can a diversity of non-equilibrium states contribute to ecosystem resilience under fluctuating conditions?

Many (perhaps too many; Green & Chapman 2011) indices of species diversity have been proposed (Washington 1984, Magurran 2004, Gray & Elliott 2009). The simplest are (1) empirical, such as the number of species scaled to the number of individuals in a sample. Others derive from: (2) models for species-abundance relationships (McGill et al. 2007); (3) the information content of a community, or a representative sample, containing a set of taxa of quantifiable abundance; or (4) the probabilities of different sorts of inter-organism encounter (Hurlbert 1971). Type (3), exemplified by the index of Shannon (1948), applies to any set of objects, and essentially treats variety in a system as if it were a message containing information about the system. Hurlbert (1971) argued in favour of (4) on the grounds that, whereas the message might be meaningful to ecologists, what was important to organisms was whether the next encounter would be with a mate, food item, or predator.

Contrary to Hurlbert’s argument, a diversity statistic based on species abundances or information theory can aggregate much of the fine grain of organism-level interaction, and might be deemed useful if it could be shown to correlate with, for example, resilience. Disappointingly, most meta-studies fail to find relationships between standard species diversity measures and ecosystem functions that are consistent over a variety of ecosystems (e.g. Hooper et al. 2005). Ives & Carpenter (2007) concluded that ‘diversity-stability relationships’ were complex, and that anthropogenic changes often affect stability and diversity simultaneously. Hooper et al. (2005) (Table S5 in Supplement 3) cited mostly positive effects of biodiversity on terrestrial ecosystem services, on ‘production and nutrient retention’ (which we equate with vigour), and on stability. They therefore argued that the diversity of functional traits was more important than the diversity of species. Likewise, Bengtsson (1998) argued that, for the management and development of sustainable ecosystems, it is probably more important to understand the linkages between key species or functional groups and ecosystem function, rather than focusing on species diversity.

Folke et al. (2004) distinguished functional-group diversity and functional-response diversity. The former refers to diversity between sets of species (not
necessarily taxonomically related) that carry out similar roles within an ecosystem, and hence quantifies the variety of ecosystem processes. Loss of a functional group results in a major change in ecosystem functioning. Functional-response diversity refers to variety in the ‘response to environmental change among species that contribute to the same ecosystem function’ (i.e. within a functional group; Folke et al. 2004, p. 570), and provides ‘a degree of ecological insurance against ecological uncertainty’ (Hughes et al. 2005, p. 383). However, if all species within a functional group respond similarly to pressures, then higher biodiversity will not afford additional protection. Estuaries exemplify naturally species-poor systems that are fully functioning and resilient (Elliott & Whitfield 2011). Osmotic stress excludes many species, but the few that can flourish here are those that are tolerant of disturbance, and it appears that they can supply all necessary ecosystem functions. Adaption to pressure might take place within species’ populations rather than by changes in the species assemblage as might occur in more species-diverse systems.

We conclude that functional-group diversity is the key component of ecosystem structure, that it is qualitative as much as quantitative, and that it can be thought of as a set of dimensions (i.e. axes in state space) relevant to a particular biome. Supplement 4 exemplifies this for soft-bottom communities in coastal waters.

**Ecosystem change and regime shift**

Marine ecosystem services provide benefits to human communities, valued at about 20 trillion US$ per year in 1994 (Costanza et al. 1997). A powerful argument for understanding, evaluating and managing marine ecosystem health is the link from health and resilience to ecosystem function and services. Ecosystems and their services change naturally, but the rate of change seems to have increased as a result of human activity in the ‘Anthropocene’ (Crutzen & Stoermer 2000, Crossland et al. 2005). Some alterations within ecosystems impact directly on services, but the cliff metaphor (Fig. 3) suggests the additional risk that changes can compound—leading to a ‘tipping point’ beyond which there is a partial collapse of existing system organization and a change to a new configuration. The new condition may provide fewer, or different, services; and the transition may prove difficult for social groups or economies that depend on particular services.

Such transformations of ecosystems are increasingly referred to as regime shifts, defined as ‘relatively sudden changes between contrasting, persistent states of a system’ (deYoung et al. 2008) or ‘sudden changes in ecosystem structure that can be detected across several ecosystem components’ (Spencer et al. 2011). They are said (Scheffer et al. 2001) to have been observed in major terrestrial and marine ecosystems. The latter include: coral reefs (Nyström et al. 2000); the north Pacific Ocean (Hare & Mantua 2000); Ringkøbing Fjord (Petersen et al. 2008); and the North Sea (Reid et al. 2001, Beauchard 2004, Weijerman et al. 2005, Spencer et al. 2011). In the northwestern Atlantic Ocean, from the Gulf of Maine to the Grand Banks of Newfoundland, there have been major changes in fisheries and ecosystems (Rose 2003, Buchsbaum et al. 2005, Bundy & Fanning 2005), with consequences that include reduced diversity of harvest and a simplified food web structure. These ‘may increase risks of ecological and economic disruptions’ (Steneck et al. 2011).

What are claimed as steps might be artefacts of the numerical methods used in analysis (Spencer et al. 2011). Nevertheless, large changes do occur, and what seems crucial from a human perspective is that the ecosystem has reached a new condition, qualitatively different from that in which it was originally found; that the new state is persistent; and that it provides different services.

