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**Historical records of  
hypoxia**

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# Historical records of coastal eutrophication-induced hypoxia

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## Abstract

Under certain conditions, sediment cores from coastal settings subject to hypoxia can yield records of environmental changes over time scales ranging from decades to millennia, sometimes with a resolution of as little as a few years. A variety of biological and geochemical proxies derived from such cores have been used to reconstruct the development of eutrophication and hypoxic conditions over time. Proxies based on 1) the preserved remains of benthic organisms (mainly foraminiferans and ostracods), 2) sedimentary features (e.g. laminations) and 3) sediment chemistry and mineralogy (e.g. presence of sulphides and redox-sensitive trace elements) reflect conditions at or close to the seafloor. Those based on 4) the preserved remains of planktonic organisms (mainly diatoms and dinoflagellates), 5) pigments and lipid biomarkers derived from prokaryotes and eukaryotes and 6) organic C, N and their isotope values reflect conditions in the water column. However, the interpretation of these proxies is not straightforward. A central difficulty concerns the fact that hypoxia is strongly correlated with, and often induced by, organic enrichment (eutrophication), making it difficult to separate the effects of these phenomena in sediment records. The problem is compounded by the enhanced preservation in anoxic and hypoxic sediments of organic microfossils and biomarkers indicating eutrophication. The use of hypoxia-specific indicators, such as the trace metals molybdenum and rhenium and the bacterial biomarker isorenieratene, which have not been used often in historical studies, may provide a way forward. All proxies of bottom-water hypoxia are basically qualitative; their quantification presents a major challenge to which there is currently no satisfactory solution. Finally, it is important to separate the effects of natural ecosystem variability from anthropogenic effects. Despite these problems, in the absence of historical data for dissolved oxygen concentrations, the analysis of sediment cores can provide plausible reconstructions of the temporal development of human-induced hypoxia, and associated eutrophication, in vulnerable coastal environments.

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# 1 Introduction

Nutrient enrichment leading to eutrophication, and in some cases hypoxia, is one of the profound impacts inflicted on coastal ecosystems by human activities. These impacts began thousands of years ago (Jackson et al., 2001), but have increased in frequency and intensity since the middle of the 20th Century. Because marine eutrophication has only been acknowledged seriously since the mid-1980s (Nixon, 1995; Boesch, 2002), few long time-series environmental datasets are available from marine coastal settings (Clarke et al., 2003, 2006). Datasets extending back several decades (e.g. Justić et al., 1987; Justić, 1991), together with the comparison of historical and modern surface sediment samples (Alve and Murray, 1995; Rabalais et al., 1996; Thomas et al., 2000; McGann et al., 2003; Tsujimoto et al., 2006a), provide key information about natural, climate-induced variability and human influences that is necessary for the sustainable management of coastal habitats (Jackson, 2007). Historical records, e.g. dates of agricultural settlement and fisheries records, can also be useful. Such information is usually too limited, however, to provide a complete picture of past events. The analysis of proxy records preserved in sediments often provides the only way to reconstruct environmental change in areas impacted by eutrophication and hypoxia and to establish pre-impact baselines (Jackson et al., 2001; Alve, 2006). Because humans are changing coastal environments everywhere, these proxies may also provide our only glimpse of past, natural conditions and their inherent variability.

Certain conditions are necessary for the preservation and recovery of sediment records of coastal hypoxia. In addition to the availability of datable sediment cores, requirements include a relatively sheltered, low-energy setting that permits the deposition of fine-grained sediments, a sedimentation rate that is sufficient to allow for the resolution of the events of interest, and limited bioturbation and taphonomic processes that do not destroy the record (Murray and Alve, 2002; Clarke et al., 2006). These conditions can be found in estuaries, on continental shelves off the mouths of major rivers, in fjords, and areas where tidal disturbances are minimal. The time periods investi-

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gated range from decades to >2000 years, although the most intense human impacts generally occurred during the 20th Century. Where sedimentation rates are high, temporal resolution may be as little as a few years, at least for the 20th Century (Cronin and Vann, 2003; Turner and Rabalais, 1994). Studies of hypoxia in the sediment record began in the 1980s (e.g. Brush, 1984; Tyson and Pearson, 1991) but significant numbers of publications only began to appear in the 1990s and have increased substantially since 2000. They have tended to concentrate in certain areas, notably the Louisiana continental shelf, Chesapeake Bay, Norwegian fjords and the Baltic.

A variety of hypoxia and eutrophication proxies, most of them biological or geochemical in nature, have been applied to sediment records. In this review, we are concerned mainly with records that span time scales during which human influences on coastal environments have left an obvious mark. These historical records can be regarded as a facet of palaeoceanography. Indeed, many of the proxies were originally developed to reconstruct palaeoenvironments in Cenozoic oceans (Fischer et al., 1999; Hillaire-Marcel and Vernal, 2007). Where appropriate, we cite some of this geological literature, for example, that relating to the intensively studied Mediterranean sapropels (Jorissen, 1999b; Jorissen et al., 2007). In addition to human influences, natural environmental oscillations leave their imprint in sediment cores. A central issue when considering sediment records of coastal hypoxia is the close relationship between hypoxia (the degree of oxygenation) and eutrophication (which affects organic input to the seafloor). These two parameters are usually strongly correlated, and it is often not obvious which of them causes the changes observed in the sediment record. For example, foraminiferal species considered to be low-oxygen indicators are also associated with organic enrichment in the absence of bottom-water hypoxia, suggesting that they respond to changes in productivity rather than bottom-water oxygenation (Jorissen et al., 2007). Another problem is that hypoxic and anoxic bottom water conditions may cause better preservation of organic matter and biomarker proxies (Hedges and Keil, 1995), partly through the reduction of animal activities (Cowie and Levin, 2009), making it difficult or impossible to distinguish hypoxia from enhanced carbon delivery

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due to eutrophication.

The deteriorating state of many near-shore habitats makes the study of hypoxia an urgent task. SCOR Working Group 128 aimed to synthesize the state of the science for the following aspects of coastal hypoxia: 1) its prevalence and spatio-temporal variability, 2) natural and human causes, 3) its effects on biogeochemistry and ecology, and 4) the resistance, resilience and recovery of ecosystems. Historical records preserved in sediment cores reveal information that is relevant to all of these goals. Unravelling the ecological history of vulnerable coastal environments is crucial for their management and for setting restoration and remediation targets. To paraphrase Johnson (2007), palaeoecological studies provide baselines for forecasting the consequences of environmental change, the best way to assess ecological responses to climate change, and a basis for studying events and processes for which there is no modern analogue. This review addresses biological, chemical and sedimentary indicators that relate to either bottom-water hypoxia or eutrophication, or to a combination of these two phenomena. With the objectives of WG128 in mind, our goals are 1) to identify indicators and proxies that reveal the deterioration and recovery of ecosystems over time, 2) explore the effects of environmental change on ecosystems, 3) identify ways to distinguish natural from anthropogenic influences, and 4) suggest potentially useful new proxies and gaps in our knowledge. First, we review indicators for hypoxia and eutrophication based on benthic organisms and diagenetic processes (Sect. 3). Water-column derived proxies are presented in Sect. 4. Following the discussion (Sect. 5), we summarize our findings and identify new avenues of research.

## 2 Terminology

Geologists and biologists apply different terminologies to describe degrees of oxygen depletion and the various resulting biofacies (Tyson and Pearson, 1991; Jorissen et al., 2007). Geologists distinguish between oxic, suboxic, anoxic and euxinic (sulphide bearing) environments and biologists between oxic, hypoxic and anoxic environments;

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the term “exaerobic” was introduced for a biofacies characterised by laminated sediments combined with an accumulation of shelly fossils, reflecting short-term colonization of the seafloor by opportunistic species (Savrda and Bottjer, 1991) (Fig. 1). Some geochemical proxies distinguish between environments with and without hydrogen sulphide in the bottom waters.

### 3 Sedimentary indicators of hypoxia and eutrophication

#### 3.1 Faunal indicators

Microfossils derived from prokaryote, protistan and metazoan organisms have been used for many years as indicators of palaeoenvironmental conditions in the geological past (Lipps, 1993). Their application to historical time scales is a more recent development that falls within the scope of environmental micropalaeontology (Martin, 2000; Haslett, 2002). Although many dominant macro- and meiofaunal groups (e.g. polychaetes and nematodes) are almost never preserved intact, some small benthic organisms, notably the Foraminifera and Ostracoda, have outstanding fossil records and may be used as proxies for the responses of benthic communities to past environmental changes. A main advantage of microfossils over larger macrofossils is that numerous complete specimens can be recovered from relatively small sediment samples. One disadvantage is that they are easily transported and the possibility exists that assemblages include a significant allochthonous component. Moreover, fossil assemblages are normally time-averaged. Extended anoxic periods will result in sediments devoid of microfossils, except where allochthonous elements have been transported laterally. If the study area has been subject to one or more short re-oxygenation phases, the time-averaged deposit will contain autochthonous faunal remains. In many cases, especially when the time period under consideration has been dominated by anoxia, the microfossils present will be typical for the most oxygenated conditions which occur at the site. An alternation of anoxia with varying degrees of hypoxia will result in a com-

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plex faunal mix, making it impossible to unravel the exact oxygenation history from the time-averaged sample. Historical studies of oxygen depletion often disregard this problem.

In this section, we consider benthic faunal indicators which primarily reflect conditions in and on the sediment and not necessarily in the overlying water column. In most coastal sediments oxygen is depleted within the first 1–2 cm of the sediment column. Sediments underlying anoxic bottom waters differ from anoxic sediments underlying normal, oxygenated bottom waters because they contain different benthic communities and re-oxidation of reduced substances does not occur. This has major consequences for sediment energy balance and metabolism (e.g., the significance of chemoautotrophic communities) and nutrient exchange across the sediment-water interface (efficient recycling or retention of nitrogen and phosphorus) (Middelburg and Levin, 2009). As indicated above, the separate effects of hypoxia and eutrophication are difficult to disentangle, and it is often unclear which is exerting the predominate influence on assemblages of indicator species.

### 3.1.1 Foraminifera

Following recent molecular phylogenetic studies (Schweitzer et al., 2007), we consider the order Rotaliida to include groups such as the buliminids, bolivinids and cassidulinids, which constituted a distinct order, the Buliminida, in some earlier classifications.

Jorissen et al. (2007) give an overview of faunal proxies based on foraminiferans, the most widely-used benthic taxon for palaeo-reconstructions of bottom-water hypoxia and other environmental parameters (Murray and Alve, 2002; Murray, 2006). These testate protists are distributed in all marine environments, are sensitive indicators of environmental conditions including hypoxia, and have an outstanding fossil record. Foraminiferans are particularly abundant on the seafloor off major rivers and in other productive areas (Phleger, 1976; Van der Zwaan and Jorissen, 1991). Some species have physiological mechanisms that enable them to withstand severe hypoxic and even anoxic conditions (Moodley et al., 1997; Bernhard and Sen Gupta, 1999;

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Risgaard-Petersen, 2006), although all species disappear when anoxia is prolonged (Alve, 1990, 1991; Schmiedl et al., 2003). The disappearance of foraminiferans may be linked to the toxic effects of hydrogen sulphide. Their absence in some heavily polluted areas may also be caused by heavy metals (Cearreta et al., 2002) or lowered pH (Green et al., 1993; Mojtahid et al., 2008).

Foraminiferans living in hypoxic environments exhibit a number of characteristics. Assemblages are typically of low species richness and dominated by a few hypoxia-tolerant species. Hyaline taxa (rotaliids), particularly chilostomellids, nonionids, uvigerinids, buliminids and bolivinids, usually predominate while agglutinated and porcelanous (miliolid) species are uncommon. The rotaliids often have thin-walled tests with flattened, elongate biserial/triserial, planispiral/lenticular or globular morphologies (Sen Gupta and Machain-Castillo, 1993; Bernhard and Sen Gupta, 1999). Two typical hypoxia-tolerant genera, *Globobulimina* and *Chilostomella*, include deep-infaunal species that live around and immediately below the zero oxygen level in sediments with a well-oxygenated sediment/water interface (Jorissen, 1999a). Species of the *Bolivina/Brizalina/Uvigerina* group have traditionally been regarded as infaunal, in contrast to trochospiral rotaliids such as *Cibicidoides*, which are often considered to be epifaunal (Corliss, 1985; Jorissen, 1999a), although there is no compelling evidence that these two groups occupy distinct microhabitats.

Benthic foraminiferans have been used to trace the development of bottom-water hypoxia in fjords, estuaries, and off major rivers such as the Mississippi and the Po (Sen Gupta et al., 1996; Murray and Alve, 2002). Environmental changes have been inferred from a variety of faunal characteristics. Depressed species richness and diversity, combined with increasing dominance by single species, are often signs of increasing oxygen depletion (Phleger and Soutar, 1973; Gooday et al., 2000; Schmiedl et al., 2003; Brunner et al., 2006). Fluctuations in the abundance of foraminiferal tests in sediment records reflect organic matter flux to the seafloor, as long as sedimentation rates are fairly constant (Herguera and Berger, 1991). Both abundance and diversity decreased dramatically in some parts of Frierfjord, Norway, coincident with the onset of the mod-

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ern industrial period around 1870 (Alve, 2000). These parameters also declined during the 20th Century in cores from the Mississippi Delta Bight (Blackwelder et al., 1996; Sen Gupta et al., 1996; Platon and Sen Gupta, 2001; Platon et al., 2005). In other cases, however, strongly hypoxic areas have extremely high foraminiferal densities, although diversity remains depressed (Phleger and Soutar, 1973; Gooday et al., 2000). Large standing stocks of a few hypoxia-tolerant species may reflect the disappearance of macrofaunal and meiofaunal predators combined with an abundant food supply. On the Pakistan margin of the Arabian Sea, foraminiferans replace the metazoan macrofauna as the principal consumers of organic matter at oxygen concentrations below 5–7  $\mu\text{mol/l}$  (Woulds et al., 2007).

Many studies in coastal environments subject to hypoxia report an increase over time (i.e. upcore) in the proportion of hypoxia-tolerant species (mainly rotaliids) and a corresponding decrease in hypoxia-intolerant species (most miliolids and agglutinated species) (Nelsen et al., 1994). A core from the Northern Adriatic Sea southeast of the Po River delta revealed a gradual change from an assemblage typical of unstressed, oxic environments, including epiphytic habitats, to one that indicated a food-rich, hypoxic environment (Barmawidjaja et al., 1995). The buliminid *Stainforthia fusiformis* is a particularly useful indicator of hypoxia in Norwegian fjords (Alve, 2003; Husum and Alve, 2006). A transition from dominance by *Cassidulina laevigata* in sediments deposited before the establishment of modern industries (pre 1870) to dominance by *S. fusiformis* in the later part of the 20th Century has occurred in Frierfjord (Alve, 2000). In Drammensfjord, fluctuations in the benthic environment over a 1500-year period are indicated by species typical of oxygenated habitats (e.g. *Adercotryma glomeratum*, *Cassidulina laevigata*, *Cribrostomoides kosterensis*, *Elphidium excavatum*, *Nonion labradorensis*) and oxygen-deficient habitats (*Spiroplectammia biformis*, *Stainforthia fusiformis*) (Alve, 1991). Conditions in this fjord have deteriorated since the middle of the 19th Century, culminating in barren, partly laminated sediments deposited during the last 15–20 years. Similar trends were apparent in two cores from the Lower St. Lawrence Estuary, Canada (Thibodeau et al., 2006). Since ~1960, the accumu-

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lation rate of dead benthic foraminiferal tests increased, together with the appearance and progressive increase in abundance of two buliminid species, *Brizalina subaenariensis* and *Bulimina exilis*. These changes were interpreted to indicate a rise in the flux of organic carbon to the seafloor, accompanied by increasingly hypoxic conditions.

5 The enhanced abundance of these species may also reflect the disappearance of less tolerant competitors.

*Ammonia* species (Fig. 2) typically dominate foraminiferal faunas in brackish (mesohaline) and polyhaline coastal and estuarine environments where they can tolerate severe hypoxia and even, for a time, anoxic conditions. Thomas et al. (2000) compared samples collected in the early 1940s and early 1960s with those collected in the late 1990s in the western part of Long Island Sound. They reported an increase in the relative abundance of *Ammonia beccarii*, a decrease in the abundance of *Elphidium excavatum clavatum*, and a decrease in foraminiferal diversity. These changes may reflect seasonal anoxia, eutrophication related to sewage inputs, or other forms of environmental stress. Tsujimoto et al. (2006b) observed a striking increase in the abundance of *Ammonia beccarii* and two agglutinated species, *Trochammina hadai* and *Eggerella advena*, in a core (OBY) from the innermost part of Osaka Bay, Japan, starting at a level in the core corresponding to ~1920. They attributed this trend to increased bottom-water hypoxia related to eutrophication. Since ~1990, however, *A. beccarii* has undergone a sharp decrease in abundance, while *E. advena* and a fourth species, *Uvigerinella glabra*, which was previously rare, have increased. The authors suggest that this latest faunal shift may be related to a change from a dinoflagellate-dominated to a diatom-dominated food supply. Three cores from a transect extending from the OBY site away from the mouth of the Yodo River also exhibited increases in *E. advena*, *A. beccarii* and *T. hadai*, but these were generally more subdued than in core OBY and started later (Tsujimoto et al., 2008). Other species have declined in importance at the two inner sites, starting in the early 1990s in the case of OBY and after the 1940s to 1960s in the case of OS3.

