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The Western Channel Observatory: a century of physical, chemical and biological data compiled from pelagic and benthic habitats in the western English Channel

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Abstract. The Western Channel Observatory (WCO) comprises a series of pelagic, benthic and atmospheric sampling sites within 40 km of Plymouth, UK, that have been sampled by the Plymouth institutes on a regular basis since 1903. This longevity of recording and the high frequency of observations provide a unique combination of data; for example temperature data were first collected in 1903, and the reference station L4, where nearly 400 planktonic taxa have been enumerated, has been sampled on a weekly basis since 1988. While the component datasets have been archived, here we provide the first summary database bringing together a wide suite of the observations. This provides monthly average values of some of the key pelagic and benthic measurements for the inshore site L4 (50°15.00' N, 4°13.02' W; approx. depth 55 m), the offshore site E1 (50°02.00' N, 4°22.00' W; approx. depth 75 m) and the intermediate L5 site (50°10.80' N, 4°18.00' W; approx. depth 58 m). In brief, these data include the following: water temperature (from 1903); macronutrients (from 1934); dissolved inorganic carbon and total alkalinity (from 2008); methane and nitrous oxide (from 2011); chlorophyll *a* (from 1992); high-performance liquid chromatography (HPLC)-derived pigments (from 1999); <20 µm plankton by flow cytometry, including bacteria (8 functional groups from 2007); phytoplankton by microscopy (6 functional groups from 1992); microplankton and mesozooplankton from FlowCam (6 groups from 2012); *Noctiluca* sp. dinoflagellate (from 1997); mesozooplankton by microscopy (8 groups from 1988); *Calanus helgolandicus* egg production rates (from 1992); fish larvae from the Young Fish Trawl survey (4 groups from 1924); benthic macrofauna (4 groups from 2008); demersal fish (19 families from 2008); blue shark, *Prionace glauca* (from 1958); and 16S alpha diversity for sediment and water column (from 2012). These data have varying coverage with respect to time and depth resolution. The metadata tables describe each dataset and provide pointers to the source data and other related Western Channel Observatory datasets and outputs not compiled here. We provide summaries of the main trends in seasonality and some major climate-related shifts that have been revealed over the last century. The data are available from the Data Archive for Seabed Species and Habitats (DASSH):

<https://doi.org/10.17031/645110fb81749> (McEvoy and Atkinson, 2023). Making these data fully accessible and including units of both abundance and biomass will stimulate a variety of uptakes. These may include uses as an educational resource for projects, for models and budgets, for the analysis of seasonality and long-term change in a coupled benthic–pelagic system, or for supporting UK and north-eastern Atlantic policy and management.

1 History

Sustained observations of the marine environment are vital to understand marine ecosystem functioning and climate change responses (O'Brien et al., 2017; Richardson, 2008). Over seasonal timescales, high-resolution observations allow one to develop an understanding of community succession and seasonality (Smyth et al., 2010, 2014); moreover, over multiple decades, they allow us to tease out the effects of local variability and anthropogenic stressors from the longer-term signal of climate change (Edwards and Richardson, 2004; Ratnarajah et al., 2023). Paradoxically, however, many sampling programmes are funded for only 3–4 years and, despite the urgency of understanding climate change responses, time series are threatened at a global scale (Vucetich et al., 2020). This makes it even more important to make data from existing long time series findable, reusable and as well documented as possible.