As Scheffer et al. (2001) point out, ‘the notion that ecosystems may switch abruptly to an alternative state emerged from theoretical models’. Solutions to dynamic models with 1 or 2 equations show a range of responses of system state variables to linear increases in forcing (May 1977, Collie et al. 2004). These responses may be ‘smooth’ (i.e. linearly proportional), ‘abrupt’ (i.e. showing a runaway response) or ‘discontinuous’ (in which the system skips to a non-adjacent state), depending on parameter values in the equations. Boundary exchange can suppress such discontinuities (van de Koppel et al. 2008). These models provide insights into ecological dynamics, but may be too simple to quantify real ecosystem behaviour. Most theoretical thinking about regime shift has used conceptual rather than numerical or analytical models, such as the cliff and landscape metaphors in Figs. 3 & 4.

Under both landscape and cliff metaphors, a system’s approach to the point at which regime shift occurs is associated with a breakdown of resilience. In the most general, systemic terms, the breakdown is a temporary loss of negative (stabilizing) feedback loops, so that for a time the system becomes domi-
nated by positive (disruptive) feedback before settling into a new basin of attraction (either in a metaphorical landscape or in state space). As considered in the preceding subsection, more concrete explanations might involve loss of functional and response diversities, erosion of trophic networks or disturbances to ecosystem metabolism, associated with the symptoms of ecosystem pathology (Tables S2 & S3 in Supplement 2). A purely empirical explanation would centre on the loss of key stabilizing components; for example, explaining the decline of Alaskan kelp forests by increased echinoderm grazing following a decline in sea-otters (Estes & Duggins 1995), which was perhaps due to increased predation by killer whales deprived of their former prey, the great whales (Estes et al. 2004).

Substantial changes have already occurred in coastal ecosystems as a result of human removal of most large marine vertebrates, including whales, dugongs, turtles and larger fish (Jackson et al. 2001, Estes et al. 2004, Thurstan & Roberts 2010). The consequences include greater risk of eutrophication because of 'microbialization of the global coastal ocean' (Jackson et al. 2001). It is not clear that such changes would fully reverse were fisheries and other pressure to be relaxed and coastal environments restored (Duarte et al. 2009), and in any case, such relaxation and restoration might be difficult to achieve. Nevertheless, the depleted systems (although changed in organization) might retain sufficient functional diversity and resilience to continue providing some services despite external pressures.

The hypothesis that altered ecosystems might be healthy systems is of considerable practical importance. Although arguably incompatible with laws, such as the WFD, which equate good status solely with that of 'reference conditions' (see 'Ecosystem health as an aggregate property', above), the hypothesis is consistent with the requirements of other laws, such as the MSFD (Borja et al. 2012), that define 'good environmental status' as that of seas that are functioning well and providing for sustainable use.

THEORY OF ECOSYSTEM HEALTH

Definition and components of ecosystem health

As shown above, there are diverse opinions about the nature of ecosystems and what is to be understood by calling them healthy. Nevertheless, there is a need for guidance on how to protect marine ecosystems. One strategy is to seek consensus amongst a group of experts (e.g. Foley et al. 2010). Such a group (the authors of this article) met in Lowestoft, UK, at a series of workshops in 2010, tasked with: (1) developing a theory of ecosystem health, and (2) suggesting how this theory might be used for evaluating the holistic state of marine ecosystems in the context of European requirements to maintain good ecological status (WFD) and good environmental status (MSFD).

Participants agreed on a definition of health that took account of both the empirical and the systemic views:

Ecosystem health depends on: the physiological health of the constituent organisms; the characteristic properties and interactions of the species present; and the emergent properties of the system comprising the biota and their environment. Healthy ecosystems can sustain services to humans. They are vigorous, resilient to externally imposed pressures, and able to maintain themselves without human management. They contain organisms and populations that are free of stress-induced pathologies, and biodiversity that includes (1) a functional diversity enabling all biogeochemical and trophic functions appropriate to the ecohydrodynamic conditions, and (2) a diversity of responses to external pressures. All expected trophic levels are present and well interconnected, and there is good spatial connectivity amongst subsystems.

Any assessment of ecosystem health in relation to this definition must be made on appropriate spatial and temporal scales, and must take into account the local ecohydrodynamic conditions and the degree of openness of the system’s boundaries.

To the original 3 health components of vigour, organization and resilience (Mageau et al. 1995, Costanza & Mageau 1999), workshop participants added hierarchy (including spatial granularity) and trajectory, to take account of ecological variability in space and time (Table 2). We could not identify simple indicators of ecosystem health. Instead, we propose the use of trajectories in state space to assess change in ecosystem condition. The definition of health, and the proposal to use trajectories, constitutes a framework for understanding marine ecosystem health which we hope will influence programmes for monitoring coastal waters and managing pressures thereon.