Karlsen et al. (2000) reported fluctuating percentages of *Ammonia* in cores from

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Chesapeake Bay spanning a 500-year time period. They linked a dramatic increase in the late 20th Century, and a corresponding decline in the abundance of *Elphidium selseyensis* and *Ammobaculites*, to environmental changes related to fertilizer use, nutrient loading and oxygen depletion. However, the *Ammonia* record yielded evidence of seasonal anoxia in some years between 1900 and 1960 and episodically in the main channel between 1600 and 1900 (Willard and Cronin, 2007) (Fig. 3), as is also evident in the ostracod record (see below). A similar relationship between *Ammonia* and *Elphidium* on the inner shelf of the Gulf of Mexico (Rabalais et al., 1996) was used by Sen Gupta et al. (1996) to propose the *Ammonia-Elphidium* (A-E) index ( $[N_A/N_A + N_E] \times 100$ , where  $N_A$  is the number of specimens of *Ammonia* and  $N_E$  the number of specimens of *Elphidium* in a sample) as an indicator of hypoxia. The A-E index possibly reflects factors that are linked to oxygen, for example, the relative availability of different food sources, in addition to oxygen itself (Brunner et al., 2006). Whatever the precise controls, this index yields plausible interpretations of historical core records, both in Chesapeake Bay (Sen Gupta and Platon, 2006) and the Gulf of Mexico (Sen Gupta et al., 1996). Historical trends were clearly evident in three cores from the inner shelf (10–30 m) of the Gulf of Mexico where hypoxia is prevalent, but not in a core from a deeper (50 m) site not subject to hypoxia where *Ammonia* and *Elphidium* were less common and the A-E index was highly variable (Sen Gupta et al., 1996; see also Brunner et al., 2006).

Other foraminiferan-based indices have been applied in somewhat deeper water where *Ammonia* and *Elphidium* are uncommon. Together with previous authors (Nelsen et al., 1994; Blackwelder et al., 1996; Platon and Sen Gupta, 2001), Osterman (2003) observed that the combined percentage of three foraminiferal species (*Pseudononion atlanticum*, *Epistominella vitrea* and *Buliminella morgani*) was highest in surface sediment samples from hypoxic areas at mid-shelf depths (30–70 m) off Louisiana. She termed this percentage the PEB Index and used it to reconstruct trends in bottom-water oxygenation over a 180-year period (Osterman et al., 2005). Osterman et al. (2007) extended coverage of the PEB Index to other parts of the Louisiana shelf

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and also examined a much longer time period in three gravity cores, including one obtained within the modern hypoxic zone. In addition to a sharp rise in the PEB Index since the 1950s, this core revealed periodic peaks over the last 1000 years. These probably reflect natural low-oxygen events related to decadal-scale increases in discharges from the Mississippi River (see also Swarzenski et al., 2007). In contrast, no trends in the PEB Index were evident in a core from 60 m water depth NE of the Mississippi Delta (Brunner et al., 2006). This index is probably only applicable locally; not all of the species concerned are widely distributed in hypoxic environments, and one of them, *E. vitrea*, is sometimes associated with well-oxygenated water masses (Pawlowski et al., 2007). Recently, Platon et al. (2005) proposed an “agglutinated-porcelanous index” (*A-P* index) as a possible palaeohypoxia indicator, based on cores from the Mississippi Delta Bight where the *A-P* index declines sharply after the early 1940s as the proportion of hyaline foraminiferans increases (see also Nelsen et al., 1994) (Fig. 4). The *A-E*, PEB and *A-P* indices are useful indicators of historical hypoxia in coastal settings, although they do not correspond to precise bottom-water oxygen values.

Possible links have been suggested between foraminiferal test morphology and bottom-water oxygenation. In a core from Yokohama Bay spanning a period of about 100 years, Toyoda and Kitazato (1996) recognised morphotypes of three dominant species (*Trochammina hadai*, *Ammonia beccarii* and *A. japonica*) with lobate and smooth outlines. Lobate tests were considered to be more typical of oxygenated conditions whereas the smooth forms were associated with oxygen depletion. Jorissen (1988) attributed similar morphotype variations in *A. parkinsonia/tepida*, *Elphidium granosum* and *E. poeyanum* from the Northern Adriatic Sea to the influence of higher organic input (leading to an accelerated growth and thinner, more lobate chambers), in addition to the effects of lower oxygen concentrations. Note that in contrast to the Yokohama Bay study, lobate chambers were associated with hypoxic conditions. Also in the northern Adriatic Sea, *Bulimina marginata* exhibited a correlation between an inferred oxygen gradient and test morphology, with morphotypes lacking spinose or-

namentation being present in the areas with highest organic input (and lowest oxygen concentrations) (Jorissen, 1988). Finally, it has been suggested that deformed *Ammonia* tests in Chesapeake Bay are associated with hypoxia (Karlsen et al., 2000).

Stable carbon isotope ratios in calcareous foraminiferal tests provide an additional tool for reconstructing historical environments, in particular organic carbon delivery to sediments. Increased organic matter inputs typically lead to more depleted carbon isotope signatures of pore water dissolved inorganic pools because respiration of organic matter adds carbon dioxide that is depleted in  $^{13}\text{C}$ . Accordingly, benthic foraminiferans extracting carbonate from this  $^{13}\text{C}$  depleted pore-water pool will be characterised by isotopically lighter  $\delta^{13}\text{C}$  values. In two cores from the Mississippi Delta Bight,  $\delta^{13}\text{C}$  values obtained from multispecies benthic foraminiferal samples, and from *Buliminella morgani*, were about 0.6–1.0% heavier before the mid-1960s compared to those derived from later sediments (Eadie et al., 1994). A similar trend was reported by Thomas et al. (2000) in Long Island Sound where calcareous tests collected in 1961 were isotopically slightly heavier than those collected in 1996/1997. The authors inferred from these results that the amount of oxidizable organic carbon had increased over this 35 year time period. They suggested that the magnitude of the shift could be used as a proxy for the occurrence and severity of past anoxic episodes in Long Island Sound.

Foraminiferans respond to a complex set of environmental parameters. In coastal environments, these include the quantity, quality and regularity of food inputs, salinity, current and wave activity, as well as oxygen (Murray, 2001, 2006). Since many foraminiferans can tolerate low-oxygen conditions, food inputs may be more important than oxygen stress. Nevertheless, because of the apparent robustness of the correlation between organic input and bottom-water oxygen conditions, at least on a local to regional scale, the foraminiferal response may indeed yield valuable information about past oxygen levels.

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### 3.1.2 Benthic Ostracoda

Ostracods are crustaceans that live in the upper 1 cm of freshwater and marine sediments where they form part of the metazoan meiofauna. Their calcified bivalved shells are commonly preserved as fossils, usually as disarticulated valves. Like foraminiferans, they are sensitive indicators of environmental conditions, including changes caused by human impacts (Ruiz et al., 2005). They respond to salinity and bottom-water oxygenation with assemblages in stressed environments often being dominated by single species (Boomer, 2002). Unlike foraminiferans, and in common with most crustacean taxa (e.g. Moodley et al., 1997), they usually are intolerant of hypoxia, although a few marine and brackish-water species are exceptions (Modig and Ólafsson, 1998; Cronin and Vann, 2003; Whatley et al., 2003; Ruiz et al., 2005; Corbari et al., 2005). *Cyprideis torosa*, one of the most abundant metazoans in coastal Baltic waters, survives in sulphidic habitats by oxidizing sulphide to thiosulphate (Jahn et al., 1996). Under hypoxia, this species can use anaerobic metabolic pathways to detoxify sulphide, allowing it to survive these extreme conditions for a considerable time. Two other Baltic species, *Heterocyprideis sorbyana* and *Candona neglecta*, may have similar capabilities (Modig and Ólafsson, 1998).

Ostracods are considerably less common than foraminiferans in sediment cores and usually play a supporting role in historical reconstructions (e.g. Alve, 1991). A number of studies, however, focus specifically on ostracods as tracers of anthropogenic environmental change in coastal environments. Alvarez Zarikian et al. (2000) analysed assemblages along a 50-cm-long core from a severely hypoxic area on the inner shelf off Louisiana, spanning the period from 1904 to 1993. One genus, *Loxoconcha*, dominated modern ostracod assemblages associated with severely hypoxic conditions at the top of the core and decreased in relative abundance downcore. The relative abundance of this species corresponded closely to the record of fertilizer application in the United States, a proxy for nutrients discharged by the Mississippi River, and suggested that the coring site became hypoxic [ $O_2=0.1-1.0\text{ mL L}^{-1}$  ( $=4.3-$

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43  $\mu\text{M}$ )] in the mid-1940s. The overall species diversity of ostracods in this core also declined in more recent sediments (Rabalais et al., 2002) (Fig. 5).

An important study by Cronin and Vann (2003) applied information on modern ostracod assemblages in Chesapeake Bay to reconstruct trends in salinity, dissolved oxygen and turbidity over a 500-year period based on a core obtained from the bay at the mouth of the Pautuxet River. One particular species, *Cytheromorpha curta*, was able to tolerate periodic anoxia as well as high turbidity and a wide range of salinities, whereas most other species required oxygenated water but tolerated different salinity and turbidity levels. Shifts in species abundances suggested natural environmental variability, including periods of hypoxia, related to climatic fluctuations during the 16th and 17th Centuries (Fig. 6a). Human impacts become apparent during the past 200 years. A striking feature of this record is the dramatic increase in the abundance of *C. curta* since the 1960s, coupled with the virtual disappearance of other ostracod species (Fig. 6b). Cronin and Vann (2003) consider the increased intensity of seasonal anoxia to be the most likely explanation for this recent faunal shift.

Yasuhara et al. (2007) described changes in ostracod abundance, diversity and faunal composition in four cores taken along a transect from the innermost part of Osaka Bay (Japan) near the mouth of the Yodo River, to the centre of the Bay. The total abundance of ostracods decreased from 1910–1920 at the innermost station (OBY) and from ~1960 at the next (inner) station (OS3) along the transect. Since ~1970, abundance has been very low (<20 specimens per 20 g dry sediment) at these two sites. Although the absolute abundances of all species decreased, the relative abundances of *Bicornucythere bisanensis*, *B. sp.*, *Cytheromorpha acupunctata* and *Loxoconcha viva* increased. These trends were attributed to eutrophication, which accelerated during the 1950s and peaked between 1960 and 1970, combined with summer hypoxia, which reached a maximum in the 1970s. Hypoxia only affected the inner part of the bay. At the outer two stations, eutrophication in the absence of hypoxia led to an increase in ostracod abundance. Rather surprisingly, these 100–200-yr sediment records revealed no major changes in ostracod diversity at any of the four stations in Osaka Bay.

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Yasuhara et al. (2003) reported similar patterns in Hiroshima Bay. Ostracod abundances decreased from the 1940s in the inner bay while the relative abundance of *B. bisanensis* increased and diversity remained relatively stable over the 100-yr record.

Direct comparisons of the responses of ostracods (metazoans) and foraminiferans (protists) to environmental impacts are rare. An illustration in Rabalais et al. (2007), reproduced here as Fig. 5, shows a similar decreasing trend in ostracod and foraminiferal diversity (Shannon-Wiener index) from 1900 to 1990 in the northern Gulf of Mexico. Over shorter time scales, oscillations in the diversity of the two taxa show some correspondence, although the match is not precise. As mentioned above, ostracod abundance in Osaka Bay began to decline at the innermost, hypoxia-influenced station (OBY) around 1910–1920 while diversity remained stable (Yasuhara et al., 2007). In the same core, foraminiferans increased in abundance from ~1920 to ~1965 before undergoing a relatively modest decrease (Tsujiimoto et al., 2006b), while diversity decreased substantially after ~1920. This suggests foraminiferans and ostracods exhibit different responses to hypoxia, with certain foraminiferal species being more tolerant than the ostracods.

### 3.1.3 Juvenile molluscs

Alve (2000) noted abundance peaks in juvenile gastropods and bivalves in the upper sections (dating from ~1870 onwards) of cores from >70 m water depth in Frierfjord, Norway. She attributed these peaks, which were accompanied by a strong dominance of the hypoxia-tolerant foraminiferan *Stainforthia fusiformis*, to the lethal affects of anoxia on the juvenile molluscs. Similar patterns were observed in Drammensfjord (Alve, 1991). Why elevated densities of recruits should occur under these conditions is not clear. Possibly, they reflect brief periods of re-oxygenation leading to a transient settlement event, as reported for the opportunistic bivalve *Abra aequalis* on the Texas continental shelf (Harper et al., 1991).

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## 3.2 Sediment laminations and trace fossils

A variety of mechanisms can generate laminations (varves) but all require that the laminated sediment fabric is not disrupted by bioturbation (Kemp et al., 1996). In marine environments, this usually happens when the macroinfauna are eliminated by severe hypoxia, as occurs in the core regions of some open-ocean oxygen minimum zones (OMZs) (van Geen et al., 2003; Levin et al., 2009), silled basins with restricted circulation (Thunnell et al., 1995), and in some fjords (Alve, 2000). Laminations are also developed off major rivers where sediment deposition rates are so high that macrofaunal populations cannot be maintained (Rabalais, unpublished observations). Modern laminated sediments are not always devoid of eukaryotic life; some harbour large numbers of foraminiferans and nematodes that cause bioturbation on a microscopic scale not apparent to the naked eye (Pike et al., 2001). Recent studies suggest that degraded laminations may persist where macroinfauna begin to reappear in the lower parts of OMZs (Levin et al., 2009).

Sequences of laminated and bioturbated sediments have been used to reconstruct Quaternary climatic fluctuations, notably along the western margin of North America (Behl, 1995; Behl and Kennett, 1996). Laminations have also been used to infer hypoxic conditions in more recent historical records. A shift from laminated to bioturbated sediments since the 1970s has been interpreted, in combination with changing foraminiferal  $\delta^{13}\text{C}$  values, as signifying reduced carbon oxidation rates and a 15–20  $\mu\text{M}$  increase in dissolved oxygen concentrations at the edges of several Southern California borderland basins. These changes correspond to a 20–30% reduction in upwelling and a 1.5–3°C increase in temperature and suggest climatic influences on productivity (Stott et al., 2000). In parts of Drammensfjord (Norway), pyrite-containing laminations indicate that hypoxic conditions have persisted throughout the last 1000 yrs (Alve, 1990, 1991; Smittenberg et al., 2005). Laminations are developed in other Norwegian fjords, e.g. Kyllaren Fjord (Smittenberg et al., 2004), and have become widespread in the Baltic Sea since the 1950s (Persson and Jonsson, 2000; Leipe et al., 2008) (Fig. 7).

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Cores from the central channel in the middle part of Chesapeake Bay are dominated by laminations, suggesting that macro-infauna have been absent for ~100 yrs (Schaffner et al., 1992).

Palaeoecologists have devised models for estimating palaeo-oxygenation from the degree of lamination and occurrence of trace fossils (Savrda and Bottjer, 1991). In modern sediments it is recognized that the abundance, body size and diversity of metazoan animals declines at very low oxygen levels (Rhoads and Morse, 1971; Levin, 2003). Thus it is reasonable to assume that the density, size and diversity of animal traces (ichnofacies) in sediments should also decline with increasing hypoxia. A number of ichnofacies models have been constructed to relate animal activities and trace morphologies to palaeo-oxygenation (Savrda and Bottjer, 1984, 1987, 1991), based largely on observations from modern hypoxic basins. Reduced density, size, depth of penetration and diversity of macrofaunal traces are documented in oxygen-deficient basins (Rhodes and Morse, 1971; Savrda et al., 1984), fjords (e.g., Nilsson and Rosenberg, 2000) and oxygen minimum zones (Levin, 2003).

Trends in body size and activity, however, are not always linearly correlated with oxygenation. The occurrence of chemosymbiotic bacterial symbionts within metazoan organisms may pre-adapt them to survive in dysoxic sediments while maintaining large body size or significant burrowing activity. The description of the exaerobic zone includes a shelly fauna comprising large individuals, possibly bivalves or brachiopods (Savrda and Bottjer, 1987). Tubificid oligochaetes in the subfamily Phalodrilidae appear to survive and bioturbate sediments at oxygen levels of  $0.02 \text{ mL L}^{-1}$  ( $=0.8 \mu\text{M}$ ) (Levin, 2003), possibly with support from their symbiotic bacteria which have multiple metabolic functions (Blazjek et al., 2005). These occurrences complicate the interpretation of laminations as they often lead to mixing of sediments under oxygen conditions thought to inhibit bioturbators. Another consideration is that in regions where hypoxia persists over evolutionary time scales, organic matter availability rather than oxygen appears to exert primary control over the development of bioturbating infaunal communities and thus the persistence of laminations. Variations in laminae formation across

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the Pakistan margin OMZ, and greater formation of laminae on the Pakistan than Oman margins, may be explained better by differences in POC flux than by oxygen levels (Gooday et al., 2009; Levin et al., 2009; Hughes et al., 2009).