The Western Channel Observatory (WCO) data contain an unprecedented collection of parameters both in terms of longevity and variety. Investigation of the marine environment in the western English Channel off Plymouth began with the opening of the Marine Biological Association (MBA) laboratory in 1888. Given the importance of the area for the pelagic fishery, the remit focused strongly on research in applied fisheries. Initial studies centred on the eggs and larvae of commercially important fish. With the advent of the International Council for the Exploration of the Sea (ICES) and a growing realization that hydrography had an influence on biological communities, plankton surveys and hydrographical measurements were soon added (Southward et al., 2005; Southward and Roberts, 1987). In the decades that followed, observations were expanded with the creation of stations E1 (50°02.00' N, 4°22.00' W) and L5 (50°10.80' N, 4°18.00' W). Sampling was interrupted during both World War I (1914–1918) and World War II (1939–1945). Funding priorities and organizational changes in the 1970s and 1980s threatened the future of long-term time series, and sampling at L5 and E1 was consequently stopped until 2002. However, in 1988 Plymouth Marine Laboratory (PML) established weekly zooplankton sampling at station L4 (50°15.00' N, 4°13.02' W), with ad hoc funding and no formal support. Sampling for phytoplankton community composition and abundance, egg production, and environmental variables followed from 1992 onwards. The WCO was founded in 2005 to bring these valuable time series together. The WCO provided a platform for a wider array of

parameters to be initiated – for example, the benthic survey from 2007, in situ automated buoys at L4 and E1 (initially supported by Natural Environmental Research Council, NERC, and then the Met Office), the Penlee Point Atmospheric Observatory (PPAO) from 2014, and Smart Sound Plymouth from 2021 (Fig. 1).

The stations around Plymouth, now known as the Western Channel Observatory, have supported major innovative work – for example, pioneering work on plankton as indicators (Russell, 1935), the measurement of nutrients and primary production (Boalch et al., 1978), early work on fatty acids and the importance of food quality for zooplankton (Conover and Corner, 1968; Pond et al., 1996), and the use of molecular biology tools to provide insight into the seasonal dynamics of viral and bacterial plankton (Lindeque, 2023; Gilbert et al., 2009; Schroeder et al., 2003). These works, including the development of intertidal research and data not covered here, can be found in the historical review of Southward et al. (2005). Later special journal issues cover the 20th and 25th anniversaries of regular sampling at L4 and are described in Harris (2010) and Smyth et al. (2015), respectively. We refer the reader to these for the historical context of the observations that we summarize here.

2 The WCO environment

The two main marine stations of L4 and E1 both exhibit strong seasonal signals and are tidally influenced (Smyth et al., 2015). Both become stratified typically after April, continuing through the summer months and lasting until late September. Station L4 is classified as a coastal site and is periodically influenced by flood water discharge from the Tamar and Plym rivers (Rees et al., 2009). However, at a depth of approx. 55 m and 13 km offshore, it is not as prone to localized inshore effects and is classified as “transitionally stratified” (Pingree, 1980). The deeper station E1, 40 km offshore and approx. 75 m deep, is less influenced by coastal water influx and is classified as an open shelf station that is seasonally stratified. The intermediate station, L5, was much sampled in early years and is just west of Eddystone reef. These stations experience classic, albeit highly variable, seasonal production cycles with spring and autumn phytoplankton blooms. Figure 2 compares the key WCO sites in relation to the wider summer pattern of stratification (Fig. 2a) and to the longer trend of climatic cycles across the North Atlantic (Bode, 2023), highlighting a recent phase of intense warming over the last 4 decades (Fig. 2b). These environ-



Figure 1. Location of Western Channel Observatory (WCO) sampling stations.

mental changes and the response of the biota are described more fully in Sect. 5 using plots derived from our summary database.

3 Objectives

The individual datasets of the WCO are valuable, but differing levels of reporting and formatting hamper their use and prevent integration. Many are currently available through data repositories such as the British Oceanographic Data Centre, <https://www.bodc.ac.uk/data/> (last access: 29 November 2023) and the Data Archive for Seabed Species and Habitats, <https://www.dassh.ac.uk/> (last access: 29 November 2023); however, some are lodged with indi-