Extent and granularity of an ecosystem

The first task in applying the framework is to define an ecosystem’s extent and describe its boundaries: where they are, what crosses them, and what is included within them. The boundaries may be fixed
Table 2. The proposed 5 components of ecosystem health

<table>
<thead>
<tr>
<th>Component</th>
<th>Definition</th>
<th>Explanation</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Structure or organization</strong></td>
<td>The types and arrangements or interconnections of the components of a system</td>
<td>If the system is thought of as a net, then this component comprises both the nodes and the links—the components and their connections; however, the nodes correspond to functional groups rather than species, and thus structure is not related to biodiversity as measured by number of species or related indices, but to functional diversity and (trophic) connectance.</td>
</tr>
<tr>
<td><strong>Vigour</strong></td>
<td>The ability of a system to maintain or renew its organization, drawing on production</td>
<td>Vigour concerns the functioning of the network—if that is seen as a set of pipes, then vigour concerns the fluxes through the pipes, and so would include primary production, nutrient cycling, and balance terms such as net production, or ratios such as that of production to respiration or allochthonous to autochthonous production; it might also include reproductive vigour and successional vigour, the potential for the biotic community to recolonize a region that has suffered disturbance.</td>
</tr>
<tr>
<td><strong>Resilience</strong></td>
<td>The capacity of a system to maintain its integrity under pressure</td>
<td>Resilience emerges (in the systemic view) from organization and vigour, and, empirically, may depend in part on key species and on functional response diversity; it is an ecosystem’s capability to maintain its functions and structure under external pressure, either by resistance to the pressure, recovery from its effects, or adapting to it; when this capability is exceeded then there is a regime shift.</td>
</tr>
<tr>
<td><strong>Hierarchy and granularity</strong></td>
<td>The distribution and interconnection of sub-systems and subregions within the ecosystem</td>
<td>Hierarchy includes both (1) the ecological equivalent of ‘subsidiarity’, the existence of subsystems that can to some extent function independently (e.g. pelagos, benthos), and (2) granularity, the existence of spatially distinct subregions that can re-seed adjacent subregions following disturbance; spatial connectivity refers to the links between these subregions; all these aspects (sometimes linked under the title of panarchy) might contribute to resilience.</td>
</tr>
<tr>
<td><strong>Trajectory</strong></td>
<td>A sequence of system states plotted in state space</td>
<td>Refers to change in the internal description of an ecosystem: the way in which system state, including structure and vigour, changes with time, resolved on an appropriate scale; trajectory, plotted in state variable space, and related to pressures, is diagnostic in that it allows resilience to be estimated, and could be prognostic (of decreasing or recovering health) if there was a theory or model that allows a domain of good health (high resilience) to be located in a state space diagram.</td>
</tr>
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</table>

by natural features, but are typically drawn by human custom or legislation. European examples include the ‘waterbodies’ of the WFD and the large marine ‘subregions’ of the MSFD (Borja et al. 2010).

The second task concerns how to deal with hierarchy: the existence of functionally distinct subsystems (such as pelagos and benthos) and spatial patchiness. Granularity can arise in the benthos from a patchwork of sediment types (Künitzer et al. 1992), or in the plankton from differing spatial and temporal patterns of mixing and stratification (Pingree & Griffiths 1978, Tett et al. 2007). It can also result from internal dynamics, such as the local settlement, growth and mortality of a cohort that temporarily dominates a region of benthos (Pearson & Barnett 1987), or from gradients of anthropogenic pressure, as with increasing distance from a fish farm (Brown et al. 1987, Nickell et al. 2003) or an urbanized coast (Garmendia et al. 2011). As discussed, resilience may in part derive from spatial connectivity and the possibility of one sub-region reseeding another after local collapse. Connectivity can be physical, for example when larvae are transported by water movements, or biological, for example, migration to exploit seasonally varying food supply. Life cycle closure theory (Sinclair 1987, Sinclair & Iles 1989) suggests that species’ populations adapt their migratory and reproductive behaviour to prevailing circulation patterns (Peterson 1998).

A variety of methods are available for observing spatial variability in marine ecosystems (Table S7 in Supplement 5). Such variation, if analysed in terms of ecosystem health, can be related to gradients of pressure and thus used to understand and manage the pressures that disturb health (COM 2010). In addition, we anticipate that the use of these methods will lead to a better understanding of the role of spatial variation in holistic ecosystem function. For now, we propose to smooth out such variability by spatially aggregating or averaging, and to deal with hierarchy by seeing it as part of system organization and hence by a suitable choice of system state variables.

**Organization and vigour**

The conclusions of the review were that key aspects of ecosystem organization are (1) spatial heterogeneity and connectivity, (2) functional diversity and response diversity, (3) trophic connectance, and (4) the existence of co-adapted species and stabilizing feedbacks. Vigour is the ability to maintain or renew organization (Costanza 1992). At the level of the system, it concerns biogeochemical fluxes, espe-
specially of compounds of carbon, nitrogen and phosphorus, and associated biochemical energy. At the level of species, it concerns population maintenance, requiring not only reproductive success but in some cases also adequate spatial connectivity. At the level of the community are successional processes, which in marine systems may include seasonal plankton cycles (Margalef 1978) or benthic recovery from disturbance (Pearson & Rosenberg 1976), and which are illustrated as the doughnut-shaped bundles of trajectories in Fig. 1. Finally, there is the idea of balance to be considered: not a static balance in which every part of the food web is in harmony, but a dynamic and panarchic balance in which populations fluctuate and are replaced, allowing ecosystems to adapt to pressures.

Understanding these issues, and their contribution to ecosystem resilience, is a crucial challenge for those studying community ecology. Some progress has been made since Holling’s (1973) pessimistic conclusion that estimating resilience, by measuring the extent of the ‘domain of attraction’ in state space, will ‘require an immense amount of knowledge of a system and it is unlikely that we will often have all that is necessary’ (p. 20). Salihoglu et al. (2013) report progress in coupling community ecology with observations and modelling of biogeochemical cycles in large marine ecosystems. Nevertheless, it is not yet possible to state, in quantitative terms, a functional relationship between organization, vigour and resilience.