### 3.3 Chemical and mineral indicators

#### 3.3.1 Sulphur and sulphides

High concentrations of sulphur and sulphide-forming metals in hypoxic and anoxic sediments are the basis of a number of proxies for bottom-water oxygenation (Berner, 1984). In the absence of oxygen, labile organic matter is degraded by prokaryotes utilizing alternative electron acceptors such as nitrate, metal oxides and sulphate, of which the latter is by far the most abundant. Sulphate reduction results in the formation of hydrogen sulphide which either reacts with reactive iron minerals or is oxidized chemically and particularly biologically by a diverse community of sulphide oxidizing bacteria. Oxygen limitation restricts the re-oxidation of hydrogen sulphide in sediments underlying hypoxic and anoxic bottom-waters and iron sulphide mineral formation therefore represents the main sink of hydrogen sulphide (Berner, 1994; Middelburg, 1991). Pyrite is normally the dominant iron sulphide mineral formed, but this usually involves acid volatile precursors (AVS, acid volatile sulphides) such as greigite.

One proxy for bottom-water oxygenation is based on the presence or absence of a correlation between organic carbon and reduced sulphur in sediments. In normal marine sediments underlying oxic bottom waters, organic carbon and sulphur are well correlated because a certain carbon delivery relates to a proportional preservation of organic carbon and reduced sulphur (Berner, 1984). In sediments underlying sulphide-rich waters there is also iron sulphide formation occurring in the water column with the result that sediment organic carbon and sulphur contents are unrelated (Berner, 1984; Middelburg, 1991; Passier et al., 1996). The preservation of sulphur has increased in sediments deposited in Chesapeake Bay during recent centuries, together with total organic carbon and total (mainly organic) nitrogen (Cooper and Brush, 1991, 1993)

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(Fig. 8c). These trends reflect increasing eutrophication and hypoxia and are linked to an increase in the formation of pyrite.

Mineral texture is the basis for another pyrite-related proxy (Wilkins et al., 1996; Passier et al., 1997). The formation of pyrite framboids (aggregations of small grains) apparently requires the presence of limited amounts of oxygen. Their size appears to depend on whether they were formed near the oxic-anoxic interface in the water column and subsequently settled to the seafloor (smaller framboids,  $\leq 5 \mu\text{m}$  diameter) or at the oxic-anoxic boundary in the sediment (larger framboids) (Brunner et al., 2006). Moreover, pyrite formed slowly in fully anoxic sediments may be euhedral (Passier et al., 1997). At a coring site northeast of the Mississippi River Delta, the size of the framboids suggested that they were formed mainly in the sediment and that the bottom water had been oxygenated throughout the 20th Century (Brunner et al., 2006).

Another measure, the ratio between acid volatile sulphur (AVS) and chromium reducible sulphur (CRS), essentially the ratio between sulphur as FeS and sulphur as pyrite, is based on the concept that the conversion of the precursor AVS to CRS (pyrite) requires an oxidant and time (Middelburg, 1991). In a core from Chesapeake Bay, the AVS:CRS ratio increased sharply around 1934, was stable between 1934 and 1984, and increased again thereafter (Zimmermann and Canuel, 2000, 2002). This trend was coincident with that of biogenic silica, an indicator of planktonic diatom production (see below), and was interpreted as indicating a worsening of anoxic episodes during recent decades. The ratio of AVS to CRS is useful but may give biased information if sediment accumulation rates fluctuate.

Probably the most reliable proxy for historical bottom-water oxygenation is the degree of pyritization (DOP=the ratio of pyritic Fe to total reactive Fe) or its equivalent the degree of sulfidation (DOS=the ratio of pyritic Fe and AVS-Fe to total reactive Fe). This has been used to distinguish between oxic and hypoxic environments in ancient sediments (Raiswell et al., 1988) and is applicable to modern sediments as well (Middelburg et al., 1991; Wijsman et al., 2001a). Cooper and Brush (1991) used this method to trace environmental changes in Chesapeake Bay (Fig. 8d). The degree of

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pyritization showed a good correspondence with evidence for eutrophication derived from the diversity and composition of diatom assemblages (Cooper and Brush, 1993).

Finally, sulphur stable isotope signatures also have potential as indicators of hypoxic conditions. During sulphate reduction there is significant isotope discrimination so that the hydrogen sulphide formed is strongly depleted in  $^{34}\text{S}$  relative to sulphate. In sediments underlying oxic bottom-waters, a major fraction of the sulphide generated is re-oxidized in multiple steps to sulphate. During this re-oxidation, in particular during the disproportionation reactions, further isotope fractionation takes place. As a consequence, sulphide isotope fractionation increases if sulphate reduction is followed by disproportionation reactions in the presence of oxidants (Canfield and Thamdrup, 1994). Distinct increases in  $\delta^{34}\text{S}$  values in cores from Charlotte Harbor (Florida) seem to reflect an intensification of reducing conditions due to lower bottom-water oxygenation and increased organic inputs to the sediment (Turner et al., 2006). Wijsman et al. (2001b) studied sediments in a zone at the shelf-edge of the Black Sea where oxygen conditions changed with time. They observed high variability in  $^{34}\text{S}$  values of pyrite in sediments and attributed this to fluctuations in the oxygen concentrations of bottom waters or varying sediment accumulation. During oxic periods, sulphide re-oxidation processes became important resulting in low AVS to CRS ratios and light  $\delta^{34}\text{S}$  values. Anoxic conditions in the bottom waters overlying shelf-edge sediments were reflected in enhanced AVS to CRS ratios and heavier sulphur isotope values.

### 3.3.2 Glauconite

The formation in sediments of the mineral glauconite, a hydrous silicate of potassium, iron, aluminum, or magnesium, is believed to be accelerated under low-oxygen or anoxic conditions in shelf sediments (Br  h  ret, 1991; Nelsen et al., 1994). The percentage of glauconite grains in the  $>63\ \mu\text{m}$  fraction of a core from a hypoxia-influenced site in the Gulf of Mexico increased during the late 1930s and early 1940s. The proportion of hypoxia-tolerant foraminiferans (Nelsen et al., 1994) increased at the same time, suggesting that glauconite may be an indicator of anthropogenically-forced hypoxia.

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### 3.3.3 Magnetic properties of sediments

Magnetic minerals provide a powerful tool to determine the dissolved oxygen status of marine sediments. In fully oxic environments, the magnetic characteristics of the mineral assemblage are mostly derived from coarse detrital particles, such as hematite and magnetite, with a very limited contribution from authigenic and fine-grained ferromagnetic minerals. Under hypoxic and/or anoxic conditions, however, remagnetization takes place. This results from the precipitation of biogenic or authigenic fine-grained magnetic minerals, e.g. greigite and pyrrhotite (precursors of pyrite), which form as byproducts of microbial metabolism (Robinson and Sahota, 2000; Liu et al., 2004; Larrasoña et al., 2007). Sulphide minerals such as greigite could indicate the presence of oxygen-depleted bottom water during their formation (see above). However, these magnetic sulphide minerals are usually ephemeral and thus cannot provide reliable historical records. Moreover, they sometimes develop as a result of late diagenetic processes that occur some metres below the sediment/water interface, and therefore do not reflect the environment in which the sediments were deposited (Lui et al., 2004; Roberts and Weaver, 2005).

Under hypoxic conditions, Fe and Mn oxides, including hematite and magnetite, may be used for respiration by anaerobic bacteria or be consumed by reactions with sulphides. Moreover, new magnetic minerals may form as a result of Fe and Mn cycling. Consequently, the ratio between different magnetic properties, such as mass-specific magnetic susceptibility ( $\chi$ ), anhysteretic remnant magnetization (ARM) and saturation isothermal remnant magnetization (SIRM), show distinct contrasts between oxic and anoxic sediments (Robinson and Sahota, 2000). These properties, together with enrichment in some redox-sensitive elements (see below), have been proposed as indicators of poor bottom-water ventilation (i.e. hypoxic conditions) during the formation of Mediterranean sapropels (e.g. Larrasoña et al., 2003). Rhoads et al. (1991) provide evidence from TEM micrographs of sediments on the Peru shelf that the inventory of biogenic magnetite is associated primarily with dysaerobic oxygen conditions [0.1–

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0.5 mL L<sup>-1</sup> (=4.3–21.5 μM)] and is 3–5 times lower in aerobic and anaerobic sediments. As far as we are aware, these approaches have not been applied to reconstruction of oxygenation regimes from historical records in coastal settings.

### 3.3.4 Trace elements

5 Trace elements that are enriched in sediments deposited under hypoxic or anoxic conditions are widely used in palaeoceanography to infer the history of bottom-water oxygenation (Bruland and Lohan, 2003; Calvert and Pedersen, 2007; Tribovillard et al., 2006). The underlying premise is that total trace element concentration can be partitioned into a detrital contribution (often estimated via Al or by comparison with average  
10 shale) and an excess contribution due to hydrothermal, biogenic and authigenic processes. Trace element enrichment of non-hydrothermal sediments can be attributed to either delivery of organic matter and associated biogenic phases (Ba, Cd, Ni, Zn, Cu) or to diagenetic enrichment due to the redox chemistry of the element involved (e.g. Re, Mo, V, U). Trace element such as Ba, Cd and Zn show nutrient type behaviour  
15 in the water column, are delivered to sediments with organic matter and can thus be used as a proxy for eutrophication, provided their signal is preserved (Tribovillard et al., 2006). Excess Ba is mainly found in barite (barium sulphate), the preservation potential of which may be low in anoxic sediments with sulphate reduction. Ni and Cu are well preserved in anoxic sediments because they form sulphides and can thus be used  
20 as proxies for organic matter abundance, but no applications in the historical record are known.

Trace element such as vanadium, uranium, molybdenum and rhenium have their own redox chemistry and if their reduced form is less soluble than the oxidized form they become enriched in hypoxic and anoxic sediments (Middelburg et al., 1991; Tribovillard  
25 et al., 2006). Vanadium, uranium and rhenium are reduced under less reducing conditions (i.e. suboxic) than molybdenum that requires sulphidic conditions. This differential behaviour provides opportunities to distinguish between suboxic/anoxic versus euxinic

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environments.

Molybdenum is one of the most widely applied trace metals in palaeo-environmental reconstructions. Off Cape Blanc on the northwestern African coast, the Mo/Al record showed a perfect correlation with the cumulative percentage of deep-infaunal foraminiferal species, considered to be low-oxygen indicators, over the last 70 000 years (Martinez et al., 1999). The sedimentary accumulation of Mo is proportional to sulphide concentrations (i.e., free H<sub>2</sub>S). It is regarded as a proxy for water-column anoxia and is not enriched in suboxic sediments that are devoid of sulphide (Crusius et al., 1996). However, the mechanisms by which Mo becomes concentrated in sediments are not fully understood. Under oxic conditions, it is present in the form of MoO<sub>4</sub><sup>-</sup> ion, and can be adsorbed to particulate manganese oxides formed under oxic conditions. These manganese oxides tend to be formed at the oxic/anoxic interface in the water, and can sink into the anoxic zone, carrying molybdenum with them. Once in the suboxic or anoxic zone, the manganese oxides will dissolve, releasing Mo to the water column or porewater (depending on water depth and location of the oxic/anoxic interface). Subsequent fixation in the sediment may involve the formation of organic thiomolybdenates (sulphidic compounds) and inorganic Fe-Mo-S cluster complexes (Adelson et al., 2001). Thus Mo enrichment tends to be associated with the presence of an anoxic water column.

Molybdenum and vanadium have been used in historical reconstructions, albeit rarely (Table 1). Adelson et al. (2001) showed that Mo levels in two cores (55 and PC6) from the upper reaches of the main stem of Chesapeake Bay exhibited considerable variability but were highest in sediments deposited after 1960 (Fig. 9). This was interpreted to indicate increasingly intense seasonal oxygen depletion, consistent with evidence from other sources for rising eutrophication in the mesohaline part of the Bay. This trend was not observed by Zheng et al. (2003) in two cores from an area ~25 km south of PC6. Instead, they found that Mo levels exhibited variations that were inversely correlated with proxies for palaeosalinity during the 20th Century. They suggested that anoxia had fluctuated naturally as a result of climatically-driven varia-

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tions in freshwater discharge over time. In a core from the Louisiana shelf spanning a 1000-year period, V was depleted at levels where peaks in the foraminiferal PEB index indicated hypoxia (Swarzenski et al., 2008). These pre-anthropogenic hypoxic events were probably related to increased discharge from the Mississippi River, caused by climatic fluctuations.

On the western Indian continental shelf, the hypoxia that is formed naturally during late summer and autumn may have intensified in recent years (Naqvi et al., 2000, 2006a) due to increased inputs of human-generated nutrients. Here, the Mo/Al mass ratio has been found to vary over a narrow range ( $8.5\text{--}11\times 10^{-4}$ ) during the last 250 years, as revealed by the analyses of two cores taken from the inner- and mid-shelf regions off Goa (D'Souza, 2007). The ratio, however, is much higher than the average crustal value of  $0.18\times 10^{-4}$  (Taylor, 1964). This implies prevalence of anoxic conditions in sediments throughout the depositional period. In two recent studies on the Indian shelf off Goa, based on three sediment cores from the zone of present seasonal anoxia and covering the last ~200, 250 and 700 years, the V/Al ratio varied between 1 and  $14\times 10^{-4}$  (D'Souza, 2007; Agnihotri et al., 2009). While this range is less than the average crustal value ( $16.4\times 10^{-4}$ ; Taylor, 1964), the highest ratios were generally recorded in the upper portions of the cores (i.e. the last four decades), corresponding to a period of record high productivity. An increase in the Cu/Al ratio was also recorded during the same period (Agnihotri et al., 2009).

The trace element chemistry of sediments of Kau Bay (Indonesia), a 470-m deep hypoxic to euxinic basin separated from the Pacific by a shallow sill, has been studied in detail (Middelburg et al., 1989, 1991). These organic-rich sediments underlying hypoxic bottom-waters are enriched in V and in particular Mo. The Mo and V enrichments showed some well-developed maxima reflecting periods of stagnant bottom water conditions (Middelburg et al., 1989) (Fig. 10). These maxima suggested that the frequency of bottom-water ventilation in Kau Bay was related to ENSO events.

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## 4 Water-column derived indicators of eutrophication and hypoxia

### 4.1 Planktonic indicators of eutrophication and hypoxia

The main indicators of eutrophication are diatoms and dinoflagellate cysts. Their remains, and chemical markers derived from them, can be used as proxies for changes in phytoplankton biomass and community composition and hence primary productivity (Rabalais et al., 2004, 2007). Because hypoxia is often associated with eutrophication, we consider these indicators in some detail. The preservation of diatoms and dinoflagellate cysts is also enhanced in oxygen-depleted sediments. Possible multiproxy approaches to distinguishing between enhanced productivity and enhanced preservation are discussed below (Sect. 5.2).

#### 4.1.1 Diatoms

Diatoms are predominantly planktonic, photosynthetic protists in which the cell is encased by a siliceous frustule composed of two valves. They are found in all aquatic environments within the photic zone and generally thrive under eutrophic conditions (Round et al., 1990). Increases in diatom abundance are often an early sign of eutrophication (Smith, 2003). Crosta and Koç (2007) review the use of diatoms as palaeo-indicators. Changes in species composition reflect, among other factors, salinity, sedimentation, turbidity and nutrient availability (Cooper 1995; Rabalais et al., 1996; Parsons et al., 1999, 2006). In the Northern Gulf of Mexico, there was a decline in the dominance of heavily silicified diatoms and a corresponding increase in lightly silicified species during the second half of the 20th Century (Rabalais et al., 1996). The rising absolute and relative abundance since the 1950s of *Pseudo-nitzschia* (Dortch et al., 1997), a lightly silicified diatom that does not preserve well in sediments, has been linked to increased eutrophication combined with silica limitation (Parsons et al., 2002). Although diatom preservation may be enhanced in anoxic sediment with reduced densities of bioturbating animals (McMinn, 1995), this factor does not seem to

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be responsible for patterns in the diatom record for the northern Gulf of Mexico (Parsons et al., 2002).

In four cores from a part of Chesapeake Bay where anoxic periods are most prevalent and severe, Cooper and Brush (1993) and Cooper (1995) detected a decrease in diatom diversity beginning around 1760 that became particularly marked in recent sediments. This resulted from a decline in the ratio of benthic or epiphytic (pennate) species to planktonic (centric) species (Fig. 8a). They attributed these trends to a number of possible factors, including eutrophication and increasing anoxia. Similar changes in diatom assemblages, also linked to eutrophication, are reported from the southern Baltic Sea (Andrén, 1999; Andrén et al., 1999) and Laajalahti Bay, Finland (Kauppila et al., 2005).

Qualitative analyses of diatom assemblages cannot be used to reconstruct actual nutrient concentrations, but these can be inferred from transfer functions based on percentage data for modern species (Clarke et al., 2003; Crosta and Koç, 2007). This approach has been used in lakes for some decades (Hall and Smol, 1999) and has been applied recently to reconstruct total nitrogen (TN) concentrations in three marine settings, Roskilde Fjord (Denmark), Laajalahti Bay (Finland) (Clarke et al., 2006) and Mariager Fjord (Denmark) (Ellegaard et al., 2006). At the Finnish site, TN values inferred from diatom data track actual monitored values well, except during the most eutrophic periods in the early 1970s when they consistently underestimated total dissolved nitrogen. In Mariager Fjord, the results suggested a progressive increase in TN since 1900, with reconstructed values being in good agreement with actual TN values between 1985 and 2000.