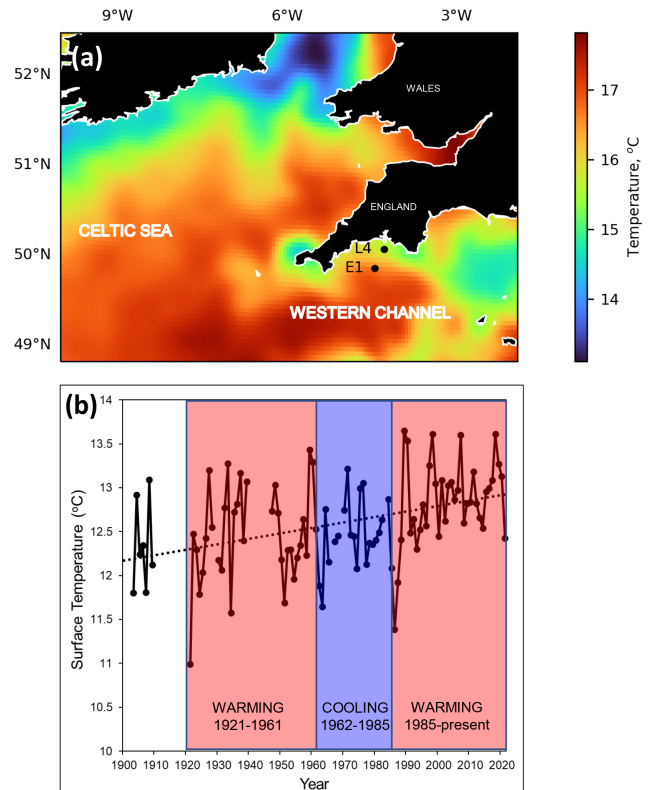


Figure 2. Wider-scale spatial and temporal context for the Western Channel Observatory. Panel (a) displays the wider setting of the L4 and E1 stations in the western English Channel, in relation to summer sea surface temperature July 2016 (Merchant et al., 2019); cold colours represent tidally mixed areas, whereas warm colours represent summer stratified areas with a seasonal summer thermocline. L4 stratifies in summer and is defined as transitionally stratified, whereas E1 is an open shelf and is defined as seasonally stratified (Pingree, 1980). Panel (b) shows the annual surface temperature records at station E1 spanning 1903–2021. Due to missing data in some months of the early years, annual means were calculated here as averages of February, May, August and November. Missing months were interpolated as the mean respective month over the whole time span. Years with more than 2 of the 4 missing months are not plotted here. The dotted line is the least squares linear regression over the whole time span. Three thermal epochs are coloured here based on the interpretation of Southward et al. (2005). They defined periods of warming from 1921 to 1961, followed by a cooling era from 1962 to 1985 and then a warming period from 1986 onwards.

vidual scientists. To improve their overall utility, the various component datasets need to be brought together into a single format. We have done this here for the first time; however, to make this project tractable, we have summarized the core datasets as monthly averages, as well as for broad functional groups. This level of resolution (coarser than some of the measurements, which can be weekly and for individual species) was chosen as a first step to allow timely completion of this initiative and to provide a summary database that

combines many diverse data sources. This data paper combines most of the key variables that have good seasonal or longer-term coverage in a single spreadsheet (Fig. 3). Specialists who wish to access the underlying high-resolution observations, who wish to access data for individual species, who require the most recent data available, or who require other datasets not summarized here are directed to our WCO data catalogue: <https://www.westernchannelobservatory.org.uk/data.php>. (last access: 29 November 2023). This catalogue provides sampling details, DOIs of the most recent versions and points of contact for specific datasets. Additional information is also available in Tables A1 and A2.

This data paper is aimed towards scientists who may not need weekly resolution or species-specific data but who wish to compare the monthly averaged physical, chemical and biological data. Biological data are provided in units of both abundance and biomass in order to enhance their utility for modelling. We have also made the spreadsheets as user-friendly and simple as possible to be of help as an educational resource at the undergraduate level. This data paper describes the database (Sect. 4); illustrates its utility to examine seasonality and longer-term trends while also summarizing previous work on these topics (Sect. 5); provides a broader-scale context for the WCO (Sect. 6); and finally provides practical advice on the strengths, limitations, and how to use and cite this database (Sects. 7 and 8).