Trajectories in state space

Given the lack of sufficient theory-based understanding of the holistic functioning of ecosystems, we opted for a more pragmatic approach. This involves the analysis of trajectories in state space, and is compatible with both empirical and systemic views.

Movement in state space has several components (Fig. 1): repeating or semi-cyclical variation, medium-term variability, and long-term movement relative to a reference condition. State space plots do not include a time-axis, but variation such as that associated with seasonal cycles of plankton will show as loops, which might be seen as part of ecosystem organization (Tett et al. 2007). Similar semi-closed trajectories might result from succession (and return to climax) after disturbance. Although the phenology of seasonal change is itself important (Racault et al. 2012), the topic of cyclical variation is outside the scope of the present review, and we smooth it by plotting annual mean states. ‘Medium-term’ variability is the difference between successive means or the deviations of these means from a long-term trend.

Estimation of such a trend requires a reference point from which (or to which) distance can be calculated. For present purposes this reference point is arbitrary, and need not be identified with formerly pristine conditions or with any desired ecosystem state.

Scalar distance in state space

Simple mathematical systems and controlled laboratory experiments can fully describe system state by 1 or 2 variables, and can control all except 1 dimension of pressure; but this is unlikely to be true of natural marine ecosystems and the multiple pressures to which they are exposed. Yet it aids explanation and understanding if, as in Fig. 3, multidimensional changes in ecosystem state and in pressure can be shown as single variables. Such a simplification can be made in several ways: (1) given the identification of a reference domain in state space, observations might be characterized as either within or outside this domain, as in the ‘Phytoplankton Community Index’ (Tett et al. 2008); (2) by using a principal axis obtained by multivariate (statistical) analysis (Muxika et al. 2007, Kenny et al. 2009); or (3) calculate the unidimensional Euclidian distance between points in state space (see Fig. 1, Supplement 6). It is this latter option that we adopt in the next section, to estimate distance travelled from a reference condition.

A state space approach, with orthogonal axes, implies that all state variables are of equal worth in quantifying state, and thus potentially avoids one of the difficulties in aggregating ecosystem status components. Furthermore, because the axes are orthogonal, it is unnecessary to require variables to be measured in the same units. However, it is desirable to standardize the axes in some way, for convenience in viewing state space diagrams as well as for combining movements along the axes into a scalar quantity. Logarithmic transformation of planktonic data is often adopted in the interests of normalizing variability (Barnes 1952) and has the additional advantage of approximating relative change (since ln[x] = ΔX/X for small changes) and homogenizing units prior to combination in Euclidian distance. Multivariate analyses typically go on to divide variables by their standard deviations, but this may be a step too far when assessing movement in state space: it may be desirable to give more weight to variables that are relatively more changeable.
Pressure-state model based on an elasticity metaphor

It is an axiom of the DPSIR (and similar) frameworks that a change in pressure causes a change in ecosystem state. According to the systemic view of ecosystem health, the extent of the state change for a given pressure change is a function of the resistance component of the system’s resilience (Walker et al. 2004). Our proposal is to relate change in ecosystem state to change in pressure on the analogy of Hooke’s Law of elasticity in mechanical systems. It is assumed that values of the state and pressure variables have been averaged over appropriate spatial and temporal scales so as to remove effects of spatial heterogeneity and cyclical variation. We define ecosystem compliance (Fig. 3) as the ratio of state change to pressure change, i.e. as the analogue of the coefficient of elasticity in Hooke’s Law and the reciprocal of ecological resistance. An ecosystem with high resistance will comply only weakly with pressure increases, so long as the system remains (in the elasticity analogue) within its elastic limit, within a valley in the landscape metaphor (Fig. 4), or before it reaches the cliff in Fig. 3.

Scheffer et al. (2009, 2012; see also Lenton 2013) suggested that increased variability was a sign of decreasing system resilience, and hence could provide an early warning of regime shift. Such variability can be demonstrated numerically in solutions of simple models (Collie et al. 2004). Ecosystems are more complex than these models, but the mechanistic explanation can be used as an analogy: a system on the cusp of a regime shift can be thought of as being pulled in several directions by ‘attractors’ for the old and new regimes. If it is also subject to other dynamic fluctuations, it may experience large fluctuations in condition before settling down into a new regime. The elasticity metaphor for resilience also leads to the prediction (Supplement 6) that medium-term variability will increase as resilience decreases, without any corresponding increase in pressure variability. Thus, we propose to use medium-term variability as a proxy for (inverse) resilience, without further considering pressure-state relationships.

Data requirements

The proposal to track the state of an ecosystem by plotting its condition in successive periods as a sequence of points in state space does not strongly constrain the choice of state variables. Given self-standardization to ensure that axes are of equal worth, there is not even a requirement for consistency in units. Thus, it should be possible to use variables that reference system organization alongside those that refer to vigour. It is, however, crucial that the selected variables should comprise an overall view of an ecosystem, whilst avoiding redundancy.

Another requirement is for time-series that are long enough for trends to be distinguished from the medium-term variability used as a proxy for resilience in the type of analysis that we propose. The next section contains an example, not intended to be definitive, of a state space analysis of a few time-series of adequate length from a large marine waterbody: the North Sea.