Biologically bound silica (BSi) is a useful proxy for diatom production (Turner and Rabalais, 1994). In the Mississippi River Delta Bight and adjacent parts of the continental slope, BSi levels have increased since ~1980, particularly at depths between 27 and 50 m where summer hypoxia is most frequent (Turner and Rabalais, 1994; Rabalais et al., 1996; Turner et al., 2004) (Fig. 11). The increased seafloor organic loading associated with these diatom fluxes is believed to be responsible for an increase in

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the persistence and extent of bottom-water hypoxia. A similar trend is evident in cores from Charlotte Harbor estuary (Florida) (Turner et al., 2006). In Chesapeake Bay, trends in the concentration of BSi in the upper core sections are fairly modest and inconsistent (Cooper, 1995; Cornwell et al., 1996; Colman and Bratton, 2003), although Zimmermann and Canuel (2000) show that BSi values increase substantially across a sediment layer corresponding to 1934 and continue to rise thereafter to the top of the core. When BSi values are converted to fluxes (BSi concentrations x total sediment flux), much clearer trends emerge, with a 4–5-fold increase beginning 200–250 years ago (Colman and Bratton, 2003). A sharp increase in BSi flux after ~1770 is likewise evident in a core from New Bedford Harbor, Massachusetts (Chmura et al., 2004).

Several factors complicate the relationship between BSi and diatom production (Colman and Bratton, 2003), particularly in turbid coastal environments (e.g. off large river estuaries) where biogenic silica from terrestrial sources such as phytoliths can be an important source for BSi. The determination of BSi in marine sediment is also subject to considerable controversy and is affected, for example, by the leaching of non-biological silica from clay minerals (Lui et al., 2002). Correction using the Al/Si ratio and/or other chemical leaching methods may lead to either an overestimate or underestimate of BSi values.

#### 4.1.2 Dinoflagellate cysts

Dinoflagellates are autotrophic, heterotrophic, or mixotrophic protists found in marine, brackish and freshwater settings. About 10–20% of the 2000 known species produce acid-resistant resting cysts, typically 15–100  $\mu\text{m}$  diameter, which sink to the seafloor and can remain viable in bottom sediments for many years (Dale, 1996; de Vernal and Marret, 2007). These “dinocysts” are preserved in ancient sediments, mainly those deposited in coastal waters (Dale and Dale, 2002). Unlike calcareous microfossils, dinocysts are organic in composition and therefore resistant to dissolution. Being derived from planktonic organisms, they reflect surface-water conditions, including temperature, salinity and nutrients, rather than those on the seafloor (Dale, 1996).

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However, some species, many of them heterotrophic, are characteristic of eutrophic conditions, both natural and anthropogenic, and thus may be associated with seafloor hypoxia (Willard et al., 2003). Thorsen and Dale (1997) suggested that an increased proportion of heterotrophic cysts may be used as an indicator of eutrophication. Reconstructions of surface productivity based on dinocyst assemblages are presently qualitative. Quantitative estimates may be possible in the future (de Vernal and Marret, 2007), although the development of transfer functions is fraught with problems (Dale and Dale, 2002).

Dinoflagellate cysts have been used as tracers of environmental change in Holocene and Quaternary sequences (de Vernal and Marret, 2007), including recent historical periods (Dale and Dale, 2002). Dale et al. (1999) and Dale (2000) described changes in dinoflagellate assemblages in the inner part of Oslofjord over a >500 yr interval. The record clearly reflected the onset and development of eutrophication during the late 19th and early 20th Century, its peak in the 1940s and 1950s, and recovery of the system following remediation efforts in the 1970s and 1980s. One particular species, *Lingulodinium machaerophorum* (the cyst of the autotrophic species *Gonyaulax polyedra*), was strongly associated with eutrophic conditions and led to an overall two-fold increase in cyst abundance. In a similar study in Frierfjord, Dale (2000) recognised a number of phases over a 300-year core record that could be interpreted either in terms of pollution or eutrophication, although the signal was less clear than in the Oslofjord. Based on dinocysts, Sangiorgi and Donders (2004) distinguished between the lower part of a core from the Northern Adriatic Sea, representing the period between 1830 and 1924, and the upper part (1930–1989) during which there was a progressive increase in the heterotrophic/autotrophic ratio as well as the abundance of *L. machaerophorum*. Since ~1930, there has been an increase in eutrophication together with incidences of severe bottom-water hypoxia in the Northern Adriatic Sea (Justić et al., 1987; Justić, 1991). Changes in dinocyst assemblages in Mariager Fjord (Denmark) during the last century are similarly consistent with increased eutrophication (Ellegaard et al., 2006).

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Dinocysts have also been used to trace human impacts in Japanese and North American estuaries. Although there was no overall increase in cyst abundance, a core from Yokohama Port, Japan, revealed a clear shift around the 1970 level from an assemblage dominated by autotrophic dinoflagellate cysts to one dominated by the cysts of heterotrophs (Matsuoka, 1999). This coincided with the beginning of strong economic growth in Japan and probably reflected the onset of eutrophication. The first North American study was carried out by Pospelova et al. (2002) in New Bedford Harbor and Apponagansett Bay, Massachusetts, both shallow, unstratified estuaries not subject to severe hypoxia. Here, there was also no overall increase in the concentration of cysts during the last 500 years but there was much greater variability in their abundance during the 20th Century compared to earlier periods. At the same time, a sharp decrease in species richness was observed in New Bedford Harbor, the more strongly polluted locality. These trends are believed to reflect increased environmental stress arising from eutrophication and heavy metal inputs. A later comparison of dinocyst assemblages with two other palaeoproductivity proxies, pigments and BSi, revealed a succession of changes in the New Bedford Harbor ecosystem over the past few hundred years linked to forest clearance, sewage discharges, and harbour modifications (Chmura et al., 2004). Unlike Dale et al. (1999) and Sangiorgi and Donders (2004), both Matsuoka (1999) and Pospelova et al. (2002) recorded decreases in the abundance of *Lingulodinium machaerophorum* in response to eutrophication. Pospelova et al. (2002) suggest that this autotrophic species may increase as a response to oxygen depletion rather than to eutrophication. Thibodeau et al. (2006) report two- to three-fold increases in the concentration and accumulation rates of dinocysts in the upper parts of two cores from the St. Lawrence Estuary. These were probably caused by increased primary production, and possibly eutrophication, since about 1960.

Dinocysts may reflect water column oxygen concentrations as well as surface primary productivity. Willard et al. (2003) used pollen and dinocysts as proxies for regional precipitation, estuarine salinity and dissolved oxygen in Chesapeake Bay during the past 2500 years. Based on quantitative analyses of dinocyst assemblages, they iden-

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tified a period between 1400 and 1500 AD characterised by a major change in species composition (Fig. 12). They interpreted this as indicating oxygen depletion in the deep channel of Chesapeake Bay and estimated minimum annual mean concentrations at  $0.5\text{--}1.0\text{ mg L}^{-1}$  [ $0.36\text{--}0.72\text{ mL L}^{-1}$  ( $=15.6\text{--}31.2\text{ }\mu\text{M}$ )]. An important consideration in the interpretation of dinocyst records is the possibility of enhanced preservation in anoxic sediments. For example, heterotrophic species of *Protoperidinium*, which are strongly associated with high productivity, are easily degraded in the presence of oxygen but occur in large numbers in sediments deposited under anoxic or hypoxic conditions (Zonneveld et al., 1997; Versteegh and Zonneveld, 2002). Enhanced preservation in anoxic sediments extends to other organic remains, including the fish scales that are abundant in some sapropel deposits (Jorissen, unpublished observations).

## 4.2 Sediment chemistry

### 4.2.1 Organic C, N and their stable isotopes

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Areas prone to hypoxia typically exhibit an upcore increase in the accumulation of total sedimentary organic carbon (TOC). Such trends could reflect the fact that labile components in younger sediments have had less time to decompose, an increase in water column productivity, an increase in the supply of allochthonous organic matter, an intensification of oxygen depletion, or a combination of these factors (Cornwell et al., 1996; Zimmerman and Canuel, 2002). Particular debate has centred around the extent to which increases in TOC reflect increased organic flux or enhanced preservation (e.g. Cowie, 2005 and references therein).

Studies of historical records have usually linked increases in TOC to increased phytoplankton production. In Bunnefjord (the innermost part of the Oslo Fjord), TOC concentrations were highest in sediment layers corresponding to the period (1945–1982) of most intense anthropogenic eutrophication (Pinturier-Geiss et al., 2002). Along the west coast of India off Goa, large, upcore increases in TOC and TN observed in the sediments of the inner- and mid-shelf since ~1950 have been attributed to enhanced



biological production fuelled by both an intensified summer monsoon-driven upwelling and a greater runoff of nutrients from land (D'Souza, 2007; Agnihotri et al., 2009; Kurian et al., 2009). Similar increases in organic carbon accumulation rates in cores from the Southern Baltic Sea are likewise believed to reflect increased input of organic carbon due to eutrophication (Andrén, 1999; Andrén et al., 1999; Struck et al., 2000). However, although eutrophication is usually considered to be the main driver of upcore TOC increases, it is often associated with bottom-water hypoxia. In Chesapeake Bay, the TOC content of the sediment has increased over the last 200 years, with abrupt increases at certain horizons (1880–1915, 1945), probably due to enhanced productivity (Zimmerman and Canuel, 2002). TN also increased upcore in Chesapeake Bay sediments and was linearly correlated with TOC values (Cooper and Brush, 1991; Bratton et al., 2003). At the same time, other indicators (DOP and AVS/CRS, see above) suggested that hypoxia/anoxia had increased, coincident with eutrophication (Cooper and Brush, 1993; Zimmermann and Canuel, 2002). In Charlotte Harbor (Florida), Turner et al. (2006) interpreted a post-1950 rise in sediment organic carbon content in cores to indicate an increased input from phytoplankton production. Based on a multiproxy approach, they also concluded that the increased organic matter loading was responsible for an expansion in the area impacted by hypoxia.

Stable isotopes of carbon and nitrogen can yield important insights into the sources and processing of organic matter preserved in sediments. In historical records from coastal settings, the stable carbon isotope signal is usually interpreted in terms of the relative contribution of terrestrial versus marine carbon sources. Thus, trends in  $\delta^{13}\text{C}$  over a 2700-year period in Chesapeake Bay are believed mainly to reflect changes in the relative importance of estuarine productivity and terrestrial material delivered to the Bay, which are related ultimately to climatic fluctuations (Bratton et al., 2003). Voss and Struck (1997) considered increases in  $\delta^{13}\text{C}$  in the Baltic cores over the last 100 years to be caused by eutrophication linked to increased primary productivity in the marine environment and the rivers discharging into it. Ellegaard et al. (2006) reached a similar conclusion based on a core from Mariager Fjord (Denmark). C-isotope-based

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approaches have also been applied in the Gulf of Mexico (Eadie et al., 1994; Turner et al., 2006) and New Bedford Harbor, Massachusetts (Chmura et al., 2004), where they indicate an increased input of marine-source carbon in more recent years consistent with eutrophication (Fig. 13). However, over the western Indian shelf, where the organic matter is largely of marine origin, no major change in  $\delta^{13}\text{C}_{\text{org}}$  appears to have occurred during the last 1300 years (Agnihotri et al., 2009).

Stable nitrogen isotope ratios ( $^{15}\text{N}/^{14}\text{N}$ ; i.e.  $\delta^{15}\text{N}$ ) have been widely used as indicators of anthropogenic eutrophication in aquatic ecosystems (Voss et al., 2000; Cole et al., 2004). Compared to carbon isotopes, differences between the  $\delta^{15}\text{N}$  values of the terrestrial and marine-derived material are small, with the former characterized by slightly lower ratios (Wada and Hattori, 1991). However, riverine and terrestrial nitrogen is intensively processed at the land-ocean interface with the consequence that dissolved and particulate  $\delta^{15}\text{N}$  become rather heavy (e.g. Middelburg and Herman, 2007). These heavy  $^{15}\text{N}$  values are transferred up the food chain and have been used to assess the anthropogenic N loadings in coastal settings. The isotopic composition of fixed N (mostly nitrate) in the ocean is determined by its cycling within the water column and sediments (Brandes and Devol, 2002). Enrichment in  $^{15}\text{N}$  in organic matter can arise either from partial nitrate utilization (Altabet and Francois, 1994) or uptake of isotopically heavy  $\text{NO}_3^-$  arising from water column denitrification (i.e. preferential loss of  $^{14}\text{NO}_3^-$ ) (Brandes et al., 1998; Brandes and Devol, 2002; Bratton et al., 2003). Photosynthesis involving nitrogen originating from denitrification can produce a particularly strong signal, with  $\delta^{15}\text{N}$  values  $>9\text{‰}$ , much higher than those generated in oxic settings (Hendy and Pedersen, 2006). Accordingly, eutrophication with extensive nitrogen recycling of external nitrogen and hypoxia with water-column denitrification may be associated with heavier  $\delta^{15}\text{N}$  values for sedimentary organic matter. Bratton et al. (2003) attributed the trend towards heavier  $\delta^{15}\text{N}$  values since about the end of the 18th Century in Chesapeake Bay cores mainly to this mechanism. The upturn coincided with the onset of land clearance by colonists which would have increased erosion and nutrient input to the Bay. Fluctuations superimposed on this overall trend

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were probably caused by climatic oscillations (wet-dry cycles). Eadie et al. (1994), Kappila et al. (2005) and Ellegaard et al. (2006) also suggest denitrification as a possible contributor to increased sediment  $\delta^{15}\text{N}$  measured in core profiles. Rises in total organic nitrogen and  $\delta^{15}\text{N}$  values in cores from the Baltic (Voss and Struck, 1997; Voss et al., 2000; Struck et al., 2000) and Laajalahti Bay, Finland (Kappila et al., 2005) are believed to reflect eutrophication caused by riverine inputs of land-derived nitrate enriched in  $^{15}\text{N}$ .

Elevated  $\delta^{15}\text{N}$  values do not necessarily occur in all coastal hypoxic zones. In the intensely denitrifying waters of the Indian shelf, Naqvi et al. (2006b) found relatively modest enrichment of  $^{15}\text{N}$ , corresponding to a fractionation factor ( $\epsilon$ ) of 7.2–7.7‰, compared to values of 25–30‰ in the open-ocean denitrification zones (Brandes et al., 1998). Consistent with this apparently lower fractionation factor, Agnihotri et al. (2009) reported an average  $\delta^{15}\text{N}$  of only 6.6‰ for the upper 5 cm in a sediment core extending back to ~1300 AD (Fig. 14). In fact, sedimentary  $\delta^{15}\text{N}$  at this location appears to have decreased slightly but steadily over the last 150–200 years, whereas productivity proxies suggest an increase in productivity over the same period (Kurian et al., 2009). During the Maunder Minimum of the Little Ice Age, however, the  $\delta^{15}\text{N}$  value fell to ~6‰, probably caused by a decrease in productivity. Overall the isotope data from the Indian shelf casts doubts over the universal applicability of  $\delta^{15}\text{N}$  as a robust denitrification proxy.

#### 4.2.2 Biochemical indicators: pigments and lipid biomarkers

Sediment cores yield pigments derived from phytoplankton, mainly diatoms but also dinoflagellates and cyanobacteria, which can be analysed by high performance liquid chromatography (HPLC). Chlorophyll and its degradation products (phaeopigments) reflect overall phytoplankton biomass, while carotenoid pigments serve as biomarkers (molecular fossils) for particular phytoplankton taxa, e.g. fucoxanthin for diatoms and prymnesiophytes (Wright and Jeffrey, 1987) and zeaxanthin for cyanobacteria (Guillard et al., 1985). However, the differential preservation of pigments, discussed below,

means that changes in their concentrations over time cannot be translated directly into phytoplankton community composition.

In the Gulf of Mexico, upcore increases in chlorophyll *a* and phaeopigments were gradual from 1955 to 1970 and then more rapid to the late 1990s (Rabalais et al., 2004). Zeaxanthin shows similar pattern, indicating that cyanobacteria were not a major biomass component of phytoplankton prior to 1950s. The increase in chloropigments was most evident in areas that are presently most prone to hypoxia, indicating either a sharp increase in eutrophication, or hypoxia, or both these phenomena, in the latter part of the 20th Century. Cores from Charlotte Harbor showed similar pigment increases as well as BSi increases since 1950 (Turner et al., 2006). Peaks in chlorophyll *a*, its breakdown products and diatoxanthin, corresponding to high nutrient loads, were reported from Laajalahti Bay (Kauppila et al., 2005).

Lipid biomarkers (sterols, fatty acids, fatty alcohols) preserved in sediments are powerful tools for tracing environmental changes (Ficken and Farrimond, 1995; Zimmerman and Canuel, 2000, 2002; Smittenberg et al., 2004, 2005). Marine organisms, including bacteria, diatoms and dinoflagellates, produce characteristic lipid biomarkers that track changes in plankton communities in response to eutrophication. In a core from Chesapeake Bay, lipid biomarkers derived from phytoplankton and microbial sources consistently increased in concentration above the 1934 horizon (Zimmermann and Canuel, 2000). This two- to ten-fold increase was interpreted as being depositional in origin and due to increased production of plankton and microbially derived organic matter. It was concurrent with increases in biogenic silica, a proxy for diatom production. Two other cores from Chesapeake Bay also exhibited increases in lipid concentrations during the 20th Century, although the precise patterns were different (Zimmerman and Canuel, 2002). Shifts in the ratios of biomarkers for different phytoplankton taxa revealed changes in plankton composition, for example, an increase in the abundance of dinoflagellates relative to diatoms after 1915 (Zimmerman and Canuel, 2002). In Kyllaren Fjord (Norway), there has been a substantial increase in the accumulation of biomarkers characteristic of marine planktonic organisms, presumably linked to eu-

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trophication resulting either from natural processes or human activities (Smittenberg et al., 2004). Sterol records, particularly for dinosterol, preserved in a core from the Indian coast (the same one analysed by Agnihotri et al., 2008), closely resembled records for total solar irradiance, suggesting insolation control of upwelling-driven productivity (Kurian et al., 2009). Data for sterols and other productivity proxies (OC and TN), indicate that the productivity in the last few decades has been the highest seen during the last 700 years. Interestingly, while the phytol content appears to have declined recently, that of dinosterol continues to rise, possibly due to a shift in phytoplankton composition in response to eutrophication.