EXAMPLE OF THE STATE SPACE APPROACH

The North Sea ecosystem

The North Sea is a large continental shelf sea which exchanges with the northeast Atlantic Ocean across its northern margin and, indirectly, through the Strait of Dover at its southern extremity. It receives significant freshwater inputs from the Baltic Sea via the Kattegat in the east, and from major rivers such as the Rhine along its southeastern flank (Rodhe 1998). Its ecosystems are subject to pressure from fisheries, nutrient enrichment, seabed disturbance and toxic pollutants (Ducrotoy & Elliott 2008).

The North Sea can be viewed as a semi-enclosed box with a mean flushing time of about 1 yr, and thus as a single ecosystem over which trophic fluxes can be averaged (Rodhe et al. 2006). Alternatively, the sea can be sub-divided into ecohydrodynamically distinct pelagic regions (Tett et al. 1993, Rodhe et al. 2006), zones of benthic associations related to sediment type (Künitzer et al. 1992), or areas shown to be similar by empirical statistical analysis (Kenny et al. 2009).

To avoid difficulties in aggregating data from different ecohydrodynamic regions, we focus here on the northwestern part of the North Sea (Fig. 5), a region defined by the Continuous Plankton Recorder Survey as Standard Areas B2 and C2 (Reid et al. 2003). Offshore waters of this region range in depth from ~60 to 120 m deep and are thermally stratified during late spring, summer and early autumn.

Choice of state variables

Three variables were chosen to represent different parts of the food web. They were the rearing success
of black-legged kittiwakes *Rissa tridactyla*, the abundance of the copepod genus *Calanus*, and simulated net annual water column primary production. In the terms introduced in the section ‘Health as an emergent property of complex systems’ above, primary production and kittiwake rearing success might be taken as indicators of vigour, and the 3 variables collectively indicate the organization of the food web. The variables are not homogenous either in their units, or as functional entities. Nevertheless, they serve to demonstrate the state space method and to show that it does not impose strong constraints on the data that may be used.

Data for kittiwake rearing success, defined as chicks fledged pair−1 yr−1, were taken from several sources. Annual averages for all Scottish nesting sites observed by the UK Seabird Monitoring Programme (SMP) between 1986 and 2008 were taken from JNCC (2011). Most of these sites (>75% of those in Scotland) are on the Orkney and Shetland Islands and the east coast of Scotland (Mitchell et al. 2004). Five-year running means for 1956 through 1980 (Coulson & Thomas 1985) from a single kittiwake colony in North Shields (northeast England, at 55.0°N, on the southern boundary of the study region) were used to extend the series. Turner (2010) reports rearing success for the same colony in 2000.

The rearing success of kittiwakes is presumed to depend on the abundance of juvenile herring (Coulson & Thomas 1985) or sand-eels (Wanless et al. 2007) near the surface of the sea. Feeding areas were assumed to be mainly in the offshore North Sea.

*Calanus* spp. numbers were taken from Continuous Plankton Recorder (CPR) surveys (Warner & Hays 1994). These may underestimate real abundance (Dippner & Krause 2013), but what we require is relative consistency over the time-series. Annual means were calculated from counts on samples from CPR regions B2 and C2. Sample values (copepods per tow unit, i.e. in about 3 m³ of water) were interpolated for each month onto a uniform grid of 1° longitude by 0.5° latitude (Vezzulli & Reid 2003). We combined the abundances of *Calanus finmarchicus* and *C. helgolandicus*, viewing the genus as an example of a functional type, with the species providing functional response diversity.

Net annual water column primary production (g C m⁻²) was estimated from a North Sea hindcast with European Regional Seas Ecosystem Model (ERSEM; Baretta et al. 1995, Ruardij & Van Raaphorst 1995) linked (Lenhart et al. 2010, van Leeuwen et al. 2013) to the General Estuarine Transport Model (GETM; Burchard & Bolding 2002). Climatological boundary conditions (i.e. a 1 yr cycle) were used for nutrients along the simulated Atlantic margin. Annual totals of production were averaged over areas B2 and C2 (Fig. 5) from model gridpoints between 55.0°N and 60.4°N, west of a line from 4.49°E (at 55.0°N) to 3.02°E (at 60.4°N).

To obtain the co-ordinates in state space for a starting point for the trajectory, we (1) averaged CPR *Calanus* data for 1958 to 1962, resulting in a mean value of 25 copepods sample⁻¹; (2) averaged simulated production for 1958 to 1962, resulting in a mean value of 137 g C m⁻² yr⁻¹; (3) assumed a kittiwake breeding success of 1.4 fledglings nest⁻¹ based on the mature stage (around 1960) of the colony reported in Coulson & Thomas (1985).
Time-series plots

Average net primary production simulated by GETM-ERSEM was 130 g C m\(^{-2}\) yr\(^{-1}\), with 90% of values lying between 115 and 151. This is higher than the range (50 to 100 g C m\(^{-2}\) yr\(^{-1}\)) of results from other models and observations (Supplement 7). Values were slightly lower in the middle part of the time series (Fig. 6c), but there was no obvious long-term trend. This may be because the simulation used climatological northern boundary conditions. Heath & Beare (2008) calculated new production from the observed spring draw-down in nutrients in each year between 1960 and 2003, and obtained higher values—implying greater inflow of Atlantic water—during the first 2 decades.