In the context of this review, green pigmented sulphur bacteria (Chlorobiaceae) are of particular interest as biomarker sources since they are associated with sulphidic/anoxic conditions rather than with eutrophication. These strictly anaerobic organisms require light and H<sub>2</sub>S and synthesise two characteristic pigments, bacteriochlorophylls and isorenieratene. Farnesol, the esterifying alcohol of the *Chlorobium* chlorophylls has been found in the sediments of the euxinic Kau Bay (Middelburg et al., 1994). Isorenieratene and its derivatives are preserved in sediments dating back to the Ordovician (Koopmans et al., 1996). Their occurrence in some eastern Mediterranean sapropels of Pliocene age has been used to infer the existence of sulphidic/anoxic conditions in the photic zone of the water column (Passier et al., 1999; Menzel et al., 2002). The sedimentary record of isorenieratene in Black Sea sediments suggests that the photosynthetic green sulphur bacteria have been active in the Black Sea for the last 6200 years and that penetration of the photic zone by anaerobic waters is a natural phenomenon (Sinninghe-Damsté et al., 1993). In shallow water, where light penetrates to the seafloor, green pigmented sulphur bacteria may also colonise the sediment. Chen et al. (2001) used bacteriochlorophyll-*e* and its breakdown products derived from two species of green sulphur bacteria (*Chlorobium phaeovibroides* and *C. phaeobacteroides*) to trace the development of photic-zone hypoxia on the Louisiana shelf over a 100-yr period (Fig. 15). Although these anaerobic prokaryotes require light, they can survive where light levels are very low and so it is unclear whether they were living at

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the sediment-water interface or in the overlying water column. Nevertheless, Chen et al. (2001) regard these pigments as the “first direct and unbiased indicators” of bottom-water hypoxia (as opposed to sediment anoxia) on the Louisiana shelf. Trends in the bacterial pigment data were the same as those for other indicators of eutrophication and hypoxia (Rabalais et al., 2007). In Kyallen Fjord, a euxinic Norwegian fjord, isorenieratene and okenone (the latter derived from purple sulphur bacteria) indicated that anoxic conditions had persisted throughout the 400-yr record preserved in a sediment core (Smittenberg et al., 2004).

It should be noted that the microbial decomposition of organic matter, including biomarkers, operates at different rates in oxic and hypoxia settings. In the Arabian Sea, the accumulation and preservation of different biomarkers can vary by a factor up to 10 between severely hypoxic [ $O_2=0.05 \text{ mL L}^{-1}$  ( $=2.2 \mu\text{M}$ )] and oxic sediments (Sinninghe-Damsté et al., 2002). Niggemann and Schubert (2006) showed that concentrations of total fatty acids in sediments from the core of the OMZ off central Chile ( $O_2=0.01 \text{ mL L}^{-1}$ ) were three to four times higher than in sediments from below the OMZ. This suggested enhanced preservation of these compounds in hypoxic environments, although the shallower water depth in the OMZ, and hence less efficient water column degradation, may also have been a factor (Niggemann and Schubert, 2006). The sensitivities of biomarkers to oxic decomposition also vary, and their accumulation rates therefore depend strongly on exposure to well-oxygenated conditions. For example, the carotenoid fucoxanthin, biphytane diols and dinosterol/dinostanol are all degraded to a greater extent than other marine biomarkers such as the alkenones and long-chain diol/keto-ols (Sinninghe-Damsté et al., 2002). Concentrations of phytol and dinosterol are higher (six and two times, respectively) in the Arabian Sea oxygen minimum zone (OMZ) core, where the bottom water is severely hypoxic, than outside the OMZ (Schulte et al., 2000). Lycopane, a biomarker probably derived from phytoplankton, also exhibits enhanced preservation in sediments deposited in the core regions of the Arabian Sea and Peru OMZs. This observation led Sinninghe-Damsté et al. (2003) to propose the ratio between lycopane and terrestrial  $C_{31}$  n-alkane as an oxygenation

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proxy in ancient marine sediments.

### 4.3 Indirect proxies

Although changing oxygenation regimes are often linked to productivity, they can also be forced indirectly by additional factors for which we have proxies. For example, in coastal embayments such as Chesapeake Bay, hypoxia development may be closely linked to stratification, which in turn can be induced by increased precipitation and freshwater input (Cronin et al., 2000; Hagy et al., 2004). Precipitation may also exacerbate nutrient enrichments resulting from fertilizer use, and often will fluctuate considerably on multidecadal time scales. Historical records of salinity variability can reflect precipitation changes. Oxygen isotopes ( $\delta^{18}\text{O}$ ) of foraminiferal tests are often used to estimate salinity in sediments and provide precipitation reconstructions for Chesapeake Bay (Saenger et al., 2006). Land use also profoundly influences vegetation, hydrologic pathways and patterns, nutrient runoff, and precipitation, with important consequences for eutrophication and hypoxia. Proxies for land use may include pollen shifts and salinity indicators as well as the other proxies discussed above (Willard and Cronin, 2007).

### 4.4 Multiproxy approaches

The experience of palaeoceanographers suggests that different proxies often yield inconsistent results. The use of several proxies in combination usually provides more convincing reconstructions of past oceans than single indicators (Guiot and de Vernal, 2007; Tribovillard et al., 2006). Multiproxy approaches are being used increasingly in historical studies. One example is the study of Turner et al. (2006) in Charlotte Harbor estuary (Florida). They utilized a suite of biological and geochemical proxies, including TN, TOC, C:N, P, trace metals, stable isotopes (S,N,C), biogenic silica and phytoplankton pigments (Chl a, caretonoids), to reconstruct conditions in the estuary between 1800 and 2000. Together, these proxies suggested that phytoplankton production had

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increased in response to increased nitrogen inputs, leading to an expansion of the area impacted by hypoxia. This study influenced water resource management policy, helping to convince the local management agency of the need to control nutrient inputs into the estuary. Rabalais et al. (2007) presented a synthesis of previous studies in the northern Gulf of Mexico that used a range of indicators for eutrophication (diatoms and biogenic silica, organic carbon of marine origin, phytoplankton pigments) and hypoxia (the mineral glauconite, bacterial pigments, faunal indices) (Fig. 5). Their conclusions were similar to those of Turner et al. (2006); phytoplankton production has increased sharply during the 20th Century, leading to severe or persistent hypoxia which began in the 1960s and became most pronounced in the 1990s.

## 5 Discussion

### 5.1 Historical reconstructions as part of palaeoceanography

Reconstructions of environmental changes caused by human activities are an aspect of palaeoceanography. This discipline depends to a large extent on the use of proxies, or in many cases suites of proxies used in combination, to infer the environmental conditions under which sediments were deposited in former seas and oceans (Hillaire-Marcel and de Vernal, 2007). Palaeoceanographic studies often focus on deep-water, open ocean environments, but also include oxygen-depleted, coastal basins like the Santa Barbara Basin, the Cariaco Basin, or the Saanich Inlet. Time periods studied range from very short (“historical”) records to those that cover very long periods,  $10^5$  years or longer.

The historical studies reviewed in this paper are usually based on nearshore sediment records. Since they require continuous sedimentary sequences, the systems studied are often situated in protected environments, such as inlets or basins. In some cases, documentary records are available and can be compared to sediment proxy records (Chesapeake Bay for example). In other areas, human settlements have been

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sparse and historical information is less complete (Cariaco Basin). However, to a large extent, studies of human impacts are based on tracers preserved in dated sediment cores. Like those conducted in deep-water, open ocean areas, such studies are concerned with environmental changes over time, particularly in surface productivity and bottom-water oxygenation.

Historical reconstructions are typically relatively local, focused on coastal waters, and represent short time periods in which human influences might be recorded (usually centuries). Because of high sedimentation rates, coastal records can often be examined at subdecadal or even shorter (annual or seasonal) time scales (Cronin et al., 2003; Rabalais et al., 1996). For example Black et al. (2007) presented an eight-century record of water temperature (based on planktonic foraminiferal Mg/Ca ratios and oxygen isotopes) with a resolution of 1–1.5 yrs for the anoxic Cariaco Basin (450-m water depth), where the sediments are laminated and sedimentation rates can reach 100 cm/kyr. However, because of sampling limitations, there are few proxy studies at these resolutions.

In general, coastal sediments are more dynamic than the floor of the deep ocean, and shallow-water communities are therefore subject to a greater range of environmental influences than those living in the deep sea (Gage and Tyler, 1991). This creates a higher natural variability from which changes caused by human activities must be distinguished. Greater variability in coastal areas compared to deeper water may also make it more difficult to establish a dating framework for sediments. For example, sedimentation rates determined in the top 10 cm of a core cannot be automatically extrapolated to deeper layers. However, a combination of radioisotopes, including (depending on the site)  $^{210}\text{Pb}$ ,  $^{234}\text{Th}$ ,  $^7\text{Be}$ ,  $^{137}\text{Cs}$  and  $^{14}\text{C}$ , can provide more accurate estimates of sediment age within a single sediment core and help to identify confounding factors such as bioturbation.

The application of palaeoceanographic approaches to reconstruct post-industrial environments in coastal settings requires critical evaluation. Some of the proxies commonly used in palaeoceanography have been applied to historical records but oth-

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ers, for example, the magnetic properties of sediments, trace metals and bacterial biomarkers, have been either unused or under-used. Conversely, historical records from coastal areas pose specific problems, leading to the development of proxies (e.g., the *Ammonia-Elphidium* ratio) that are applicable only in shallow water.

## 5.2 Distinguishing hypoxia and eutrophication

As emphasised elsewhere in this review, it is often difficult to distinguish the effects of hypoxia and eutrophication on proxies preserved in the sediment record. Certain species and morphotypes of foraminiferans, the most widely-used benthic faunal indicators, are common in hypoxic environments. In addition to oxygen, however, they appear to respond to an enhanced food supply resulting from eutrophication, as well as an absence of predators and competitors. The remains of diatoms, dinocysts, and eukaryotic biomarkers derived from the water column, generally indicate eutrophication, but their preservation is enhanced to a variable extent in oxygen-depleted sediments. Nevertheless, a wide range of different indicators is now available and these have been applied in many coastal systems that have experienced eutrophication and worsening oxygen conditions in recent decades and centuries.

The degree of sensitivity to hypoxia and eutrophication varies among proxies (Table 2). Some of the geochemical and biochemical proxies, including redox-sensitive elements such as Mo and Re, and bacterial biomarkers (isorenieratene) for photosynthetic sulphur bacteria, appear to be hypoxia specific. However, even in these cases, interpretation is not always simple. Enrichment of Mo, for example, represents conditions at least several cm deep in the sediment rather than in the overlying water. Bacterial biomarkers may come from bacteria in the water column, or from surficial sediments in the photic zone that are in contact with overlying oxygenated water. Sediment laminations require an absence, or virtual absence, of bioturbating macrofauna and therefore provide good evidence of bottom-water hypoxia, except in areas where sedimentation rates are so high that they prevent bioturbators from colonising the sediment. The best evidence for hypoxic conditions, either in the upper water column or the bottom wa-

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ter, comes from multi-proxy studies. A combination of the above-mentioned indicators that are most strongly influenced by hypoxia with those that reflect a combination of eutrophication and hypoxia (e.g. BSi) can yield the most convincing conclusions, as well as supporting interpretations of more equivocal indicators such as foraminiferans (Fig. 5).

### 5.3 Calibration of proxies

In a thought-provoking critique, Murray (2001) drew attention to a number of problems in the interpretation of proxies based on foraminiferans, particularly indicators for surface productivity and oxygen (see also Murray, 2006). Among the points made by Murray is that the abundance of particular species is related to a particular environmental variable only close to its tolerance limit for that variable. Despite these difficulties, palaeoceanographers have suggested several quantitative, foraminiferal-based approaches to the estimation of hypoxia that could be applied to historical records in coastal settings (Jorissen et al., 2007). All of them involve the recognition of species characteristic of oxic and hypoxic environments. Murray (2006) considers the transfer function proposed by Jannink (2001) to be the “potentially most successful”. The argument underlying the proposed proxy is that the relative abundance of oxyphilic species increases as oxygen concentrations rise, because the thickness of the oxygenated, and therefore habitable, sediment layer increases. Jannink et al. (2001) used their transfer function to reconstruct oxygen values based on the 160-year sediment record of Bar-mawidjaja et al. (1995) from the North Adriatic. The results agreed well with the actual historical oxygen measurements given by Justić et al. (1987). However, according to Jorissen et al. (2007), it is more likely that these shallow-infaunal species respond to inputs of labile organic matter rather than to oxygen-related changes in habitat availability linked to oxygen concentrations. Schmiedl et al. (2003) proposed an interesting method that combined the relative proportion of oxyphilic species (“high oxygen indicators”) with a measure of species diversity. The degree of oxygenation was indicated on a scale of 0 (anoxic) to 1 (oxic) rather than by actual values. The authors applied it

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to Mediterranean sapropels with convincing results (Fig. 16).

All current foraminiferal-based oxygen proxies suffer from drawbacks (Jorissen et al., 2007), the most severe being the inversely-related effects of organic fluxes and oxygen concentrations (Levin and Gage, 1998; Gooday, 2003). Also, most foraminiferans live within the sediment and therefore reflect oxygen levels within the sediment pore-water rather than in the overlying water (Gooday et al., 2000; Murray, 2001; Jorissen et al., 2007). This can lead to deep-infaunal, hypoxia-tolerant taxa being abundant in settings where the bottom-water is fully oxic (Sen Gupta et al., 1981; Gooday et al., 2001; Fontanier et al., 2005). Jorissen et al. (2007) suggest that below a certain threshold, the proportion of deep-infaunal foraminiferans depends on the elimination of less tolerant species by hypoxia. Hence, at least in principle, the percentage of deep-infaunal taxa can be used as a quantitative estimator of bottom-water oxygenation. Jorissen et al. (2007) further suggest that for such a proxy to be reliable, it needs to be calibrated by reference to values derived from modern assemblages over a range of bottom-water oxygen concentrations between 0 and 2 mL L<sup>-1</sup> (=0–87 μM). Because benthic foraminiferal species used to reconstruct anthropogenic hypoxia in coastal settings are often still living in the same area, it should be easier to calibrate and test them according to the criteria set out by Jorissen et al. (2007).

Few other attempts have been made to quantify bottom-water oxygen proxies. Willard et al. (2003) estimated dissolved oxygen values in cores from Chesapeake Bay using a modern analogue technique based on dinocyst species (Fig. 12). Building on the biofacies approach of Savrda and Bottjer (1991), Behl (1995) recognised seven sedimentary facies in the Santa Barbara Basin, including four ranging from completely bioturbated to completely laminated. Behl and Kennett (1996) correlated these to oxygen concentrations of <0.1 mL L<sup>-1</sup> (<4.3 μM; well-laminated sediment), ~0.1 mL L<sup>-1</sup> (~4.3 μM; bioturbation limited to meiofauna, insufficient to destroy laminations), 0.1–0.2 mL L<sup>-1</sup> (4.3–8.6 μM; partial homogenisation) and >0.3 mL L<sup>-1</sup> (>13 μM; homogenised sediment) (Fig. 17). The 0.1 mL L<sup>-1</sup> bioturbation threshold is consistent with recent evidence from the Pakistan margin. However, it appears that the responses

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by benthic organisms to hypoxia are different in deep and shallow water (Levin et al., 2009), suggesting that quantitative proxies of oxygenation developed in the deep sea may not be applicable to coastal settings.

#### 5.4 Natural versus anthropogenic drivers of hypoxia

5 Areas of seafloor overlain by naturally oxygen-depleted water masses are widespread in deeper-water, e.g. the Cariaco and Santa Barbara Basins, the Black Sea, Kau Bay, and oxygen minimum zones, and greatly exceed those affected by anthropogenically-induced hypoxia (Navqui et al., 2000; Helly and Levin, 2004). In shallow water, natural decadal or multidecadal climatic shifts linked to changes in indices such as the North Atlantic Oscillation (NAO) can have a major impact, particularly on coastal ecosystems where seasonal hypoxia can develop as a result of increased water column stratification during the summer linked to increased precipitation and freshwater runoff (Saegner et al., 2006; Cronin and Walker, 2006; Willard and Cronin, 2007; Leipe et al., 2008). Although the trend in most river-dominated coastal systems is for an increase in the N and P loads over the last half century (Boesch, 2002; Turner et al., 2003a, b), hypoxia may also develop naturally off the mouths of major rivers (Van der Zwaan, 2000). When interpreting proxy records that suggest hypoxia, it is therefore important to keep in mind possible natural causes, often linked to climatic variation, in addition to human influences (Murray and Alve, 2002). Historical records, such as official documents and unoffical journals, can also be useful in distinguishing human from natural drivers of hypoxia (Jackson et al., 2001).

There are several examples of systems where anthropogenically-induced hypoxia is superimposed on natural cycles that have persisted over much longer time scales. Cronin and Vann (2003) review studies in Chesapeake Bay that used different oxygen-related proxies to identify periods of hypoxia lasting several decades, prior to the European colonization of North America (Fig. 6). In the absence of significant human activity, it is relatively easy to recognise these as natural events. They were probably related to changes in atmospheric circulation that caused climatic variations in the

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mid-Atlantic, for example, related to the North Atlantic Oscillation (Cronin et al., 2000, 2003). However, over the last 300 years, a combination of factors has modified Chesapeake Bay and the picture has become more complicated. Between 1700 and 1900, progressive eutrophication linked to forest clearance, which intensified erosion and the delivery of nutrients to the Bay, was superimposed on climatic changes involving periods of increased precipitation, run off, water column stratification and oxygen depletion. During the latter part of the 20th Century, human activities were the main cause of dramatically decreasing oxygen levels, but at times, climatic forcing probably still played a role (Cronin and Vann, 2003; Willard and Cronin, 2007). Understanding the interplay between natural climate forcing and human activities is key to conservation of habitat for two anadromous sturgeon species in Chesapeake Bay, one of which is endangered (Willard and Cronin, 2007). Faunal and geochemical proxies have revealed similar natural hypoxic events related to climatic fluctuations in sediment records from the Louisiana shelf (Swarzenski et al., 2008).

The influence of natural processes is also evident in Scandinavia, where isostatic uplift following the last deglaciation has reduced sill depth of some Norwegian fjords, inhibiting water exchange with the open sea, trapping particulate matter and promoting anoxic conditions (Husum and Alve, 2006; also Smittenberg et al., 2004; Pinturier-Geiss et al., 2002, and references therein). Other processes have been at work in the Gullmar Fjord, an unpolluted (since 1966) fjord on the Swedish west coast. Here, there has been a shift from a fairly diverse foraminiferal assemblage comprising typical shelf species also found in the adjacent Skaggerak/Kattegat to an assemblage dominated by the opportunistic, hypoxia-tolerant species *Stainforthia fusiformis* (Nordberg et al., 2000; Filipsson and Nordberg, 2004). The change to a *S. fusiformis* dominated assemblage resembled that seen in Frierfjord, Norway, where the underlying cause was clearly anthropogenic (Alve, 2000). In the Gullmar Fjord, however, the assemblage change coincided with a severe hypoxic event that occurred during the 1979/1980 winter and was sustained by an increased input of phytodetritus. A variety of climatic and oceanographic factors, including changes in the North Atlantic Oscillation leading to

predominantly westerly winds that inhibited the exchange of bottom water in the fjord and in the flow of Atlantic water into the North Sea, are believed to underlie this faunal shift (Filippson and Nordberg, 2004). In open-shelf settings, similar changes have arisen from alterations in the hydrodynamic regime related ultimately to climatic forcing, for example, on the northwestern Iberian shelf (Martins et al., 2007) and off Oregon (Grantham et al., 2004).

## 6 Future directions

1) The number of coastal areas that are oxygen-depleted to the extent that living resources are compromised continues to rise (Diaz and Rosenberg, 1995; Diaz, 2001; Diaz et al., 2004; Rabalais and Gilbert, 2008). There is an increasing need to study proxy records to recognise that these changes are occurring so that management interventions can be applied before the coastal ecosystem has declined irreparably (Rabalais et al., 2007). Areas of incipient hypoxia where the historical approach might be applicable exist off the Changjian (Yantze River) in the East China Sea (Li and Daler, 2004; Chen et al., 2007; Wei et al., 2007) and in the Southwestern Gulf of Mexico off the Coatzacoalcos River (Rabalais, 2004). Palaeoindicators of eutrophication and hypoxia in the Charlotte Harbor estuary in Florida prompted the water management authorities to begin planning improved waste-water treatment plants for expected population increases. In other areas, where industrialization and mechanized agriculture are trying to keep pace with a burgeoning population, management and remediation measures are less well developed.

2) As more and more coastal systems are impacted by human activities, either directly or indirectly through consequences such as climate change, our ability to document a “natural” coastal ecosystem becomes increasingly difficult. It is essential that these baseline conditions should be described before it is too late. The variability of existing relatively pristine systems, for example, in polar regions, should also be documented in order to distinguish between natural variation and anthropogenic impacts.

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3) Distinguishing the effects of eutrophication from hypoxia remains a difficult challenge. Many proxies reflect both phenomena; for example, increased pigment concentrations can be due to both enhanced preservation under oxygen-depleted conditions or elevated production. The use of multiple indicators, i.e. a combination of proxies related to eutrophication (e.g. diatoms, phytoplankton pigments, BSi) with hypoxia-specific proxies (redox-sensitive trace elements such as Mo, Rh, bacterial pigments, indicators derived from sulphide minerals), may help to distinguish between these two closely-related drivers of ecosystem change. Examples of this approach are provided by the studies of Zimmermann and Canuel (2002) and Turner et al. (2006).

4) Proxies require verification and calibration. Collections across a range of oxygen concentrations will help to relate bioindicators to ambient oxygen concentrations. In order to improve the reliability of faunal proxies, it is important to identify organisms that were living when collected. Fluorescent dyes may be an improvement on traditional methods such as rose Bengal staining which can yield false positives, particularly under hypoxic conditions (Bernhard, 2000). Laboratory-based experiments conducted using a range of coastal sediments, for example, to examine the geochemical responses of various metals and trace metals to lowered oxygen concentrations or anoxic conditions, provide another approach to calibration. Some proxies, including trace metals such as rhenium and the magnetic properties of sediments, have yet to be applied to records of human-induced hypoxia.

5) There is considerable scope for the development of new proxies, in addition to the calibration of existing ones. More use could be made of animal hard parts; for example, the carapaces of ampeliscid amphipods, polychaete setae, the jaws of dorvilleid polychaetes, echinoderm spines and ossicles and the shells of chemosynthetic bivalves (e.g. *Thyasira*). These and other biogenic structures may form successional sequences in response to spatial or temporal oxygen gradients in coastal and deeper-water settings (Harper et al., 1991; Diaz and Rosenberg, 1995; Rabalais et al., 2001; Rosenberg, 2001).



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**Table 1.** Studies using faunal and chemical indicators to trace human impacts in coastal environments. Lamin = Laminations; Glauconite = Glauconite; TrM = Trace metals; BSi = Biogenic silica. In area column B = Baltic; D = Denmark; F = Finland; N = Norway; S = Spain. See Brush (2001), Cronin & Vann (2003), Kemp et al. (2000) and Rabalais et al. (2007) for reviews of Chesapeake Bay and Gulf of Mexico records.

Area	Benthic indicators						Mainly water column indicators						
	Foraminifera	Ostracods	Lamin	Bacterial biomarkers	Sulphur Sulphides	Glauconite	TrM	Diatoms	BSi	Dinocysts	Bio-markers	TC, TN	Stableisotopes
<i>North America</i>													
St Lawrence Estuary	53									53		53	
Chesapeake Bay	27	20	47		17,18, 62, 67		1, 66	17,18	16,17,19,67	62	67-69	67-69, 9	68, 69, 9
Massach. estuaries										13	13,40	13	40
Long Is Sound	54									54		54	
Gulf of Mexico	8, 10, 30-33, 37-39, 48, 52	2, 44		12	10	30	52, 59	34,42	42, 58,59		43, 52	23, 52	23, 52, 59
Charlotte Harbor, Fla					60		60			60		60	60
<i>Europe</i>													
Bilbao estuary (S)	11												
Drammensfjord (N)	4		4, 50								50		
Frierfjord (N)	3										22		
Kyallenfjord (N)			49	49							49		49
Oslofjord (N)										21	36	36	
Mariager Fjord (D)								24	24	24	24	24	24
Roskilde Fjord (D)								14,15	15			14, 15	
Arkona Basin (B)													51
Oder Estuary (B)			35					5,6					51, 61
Bornholm Basin (B)													51
Gotland Basin (B)													51
Laajalahti Bay (F,B)								15, 28	15			15,28	28
North Adriatic	7							41		45			
<i>Japan</i>													
Osaka Bay	56, 57	64, 65											
Yokohama Port	55	25						46		29			
Hiroshima Bay		63											
Lake Hamana				26			26		26		26	26	
<i>India</i>													
Shelf off Goa												70	70-72, 71,72

1) Adelson et al. (2001); 2) Alvarez Zarikian et al. (2000); 3) Alve (2000); 4) Alve (1991); 5) Andr n (1999); 6) Andr n et al. (1999); 7) Barnawidjaja et al. (1995); 8) Blackwelder et al. (1996); 9) Bratton et al. (2003) ; 10) Brunner (2006); 11) Cearreta et al. (2000); 12) Chen et al. (2001); 13) Chmura et al. (2004); 14) Clarke et al. (2003); 15) Clarke et al. (2006); 16) Colman and Bratton (2003); 17) Cooper (1995); 18) Cooper and Brush (1991, 1993); 19) Cornwell et al. (1996); 20) Cronin and Vann (2003); 21) Dale (2000); 22) Dale (2000); 23) Eadie et al. (1994); 24) Ellegaard et al. (2006) ; 25) Ikeya (1995); 26) Itoh et al. (2003); 27) Karlson et al. (2000); 28) Kauppila et al. (2005); 29) Matsuoaka (1999); 30) Nelsen et al. (1994); 31) Osterman (2003); 32) Osterman et al. (2005); 33) Osterman et al. (2007); 34) Parsons et al. (2002); 35) Persson and Jonsson (2000); 36) Pinturier-Geiss et al. (2002); 37) Platon and Sen Gupta (2001); 38) Platon et al. (2001); 39) Platon et al. (2005); 40) Pospelova et al. (2002); 41) Puškarić et al. (1990); 42) Rabalais et al. (1996); 43) Rabalais et al. (2004); 44) Rabalais et al. (2002); 45) Sangiorgi and Donders (2004); 46) Sato (1995); 47) Schaffner et al. (1992); 48) Sen Gupta et al. (1996); 49) Smittenberg et al. (2004); 50) Smittenberg et al. (2005); 51) Struck et al. (2000); 52) Swarzenski et al. (2008); 53) Thibodeau et al. (2006); 54) Thomas (2006); 55) Toyoda and Kitazato (1995); 56) Tsujimoto et al. (2006a); 57) Tsujimoto et al. (2008); 58) Turner and Rabalais (1994); 59) Turner et al. (2004); 60) Turner et al. (2006); 61) Voss and Struck (1997); 62) Willard et al. (2003); 63) Yasuhara et al. (2003); 64) Yasuhara and Yamazaki (2005); 65) Yasuhara et al. (2007); 66) Zheng et al. (2003); 67) Zimmerman and Canuel (2000); 68) Zimmerman and Canuel (2001); 69) Zimmerman and Canuel (2002); 70) Kurian et al. (2009); 71) D'Souza (2007); 72) Agnihotri et al. (2009).

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**Table 2.** Indicators and their relationship to anoxia, hypoxia and eutrophication. See text for further details.

Indicator	Anoxia	Severe hypoxia (<5% saturation =0.2 mL L <sup>-1</sup> )	Moderate hypoxia (5–30% =0.2–1.2 mL L <sup>-1</sup> )	Eutrophication and increased organic flux to seafloor	Comment
<i>Benthic indicators</i>					
Foraminifera	Some species tolerant	Decrease in species number and diversity	Little apparent influence on many species	Strong influence on some species	Anoxia-tolerant species may respire nitrate. Interpretation of oxygen conditions confounded by organic enrichment.
Ostracoda	Absent	Decrease in diversity; a few tolerant species, some of which can survive sulphidic conditions	Some tolerant species	Abundance of some species may increase as a result of eutrophication	Most ostracods are intolerant of hypoxia
Bioturbation	Laminations	Incomplete laminations	Bioturbation limited to a few mm to cm at surface	Sediments usually fully bioturbated, but in Gulf Mexico, small, surface deposit-feeding opportunists that do not bioturbate are abundant	Laminations may also result from high sedimentation, regardless of overlying oxygen conditions.
Bio/ichnofacies	Traces and macrofossils absent	Shelly exaerobic facies recognised by bivalve shell detritus	Decreasing penetration of burrows with increasing hypoxia	Body and trace fossils usually common, but not in all environments	Oxygenated sediments with sufficient food resources are occupied by a mature, diverse, deep-burrowing benthos
DOP	Values >0.7	0.55–0.70			Reliable proxy for bottom-water oxygenation
AVS:CRS	+++				Useful indicator of anoxia, but depends on sediment accumulation rate



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**Table 2.** Continued.

Indicator	Anoxia	Severe hypoxia (L <sup>-1</sup> )	Moderate	Eutrophication and	Comment
Glauconite	Mineral forms under hypoxic and anoxic conditions				
$\delta^{13}\text{C}$ foram shell				Reflects uptake of photosynthetically fixed carbon	
$\delta^{13}\text{C}$ organic matter				Eutrophication leads to raised $\delta^{13}\text{C}$ values due to increased in situ marine phytoplankton production	Reflects relative contribution of terrestrial and marine carbon sources
$\delta^{15}\text{N}$		Heavy values reflect denitrification		Heavy values may reflect land derived N and eutrophication	$\delta^{15}\text{N}$ can be heavier when the nitrogen source is from regeneration (e.g. $\text{NH}_4^+$ )
$\delta^{34}\text{S}$	Increasing values indicate reducing conditions		Lower values because of active S re-oxidation cycle		
Sediment color and smell	Dark with strong smell of sulphide	Dark with strong smell of sulphide		Yellow to brown colour with smell of fresh organic matter	
Magnetic properties of sediment	Fine-grained, secondary minerals (e.g. greigite, pyrrhotite) responsible for magnetic properties			Coarse detrital minerals (e.g. haematite) responsible for magnetic properties	
<i>Water column indicators</i>					
Diatoms	Preservation may be enhanced			Overall diatom production, and ratio of planktonic (centric) to benthic (pennate) diatoms, increase with increasing eutrophication. Lightly silicified species may also increase	Good indicators of eutrophication

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Table 2. Continued.

Indicator	Anoxia	Severe hypoxia (L <sup>-1</sup> )	Moderate	Eutrophication and	Comment
Dinocysts	Enhanced preservation of organic-walled cysts destroyed in oxic conditions	A few species may be associated with hypoxia		Increases in heterotrophic cyst abundance often linked to eutrophication	Species composition reflects productivity, sea surface temperature, salinity, oxygenation and enhanced preservation of certain species under anoxic conditions
Trace metals:	Enrichment in Mo indicates anoxic, sulphidic conditions; enrichment in Rh indicates absence of free H <sub>2</sub> S	High enrichment in Mo vanadium, uranium, but relative low enrichment			In anoxic conditions, most metals exist in lower redox state (e.g. Fe <sup>2+</sup> , Mn <sup>2+</sup> ) and form insoluble sulphide compounds. In oxic conditions, metals exist in higher redox state (e.g. Fe <sub>2</sub> O <sub>3</sub> and MnO <sub>2</sub> )
BSi				Increases linked to eutrophication	Chemical proxy for diatom production
Pigments and lipid biomarkers	Enhanced preservation	DO=0.28 ml L <sup>-1</sup> leads to enhanced preservation of Chl a & phaeopigments, compared to oxic conditions Concentration of chlorins (Chl a breakdown products) in sediments increases with lower DO in water column; however degradation rates not influenced by bottom-water DO		Increased pigment and lipid concentrations generally indicate increased primary productivity	Biomarkers indicate relative abundance of different phytoplankton taxa. Low oxygen generally enhances preservation of biomarkers in the sediment.
Pigments or lipids from green pigmented sulphur bacteria (isorenieratine, B-chl, farnesol)	Indicate photic zone anoxia				Derived from strictly anaerobic organisms that require light and sulphide. B-chl in sediments; hypoxia-specific proxy.

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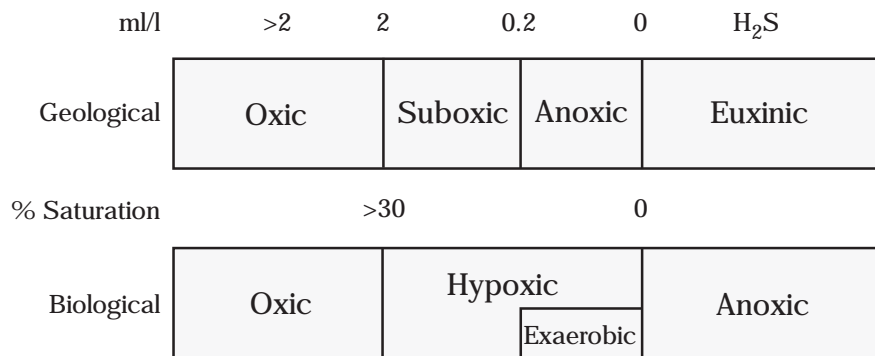


Fig. 1. Terminology used by geologists and biologists to describe oxygen depletion.