Calanus abundance (Fig. 6b) showed an early decrease with some recovery in recent decades, perhaps associated with observed changes in planktonic communities (Beaugrand et al. 2002). The kittiwake data (Fig. 6a) also show a steady decline (from the start of the SMP time-series in 1986), possibly the result of changes in the availability of its prey fishes, in turn influenced by fisheries and by climate fluctuations (Coulson & Thomas 1985, Furness & Tasker 2000, Wanless et al. 2007). The agreement between the SMP data and Turner’s (2010) measurement of rearing at the North Shields colony in 2000 lends confidence to combining early and recent kittiwake data.

The data were plotted in 3 dimensions using a Matlab script and standard 3D plotting and viewing functions. Fig. 7 shows the trajectory of annual means through the state space. No transformations have been applied at this stage. There are 3 obvious features to this trajectory. First, there is much interannual variability. Second, there has been a long-term decrease in kittiwake rearing success and in Calanus abundance. Third, consequent on the second point, the system has moved a long way from the conditions ca. 1960. There is, however, no indication of sudden change from one contrasting and persisting state to another, i.e. no evidence for a regime shift as defined by deYoung et al. (2008).

Fig. 8a shows the standardized Euclidian distance (Fig. 1) of each annual point in Fig. 7 from the arbitrary 1960 ‘reference condition’, and confirms the extent and continuity of the change. Annual scalar variability (Fig. 8b, hypothesized to be proportional to compliance and inverse resilience) was calculated as the absolute deviation from the trend-line fitted to this change. Although only the copepod data were taken from a single monitoring programme, the 11 yr running mean (labelled ‘smoothed’ in Fig. 8b) suggests an initial trend of increasing variability, followed perhaps by a weak trend of a decrease. These results are presented to exemplify the state space approach rather than to test hypotheses about change.
and so we have not carried out rigorous time-series analysis. Nevertheless, a third order polynomial explained a significant part of the variability in the data and confirmed the pattern in the running mean.

**Assessment**

The 3 state variables used in this example were intended to capture 3 elements of the trophic network in the North Sea. They were also chosen because adequately long time-series of each were available and sufficiently understood to explain and caveat the conclusions reached from the analysis. Clearly, ‘adequately long’ implies half a century or more in these waters, in order to take account of natural fluctuations and the decadal time-scales of observed ecosystem changes (Kenny et al. 2009).

Several studies have deduced a regime shift in the North Sea centred either on 1988 (Edwards et al. 1988a).
2001, Reid et al. 2001, McQuatters-Gollop et al. 2007) or on 1993 (Kenny et al. 2009). Such a shift is not evident in Fig. 7, which is more compatible with the conclusion by Spencer et al. (2011, p. 19) that ‘changes in UK marine communities appear to be dominated by gradual trends over the last two to three decades’. However, Fig. 8b suggests increasing variability (about the long-term trend in Fig. 8a) until about 1980. If this is interpreted (Scheffer et al. 2009) as a decrease in ecosystem resilience, it would be compatible with an approach to a gradual regime shift during the 1980s.

SYNTHESIS AND A LOOK FORWARDS

Holistic approach

Marine ecosystems include co-adapted species linked through trophic networks and biogeochemical cycles, with the consequence that disturbance to some species’ populations can impact on those of other species, or modify ecosystem services that depend on interactions among ecosystem components. It is necessary to understand these links, and to identify whole-ecosystem quality objectives and assessment methods, if environmental managers are to fully protect ecosystem services and ensure their sustainable use under pressure. Several integrated assessment methods have been proposed in recent years. We consider 3 examples, each with a different approach to the challenge.

Working within the same (European) policy framework as us, Borja et al. (2011) suggested an aggregative approach to GES under the MSFD. The issue here is the weights used in the combinatorial process, which must be decided by expert judgement. Halpern et al. (2012) proposed a generic ‘ocean health index’ assembled from public goals for marine ecosystem services. This may allow too much focus on particular uses of marine systems, even if the sustainability of such use depends on the maintenance of good ecosystem health, and it provides little insight into the functioning that underpins health. Closest to our proposed use of state space is the method used by Hemery et al. (2008), who took the first principal component from a multivariate analysis as an overall index of changing ecosystem state in the Bay of Biscay. Such use of principal component analysis (PCA) to estimate a scalar distance in a multivariable state space contrasts with our use of Euclidian distance.

Management of ecosystems requires explanation as well as measurement of change. Our contention is that a theory of ecosystem health—even if yet to be fully developed—can provide a basis for understanding ecosystem functioning and for monitoring and managing marine ecosystems in relation to their intrinsic worth as well as their value to human society. According to the systemic approach, persistence of an open system depends on the maintenance of functional integrity whilst processing throughputs of energy and materials. Resilience is the ability to maintain integrity despite changes in boundary conditions. It depends on the organization and vigour of ecosystems. Organization, which refers to ecosystem components and their interconnections, includes functional-group diversity and functional-response diversity, the occurrence of co-adapted species, and the existence of multiple and alternative trophic pathways and of stabilizing feedbacks. Vigour is the flow of energy and materials that maintains organization. Panarchy takes account of system heterogeneity on multiple scales in space and time, which may contribute to resilience.

State space method

Resilience is the crucial aspect of ecosystem health, and sustaining it should be the prime objective of ecosystem managers (Gunderson 2000), because resilient systems can maintain themselves against pressures and change. The challenge is to quantify its elements: resistance, latitude and precariousness (Walker et al. 2004). In the absence of sufficient ecological theory to compute these from directly observable properties, such as biodiversity (as relating to organization) and production (as a measure of vigour), we have proposed the use of a state space approach to track changes in ecosystem condition (Table 3). As discussed in Supplement 6, relating changes in state to changes in pressure might allow models for system compliance (the inverse of the resistance component of resilience) to be parameterised empirically. In this study, however, our focus has been on ecosystem state and its changes rather than on the causes of those changes.

Our method requires the identification of variables that define a state space and capture the most important aspects of ecosystem organization and vigour. The review in ‘Organization and biodiversity’ above suggested that focusing on functional diversity is most likely to provide insights into ecosystem function, and thus, ideally, that the state variables chosen for plotting should represent functional groups. The variables used in our North Sea example were some
The suggestion to plot marine ecosystem data in state-space is not new. Margalef (1978) drew a 2-D space, defined by conditions in the physico-chemical environment, to show the niches preferred by different types (lifeforms) of phytoplankter, and to theorize how the state of the phytoplankton community responded to seasonal or other changes in the physical environment. More recently, Bald et al. (2005) and Muxika et al. (2007) used state space plots, from which they extracted principal components, to characterize both the pressures on, and the ecological state of, the shallow-water benthos.

Finally, the state space method is based on a definition of ecosystem that requires identification of a region within defined boundaries. Ideally these boundaries would be natural discontinuities. In practice they are determined by management considerations, and in consequence the system thus defined may contain a patchwork of physical and chemical environments and biotic communities. In order to avoid issues resulting from such heterogeneity, we chose the comparatively homogenous northern North Sea for our example. It is smaller than the smallest region (the ‘Greater North Sea’) allowed as an assessment unit by the MSFD, and we have not considered issues relating to spatial patchiness and panarchy.

### Table 3. Summary of the proposed method for tracking change in ecosystem state and estimating resilience

<table>
<thead>
<tr>
<th>Steps of the method</th>
<th>Example used in this study</th>
<th>Issues</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. Identify (spatial extent) of ecosystem</td>
<td>Northwestern part of the North Sea (see Fig. 5)</td>
<td>Boundaries determined by natural conditions or by human custom or legislation; extent of trans-boundary fluxes</td>
</tr>
<tr>
<td>2. Identify spatial granularity and extent of repetitive temporal variability, and decide how to average or aggregate over these</td>
<td>Treated as a single spatial unit and averaged or aggregated over a year</td>
<td>Granularity and hierarchy may contribute through panarchy to resilience; ecosystem may contain several biomes; seasonal variability may be seen as organization</td>
</tr>
<tr>
<td>3. Select state variables</td>
<td>Annual net primary production (from simulation); mean annual abundance of copepods <em>Calanus</em> spp. (from Continuous Plankton Recorder [CPR] survey); kittiwake chick rearing success (mean over coastal colonies supported by the marine ecosystem under consideration) (see Fig. 6)</td>
<td>Variables should broadly represent the ‘condition’ of the ecosystem, including its organization and vigour; they should comprise time-series extending over several ‘natural periods’ for that ecosystem</td>
</tr>
<tr>
<td>4. Plot trajectory in state space and calculate Euclidian (scalar) distance from (arbitrary) reference condition</td>
<td>3D state space plot in Fig. 7; scalar distance from 1960 ‘reference condition’ in Fig. 8a</td>
<td>Euclidian distance is one method to reduce a vector distance to a scalar; other options for simplification of a n-dimensional trajectory include extraction of principal component(s) and the enumeration of points in relation to a reference envelope</td>
</tr>
<tr>
<td>5. Calculate medium-term variability about trend in state space, and use this variability as proxy for (inverse) resilience</td>
<td>Trend established by fitting third order polynomial; variability measured as annual deviation from this trend (see Fig. 8b)</td>
<td>The distinction between repetitive, medium-term and long-term variability; the differences in natural variability amongst ecosystems</td>
</tr>
</tbody>
</table>

The results reported here involved the calculation of Euclidian distance from an arbitrary reference condition, and the comparison of variability over time. This procedure is applicable to data from any type of ecosystem, irrespective of the state space coordinates of the reference condition or the system’s natural variability. However, although not required by the method, the reference condition would ideally lie within a state space domain corresponding to good health and maximum resilience, and we see an important goal for quantitative ecological theory as being the specification of such domains in a variety of estuarine and coastal ecosystem types. The identification of these domains might, we think, be achieved in part through the analysis of existing time-series of marine ecosystem states, pressures, and services, and in part through the development of generic models for relationships between pressures and ecosystem states. Ideally, these models will be of sufficient generality to apply to systems of naturally low species diversity, such as those found in estuaries (Elliott & Whitfield 2011), as well as to the more species-rich open sea ecosystems.

An interesting question concerns whether there is a unique domain of health for a given ecosystem type, i.e. under given ecohydrodynamic conditions. This would be the case for Clement’s (1936) concept of a climax community and the argument (Moss 2008)
that natural ecosystems are best adapted to their environments. It is the view taken by the WFD, which equates ‘Good Ecological Status’ with the structure and function of aquatic ecosystems that are similar to those of the same type under undisturbed conditions. In contrast, the view of systems as multiple interconnected networks suggests that they can exist in more than one stable (i.e. resilient and healthy) configuration (Krebs 1988). It is even possible that ecosystems that have become impoverished in species can function well and can demonstrate resilience as a result of adaptation within populations of generalist species, as may naturally be the case for estuaries (Elliott & Whitfield 2011). The possibility of more than one domain of good ecosystem health would seem to be allowed for within the MSFD, which for GES simply requires ecosystems to ‘function fully’ within the constraints set by the intrinsic environmental conditions and to ‘maintain their resilience to human-induced environmental change’. If each stable configuration can deliver a set of sustainable services, albeit different ones, the issue then becomes that of deciding which set is preferred by society.