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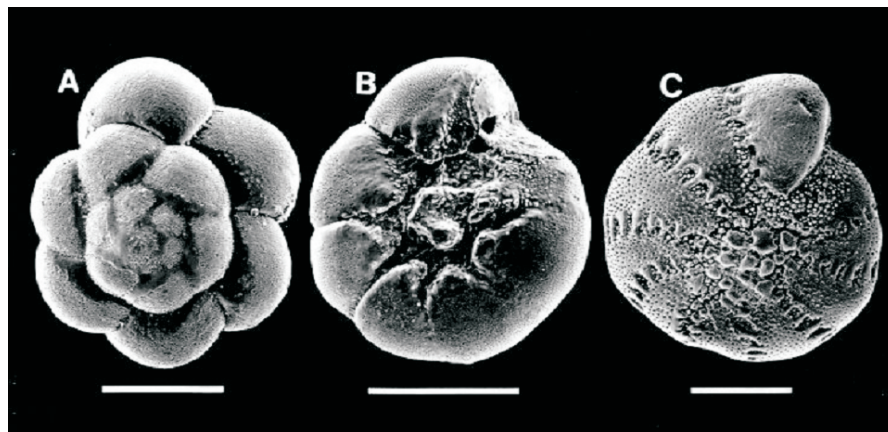
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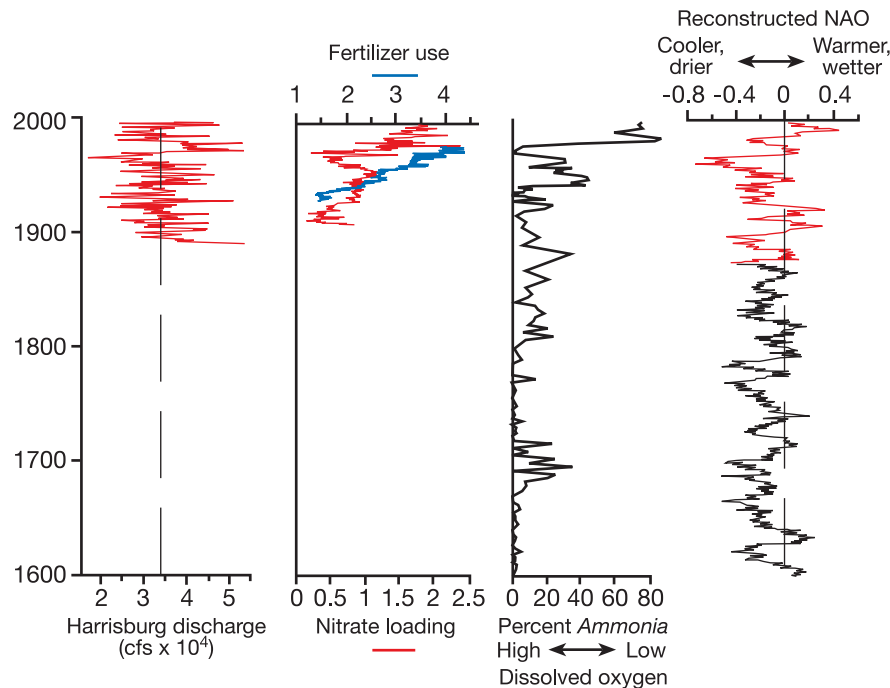


**Fig. 2.** Scanning electron micrographs of benthic Foraminifera tolerant or intolerant of hypoxia. (A–B) *Ammonia parkinsoniana*; (C) *Elphidium excavatum*. Specimens are from the Louisiana shelf, west of the Mississippi Delta. Scale bars=100  $\mu\text{m}$ . From Geology, vol. 24, B. K. Sen Gupta, R. E. Turner and N. N. Rabalais, Seasonal oxygen depletion in continental-shelf waters of Louisiana: Historical record of benthic foraminifers, pp. 227–230, 1996, copyright Geological Society of America.

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**Fig. 3.** Instrumental (red and blue lines) and proxy-based (black lines) records of **(a)** fluvial discharge at Harrisburg, Pennsylvania; **(b)** fertilizer in Chesapeake Bay watershed (blue line) and nitrate loading to Chesapeake Bay (red line); **(c)** dissolved oxygen based on percent abundance of *Ammonia* ; **(d)** North Atlantic Oscillation Index (NAO) reconstructed from tree-ring records. Redrawn from *Frontiers in Ecology and the Environment*, vol. 5, D. A. Willard and T. M. Cronin, *Paleoecology and ecosystem restoration: case studies from Chesapeake Bay and the Florida Everglades*, pp. 491–498, 2007, with permission from The Ecological Society of America.

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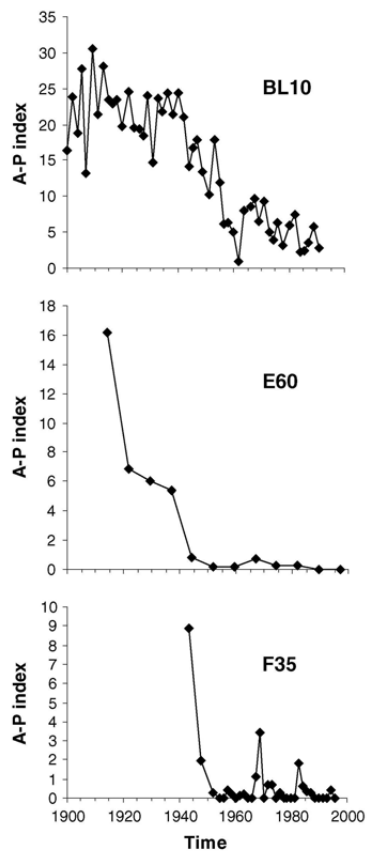
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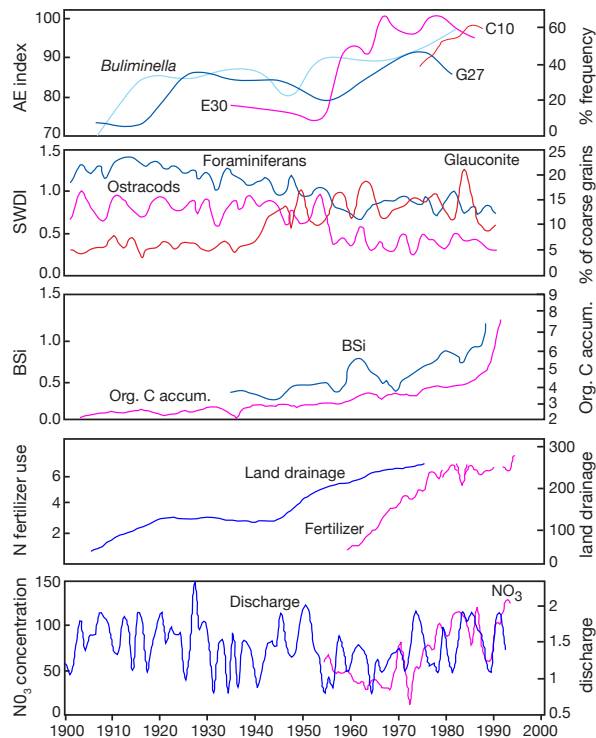


**Fig. 4.** The ratio between the abundance of agglutinated and porcelanous Foraminifera (the *A-P* Index) in three cores from the Louisiana inner continental shelf, SW of the Mississippi River delta: BL10 (29 m water depth), F35 (35 m), E60 (60 m). Reprinted from *Marine Micropaleontology*, vol. 54, E. Platon, B. K. Sen Gupta, N. N. Rabalais, R. E. Turner, Effect of seasonal hypoxia on the benthic foraminiferal community of the Louisiana inner continental shelf: the 20th century record, pp. 263–283, 2005 with permission from Elsevier.

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**Fig. 5.** Trends in faunal and environmental parameters in cores from the Louisiana coast, west of the Mississippi River delta. **(a)** The ratio between the abundance of *Ammonia* and *Elphidium* (A-E index) in three cores (C10, E30, G27), and the percentage abundance of *Buliminella* (G27, light blue line). **(b)** Shannon-Wiener diversity index (SWDI) for Foraminifera and ostracods and the percentage of the mineral glauconite among the coarser sediment grains (red line). **(c)** Frequency of biologically bound silica (BSi) and organic carbon accumulation rate in core E30. **(d)** Nitrogen fertilizer use in the Mississippi River basin and land drainage. **(e)** Nitrate concentration in the lower Mississippi River and lower Mississippi River discharge. Redrawn from Ecological Applications, vol. 15, supplement, N. N. Rabalais, R. E. Turner, B. K. Sen Gupta, E. Platon, M. L. Parsons. Sediments tell the history of eutrophication and hypoxia in the northern Gulf of Mexico, S129–S143, 2007, with permission from The Ecological Society of America.

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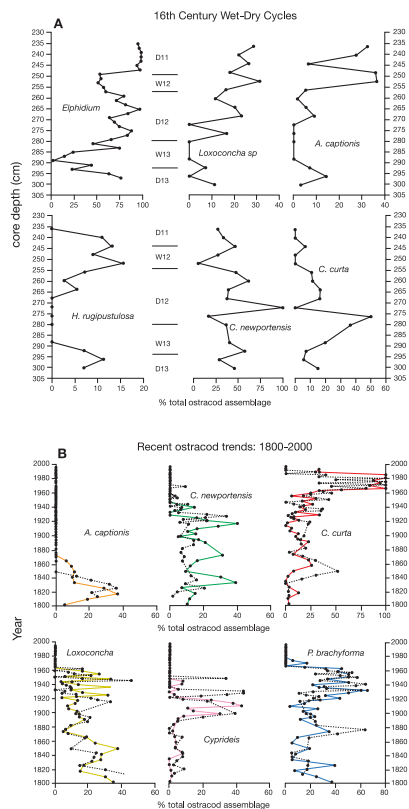
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**Fig. 6.** Ostracod records based on piston and gravity core samples from Chesapeake Bay, off the mouth of the Patuxent River (PTXT site). **(A)** Decadal-scale variability in 5 ostracod species and 1 foraminiferal species (*Elphidium* sp.) in sediments from a piston-core representing the period 1500–1650 AD, which featured three wet/dry climatic cycles (D11–13, W12–13). Distinct ostracod assemblages characterize each dry and wet period, reflecting inferred salinity, turbidity and oxygen conditions. **(B)** Variability in the abundance of 6 ostracod species during the past 200 years based on a piston core (PTXT-2-P-5, solid line) and a gravity core (PTXT-2-G-3, dashed line). Redrawn from Estuaries, vol. 26, T. M. Cronin and C. D. Vann, The sedimentary record of climatic and anthropogenic influence on the Patuxent Estuary and Chesapeake Bay ecosystems, pp. 196–209, 2003, with permission from Springer.

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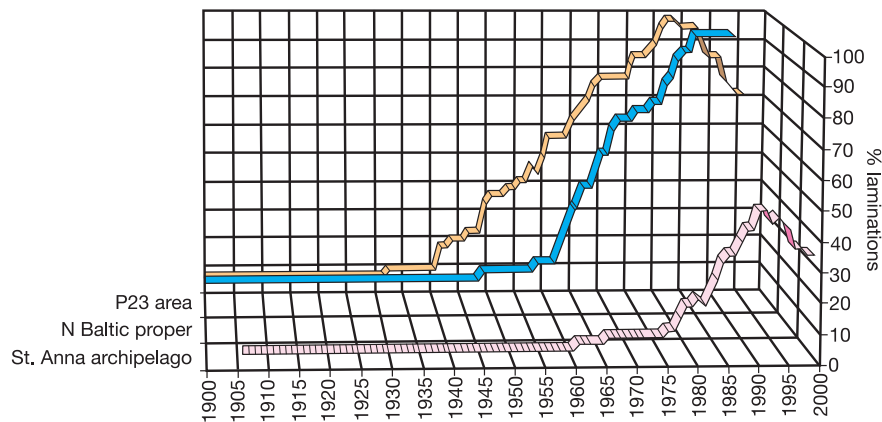
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**Fig. 7.** Laminated sediments in the Baltic Sea. The proportion of laminated sediments in relation to the total number of cores investigated in the St. Anna archipelago ( $n=48$  cores, pink line), the offshore Baltic proper ( $n=35$ , blue line) and the P23 area in the NW Baltic proper ( $n=40$ , orange line). Redrawn from Marine Pollution Bulletin, vol. 40, J. Persson and P. Jonsson, Historical development of laminated sediments – an approach to detect soft sediment ecosystem changes in the Baltic Sea, pp. 122–124, 2000, with permission from Elsevier.

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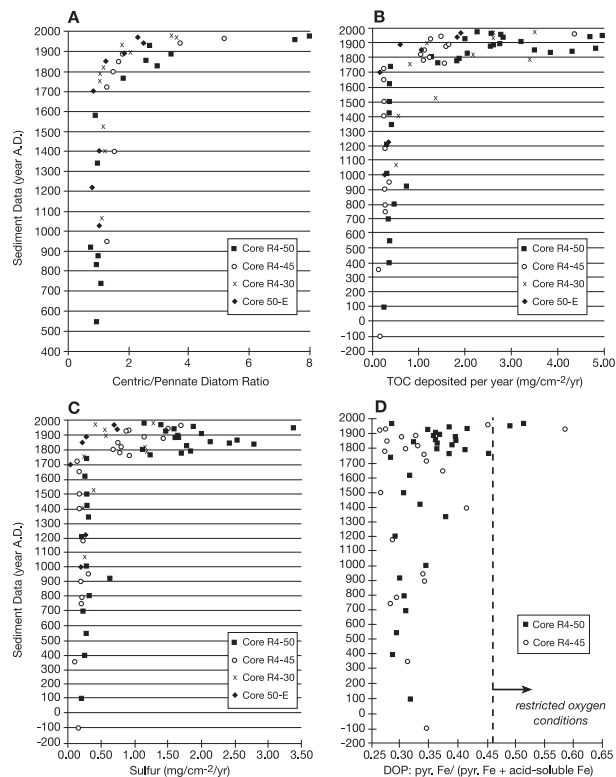
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**Fig. 8.** Long-term sediment records of eutrophication and hypoxia in cores from mesohaline part of Chesapeake Bay. **(A)** Centric:pennate diatom ratio; increase after 1940 indicate increased eutrophication. **(B)** Total organic carbon (TOC), **(C)** Preservation of sulphur, corrected for sedimentation rate. **(D)** Degree of pyritization (DOP); higher values indicate hypoxic conditions. Redrawn from *Estuaries*, vol. 16, S. R. Cooper and G. S. Brush, A 2500-year history of anoxia and eutrophication in Chesapeake Bay, pp. 617–626, 1993, with permission from Springer.

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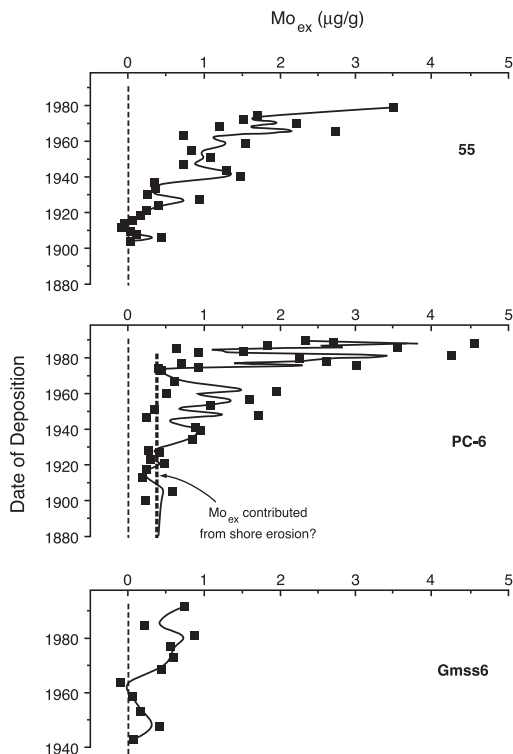
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**Fig. 9.** Excess molybdenum concentration ( $M_{ex}$ ) in three cores from the middle and upper part of Chesapeake Bay. The heavy dotted line in the central panel suggests a baseline input of  $M_{ex}$  from shore erosion. The curve shows the gross trends in the data, suggesting increasingly intense seasonal oxygen depletion. Redrawn from *Geochimica et Cosmochimica Acta*, vol. 65, J. M. Adelson, G. R. Helz, C. V. Miller, Reconstructing the rise of coastal anoxia: molybdenum in Chesapeake Bay sediments, pp. 237–252, 2001, with permission from Elsevier.