**Data requirements**

The role of biodiversity in relation to ecosystem function, and the causes and nature of regime shifts, have become better understood as increasing amounts of data and increasing numbers and lengths of time-series have become available from a variety of ecosystems. Our proposed method also needs long time-series to track changes in system state space, to estimate changes in variability as a proxy for resilience, and to seek empirical relationships with pressures. Although there is potential in remote sensing (Platt & Sathyendranath 2008, Platt et al. 2010), multi-decadal oceanic ecological datasets are rare (Edwards et al. 2010). In general, research funding is inadequate to collect sufficient data, over sufficiently long periods of time, to test hypotheses relating to changes in ecosystem health. The relevant time-scale of variability shown by our North Sea example is, at least, decadal, (see also Kenny et al. 2009). The most effective drivers of data-collection on this long time-scale are (1) fisheries management, (2) conservation of protected species, and (3) environmental protection legislation—and we suggest that data collected routinely for such purposes might also be used for research into the key issues raised in this study concerning ecosystem health (Supplement 8). Although it may be that the research questions ought to influence the choice of variables observed during these programmes, we consider that the holistic approach to ecosystems should add little to a programme’s overall cost, because the extra work required lies mainly in the numerical analysis exemplified in ‘An example of the state space approach’, above.

**Metaphors and models**

Data alone are insufficient for understanding and predicting changes in ecosystem health. Complex systems do not function in ways that can easily be understood by common sense, nor on spatial or temporal scales that map well to human polities and democratic political processes. Nevertheless, society needs explanations that can justify the management of human pressures that act on the holistic condition of marine coastal ecosystems. The idea of ecosystem health is, in its most general form, a way of translating complex system behaviour into a widely and intuitively comprehensible explanation. However, as Lackey (2001) and others have argued, the use of health as a metaphor is open to abuse through the covert insertion of sectional values. Thus, there is a need to describe system behaviour in terms that are scientifically sound (i.e. open to falsification) but sufficiently simple and transparent to communicate changes in system state and vulnerability to non-specialists. That is why metaphors for system resilience have been found useful.

This is not to deny the relevance and utility of other types of models (Table 4). Simple mathematical models with 1 or 2 state variables can demonstrate bifurcations (Collie et al. 2004), the importance of trophic connectance (Vallina & Le Quéré 2011), and the role of boundary conditions (van de Kopp et al. 2008). Complex *mechanistic models* (such as ERSEM; Barreta et al. 1995, Allen et al. 2001, Blackford et al. 2004)—in which the model building blocks represent functional groups, and the system can be replicated on spatial grids to include the effects of granularity—can provide detailed internal descriptions of ecosystems. ERSEM was used to provide the production data in ‘An example of the state space approach’, above. Improved understanding of functional diversity and functional response diversity (see ‘Organisation and biodiversity’, above) should help in allowing such models to fully represent ecosystem processes and their adaptive responses to changes in external forcing. Complex *empirical models*, such as EcoPath (Pauly et al. 2000, Christensen & Walters 2004) for
Table 4. Examples of use of models in relation to ecosystem health

<table>
<thead>
<tr>
<th>Generic type of model</th>
<th>Epistemology</th>
<th>Simple model examples</th>
<th>Complex model examples</th>
</tr>
</thead>
<tbody>
<tr>
<td>Conceptual and</td>
<td>Used to explain</td>
<td>Landscape (including cliff and basin) (Holling 1973, Krebs 1988, Scheffer et al. 2001) and elasticity (the present study)</td>
<td>Analogies (such ‘attractors’) derived from mathematical properties of systems of equations (Ives &amp; Carpenter 2007); networks (e.g. food webs based on stomach contents; Hardy 1924)</td>
</tr>
<tr>
<td>metaphorical</td>
<td>using ‘common sense’ analogies</td>
<td>(the present study) metaphors for emergent properties such as stability and resilience</td>
<td></td>
</tr>
<tr>
<td>Mechanistic</td>
<td>Assembled from qualitative hypotheses tested under controlled conditions—i.e. simulating causation</td>
<td>Lotka-Volterra and similar models (and their predictions about bifurcation, regime shift, etc. (Collie et al. 2004, van de Koppel et al. 2008, Vallina &amp; Le Quéré 2011)</td>
<td>Multi-component physically-coupled ecosystem models, e.g. European Regional Seas Ecosystem Model (ERSEM; Baretta et al. 1995, Blackford et al. 2004); complex model parameterization difficulties discussed by Cropp &amp; Norbury (2013)</td>
</tr>
<tr>
<td>Empirical</td>
<td>Fitted to data (e.g. by minimizing sum of squares or maximizing likelihood)—i.e. summarizing correlation</td>
<td>Statistical: regressions and trends; by-analogy models such as that for compliance corresponding to elasticity in the analogue of a mechanical spring</td>
<td>MultiVariate Analysis (MVA) used for detecting regime shift; Bayesian belief networks (Langmead et al. 2009); network models fitted by parameter estimation</td>
</tr>
</tbody>
</table>
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