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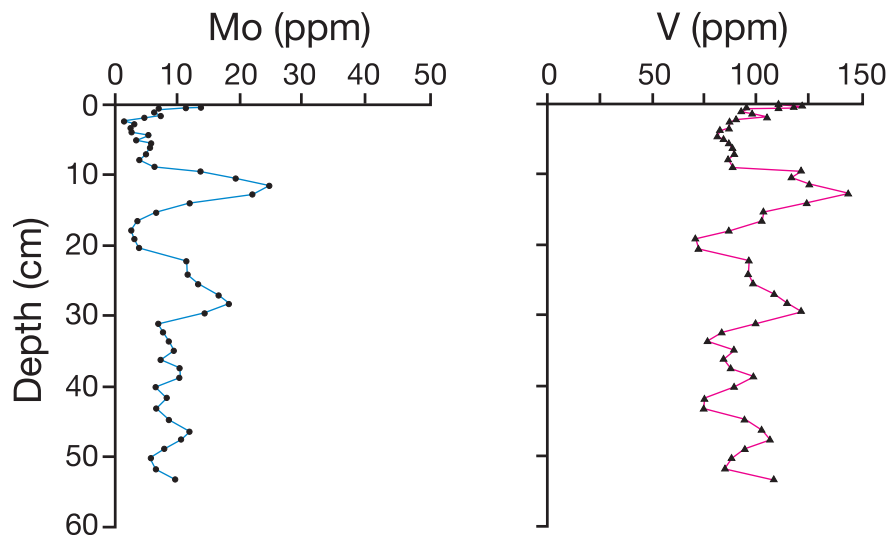
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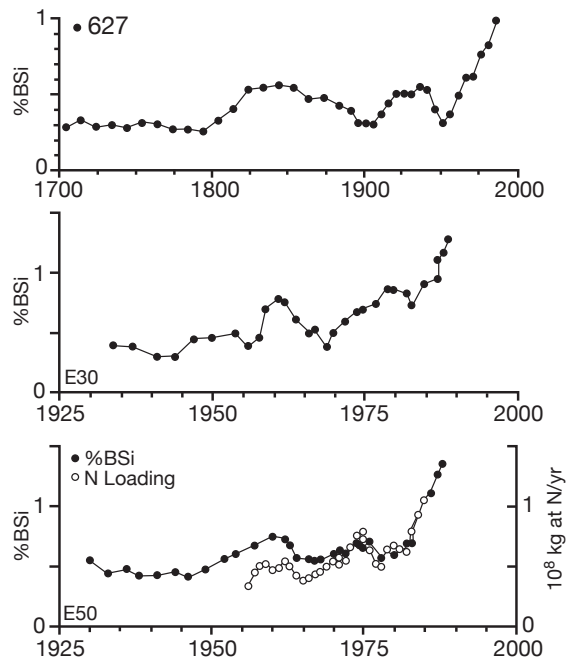
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**Fig. 10.** Mo, V profiles in sediments sampled using a box core (K9B1, 410 m water depth) in Kau Basin, northern Maluku Indonesia. Redrawn from J. J. Middelburg Ph.D. thesis.

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**Fig. 11.** Average concentrations of biologically bound silica (BSi) in sediment profiles of three dated sediment cores taken at 27–50-m water depth in the Louisiana inner continental shelf, SW of the Mississippi River delta. A 3-yr running average is plotted against time determined by Pb<sub>210</sub> dating. For Station E50, a 3-yr average nitrogen loading from the Mississippi River is superimposed on the BSi trend. Reprinted with permission from Macmillan Publishers Ltd. [Nature] (Turner, R. E., and Rabalais, N. N.: Coastal eutrophication near the Mississippi river delta, Nature, 368, 619–621), 1994.

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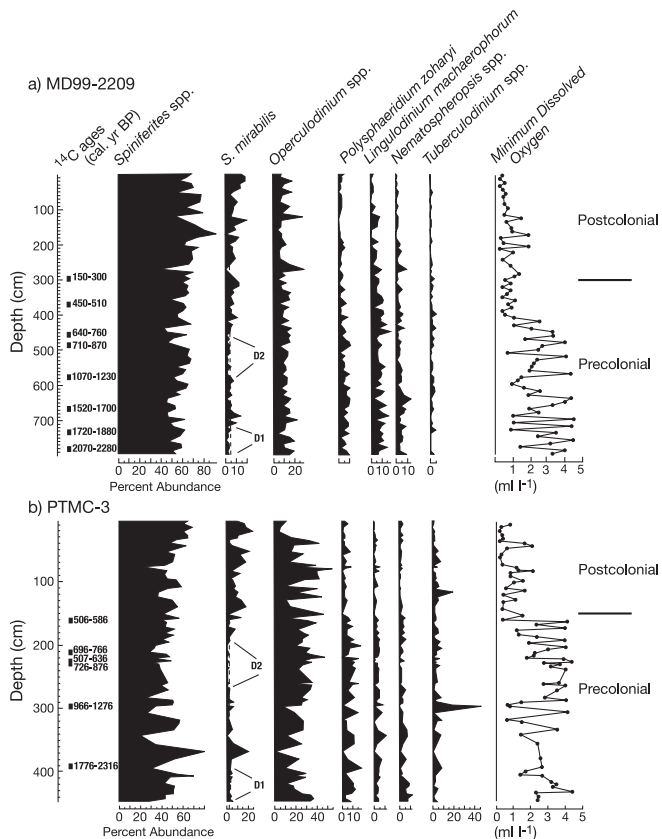
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**Fig. 12.** Percentage abundance of dinoflagellate cyst taxa in two cores from the mesohaline part of Chesapeake Bay. The dashed line indicates average precolonial abundance. Estimates of minimum annual dissolved oxygen are calculated from minimum DO values for 10 closest modern analogues. **(a)** Core MD99-2209, near mouth of Rhode River. **(b)** Core 1071 PTMC-3, near mouth of Potomac River. Redrawn from Holocene, vol. 13, D. A. Willard, T. M. Cronin and S. Verado, Late-Holocene climate and ecosystem history from Chesapeake Bay sediment cores, USA, pp. 201–214, 2003, with permission from SAGE Publications.

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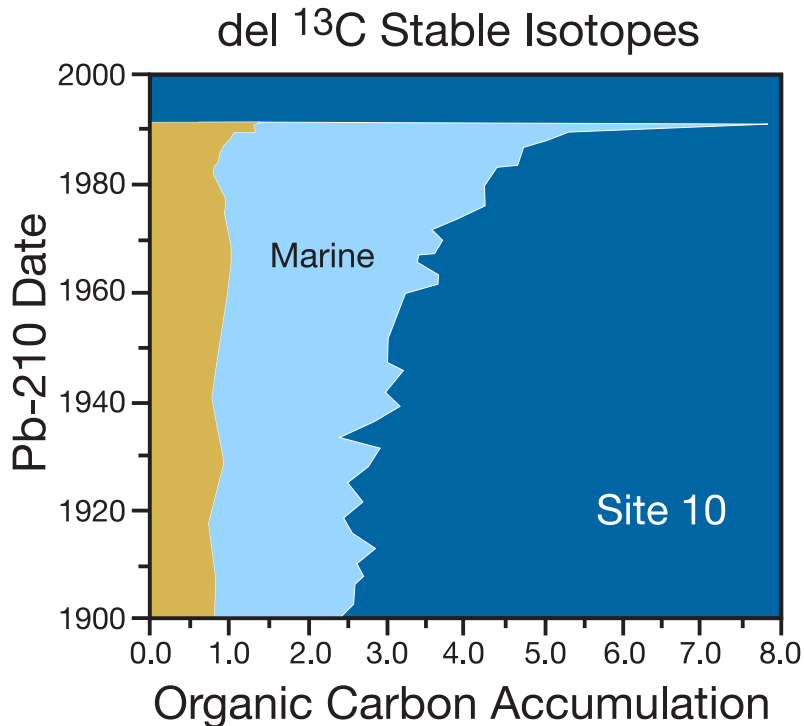
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**Fig. 13.** Organic carbon accumulation rate ( $\text{C cm}^{-2} \text{ yr}^{-1}$ ) in a box core obtained at Station 10 (29-m water depth) in a chronically hypoxic area on the Louisiana inner continental shelf, SW of the Mississippi River delta. The partitioning of organic matter into a terrestrial (yellow) and marine (light blue) component is based on  $\delta^{13}\text{C}$  data. Redrawn from *Estuaries*, vol. 17, B. J. Eadie, B. A. McKee, M. B. Lansing, J. A. Robbins, S. Metz, J. H. Trefry, Records of nutrient-enhanced coastal productivity in sediments from the Louisiana continental shelf, pp. 754–765, 1994, with permission from Springer.

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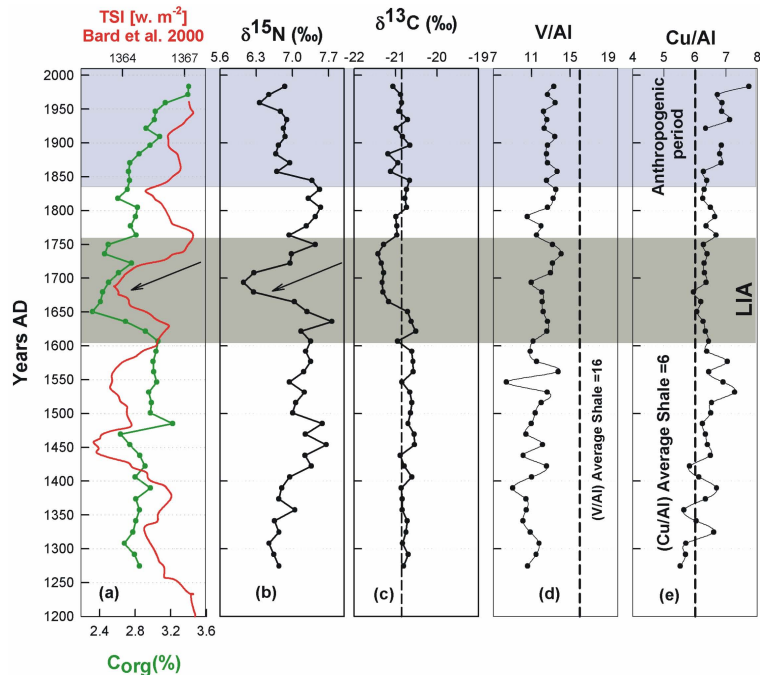
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**Fig. 14.** Profiles of selected productivity and redox proxies in the Gravity Core CR-2 from the inner shelf region off Goa (Lat. 14.8° N, Long. 74.2° E; water depth ~45 m) in the eastern Arabian Sea. **(a)** Total solar irradiance (TSI) from Bard et al. (2000) and organic carbon (C<sub>org</sub>), **(b)** δ<sup>15</sup>N of sedimentary organic matter, **(c)** δ<sup>13</sup>C of C<sub>org</sub>, **(d)** V/Al and **(e)** Cu/Al. The core was dated by C-14 and Pb-210 techniques. The arrows within the grey portion point to the Maunder Minimum of the Little Ice Age (LIA) whereas the upper portion of the core corresponding to the Anthropocene period is shown in purple (modified from Agnihotri et al., 2009).

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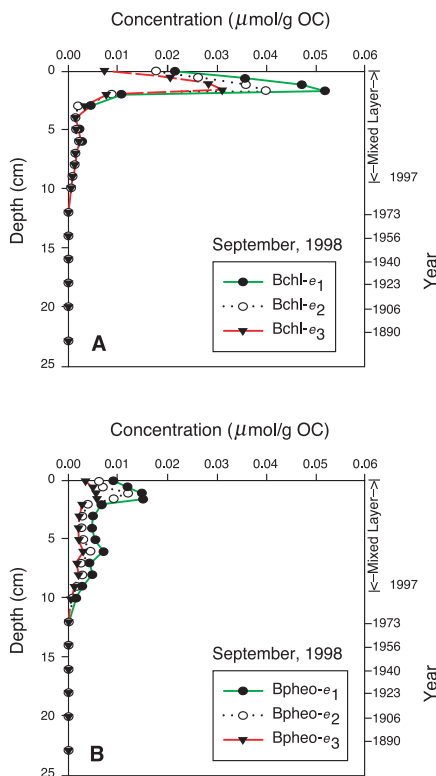
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**Fig. 15.** Down-core concentrations of bacteriochlorophylls- $e_1$ ,  $e_2$  and  $e_3$  (indicators of green sulphur bacteria) in box-core subcores from a hypoxic site (PS4 ~27 m water depth) in a chronically hypoxic area on the Louisiana inner continental shelf, SW of the Mississippi River delta. Bchl- $e$  was not detected below 10 cm depth in the core. From Organic Geochemistry, vol. 32, N. Chen, T. S. Bianchi, B. A. McKee, J. M Bland, Historical trends of hypoxia on the Louisiana shelf: applications of pigments as biomarkers, pp. 543–561, 2001, with permission from Elsevier.

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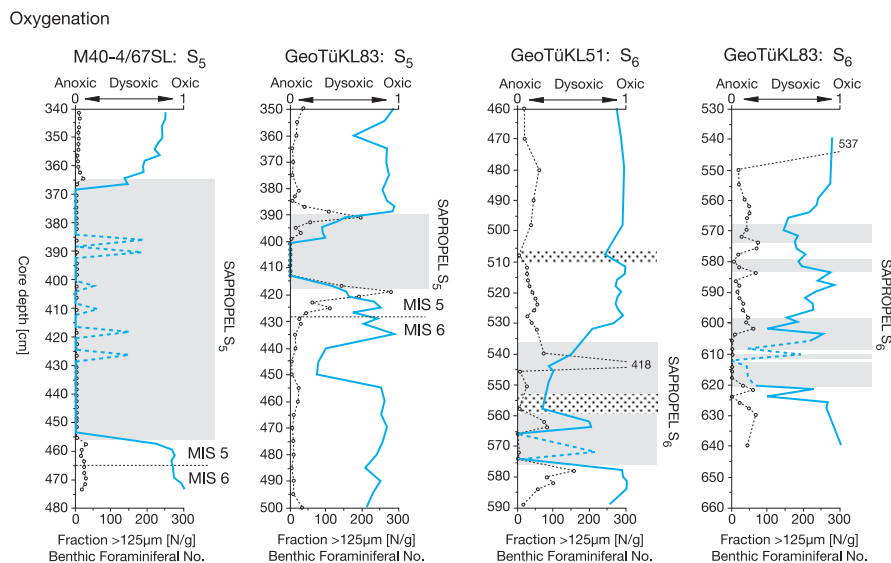
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**Fig. 16.** Application of the “Schmiedl proxy” for bottom-water oxygenation to Mediterranean sapropels. Relative changes of oxygenation (thick blue line) across sapropels S5 and S6 from the lower bathyal western (core M40-4/67SL) and middle bathyal eastern (core GeoTüKL83) Levantine Basin. Oxygen estimates are based on the ratio  $(HO/(HO+LO)+Div)U0.5$ , where HO = the relative abundance of high oxygen indicators (Miliolids - *Articulina tubulosa* + *Cibicidoidea pachydermus* + *Gyroidinoides orbicularis*), LO = relative abundance of low oxygen indicators (*Fursenkoina* spp., *Chilostomella oolina*, *Globobulimina* spp.), and normalized benthic foraminiferal diversity (H(S) normalized). Intervals where oxygen contents are likely to be overestimated due to very low benthic foraminiferal numbers (BFN, number of individuals per g of sediment) are indicated by stippled line. The BFN for the size fraction >125 μm is given for comparison (thin dotted line). Redrawn from Palaeogeography Palaeoclimatology, Palaeoecology, vol. 190, G. Schmiedl, A. Mitsuhele, S. Beck, K. C. Emeis, C. Hemleben, H. Schulz, M. Sperling, S. Weldeab, Benthic foraminiferal record of ecosystem variability in the eastern Mediterranean Sea during times of sapropel S5 and S6 deposition, pp. 139–164, 2003, with permission from Elsevier.

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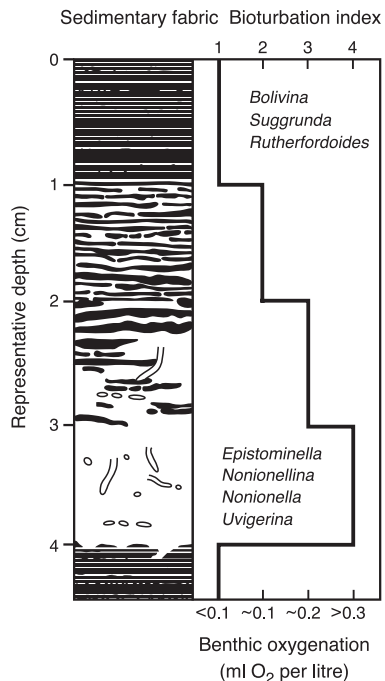
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**Fig. 17.** Bioturbation index (BI) based the development of sediment laminations; application to sediments from Santa Barbara Basin (576-m water depth), NE Pacific. BI value 1 = indicates sediments with distinct, continuous laminations. BI value 2 = diffuse, discontinuous irregular laminations. BI value 3 = slightly bioturbated with either faint, diffuse laminations or a few discrete patches of laminations surrounded by homogenized sediment, including local *Chondrites* burrows. BI value 4 = completely bioturbated, fine-grained sediments with no discernable fabric and commonly containing echinoid spines. Characteristic Foraminifera are listed for two end-member facies. The four facies recognized are believed to correspond approximately to the following oxygen concentrations:  $<0.1 \text{ mL.L}^{-1}$  ( $<4.3 \mu\text{M}$ ; facies 1), macrofaunal excluded;  $0.1 \text{ mL.L}^{-1}$  ( $4.3\text{--}8.6 \mu\text{M}$ ; facies 2), meiofaunal bioturbation causing laminations to become diffuse;  $0.1 - 0.2 \text{ mL.L}^{-1}$  ( $4.3\text{--}8.6 \mu\text{M}$ ; facies 3), enough oxygen to permit partial homogenization of sediment by fauna;  $>0.3 \text{ mL.L}^{-1}$  ( $>13 \mu\text{M}$ ; facies 4), macrofauna present. Redrawn with permission from Macmillan Publishers Ltd. [Nature] (Behl, R. J., and Kennett, J. P.: Brief interstadial events in the Santa Barbara Basin, NE Pacific, during the last 60 kyr, Nature, 379, 243–246, 1996).

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