4 Data processing

This paper consolidates 22 individual and diverse datasets using monthly averages. Data with comprehensive seasonal coverage that span at least 2 years are included. Detailed information on sampling and analysis protocols as well as data coverage can be found in Tables A1 and A2. It is essential to read the appendices before extracting data to avoid errors, for example, in distinguishing between zeros and absent data. A zero represents a parameter that was either looked for and not found (for plankton data) or was below the detection limit (nutrient data). A blank cell, by contrast, indicates there are no data available for that particular month. Most biotic data are reported both in units of abundance and biomass. The smallest plankton ($< 20 \mu\text{m}$) measured by flow cytometry are an exception. These use fixed conversion factors for the whole functional group, and they have multiple groups and depths. Therefore to remove the complexity of having many data fields that are simple multiples of others, these are reported only as abundance per millilitre. Median cell diameters are provided, thereby enabling the estimation of biomass based on the volume of a sphere and carbon values from the literature (Table 1). Median cell diameters were derived by collecting seawater samples, filtering them sequentially through a series of membrane filters, analysing the filtrates by flow cytometry, and plotting the percentage of cells remaining as a percentage of unfiltered seawater against fil-

ter pore size (Burkill et al., 1993). The FlowCam analysis of the $63 \mu\text{m}$ mesh plankton net reports biomass only. Here, along with the microscopy analysis of phytoplankton preserved in Lugol's solution and formalin, the biomass estimation is based on mean cell dimensions and taxon-specific biovolumes (Table A1). Biomass calculations for the mesozooplankton are based on measured body lengths of material from L4 and applied literature values of length–mass regressions to convert values to individual biomass. These were then multiplied by numerical abundances to derive biomass densities (McEvoy et al., 2022b). The benthic fauna biomass data are derived from blotted wet weight measured in the lab, and Cephalopoda and demersal fish families' biomass are derived by wet weight on board.

5 Results and discussion

In this section, we briefly showcase some of the key datasets, by outlining the seasonality and environmental variability, illustrating the coverage of all of the component data series at L4 (Fig. 4) and at E1 and L5 (Fig. 5). We then show selected examples of the data time series' coverage, including longer-term trends at L4 (Fig. 6) and at E1 and L5 (Fig. 7). A few other key example results are shown, including the *Calanus* spp. egg production experiments (Fig. 8) and the time–depth resolution of sampling for carbonate chemistry (Fig. 9).

5.1 Overall seasonality: L4 pelagic system

The high-resolution sampling of multiple parameters at L4 makes it an ideal site for improving the understanding of the drivers of seasonality. Figure 4 summarizes some of the key aspects of this seasonality. Briefly, L4 is a transitionally stratified site (Pingree, 1980) that is typically stratified from around May to September with surface temperatures ranging from about 9°C in March to around 16°C in August. This stratification cycle drives much of the pelagic dynamics with nutrient (especially nitrate) depletion to near-limiting levels in the upper water column during the stratified period as well as progressive reductions in dissolved inorganic carbon (DIC), methane and nitrous oxide typically until about August (Kitidis et al., 2012).

The combination of nutrients, light and grazing causes the conditions for a “classic” temperate shelf sea production cycle (Irigoien et al., 2005). Thus, there is typically a spring bloom around April–May dominated by diatoms and the prymnesiophyte *Phaeocystis*, followed by a dinoflagellate bloom in late summer and often diatoms in autumn. Importantly, however, the monthly mean values of the phytoplankton functional groups in Fig. 4 disguise the substantial interannual variability in their magnitude or floristic composition over the time series (Widdicombe et al., 2010). The pico- and nano-fractions follow slightly different dynamics, with the highest biomass values building up in the summer stratified period with maxima often in August–September

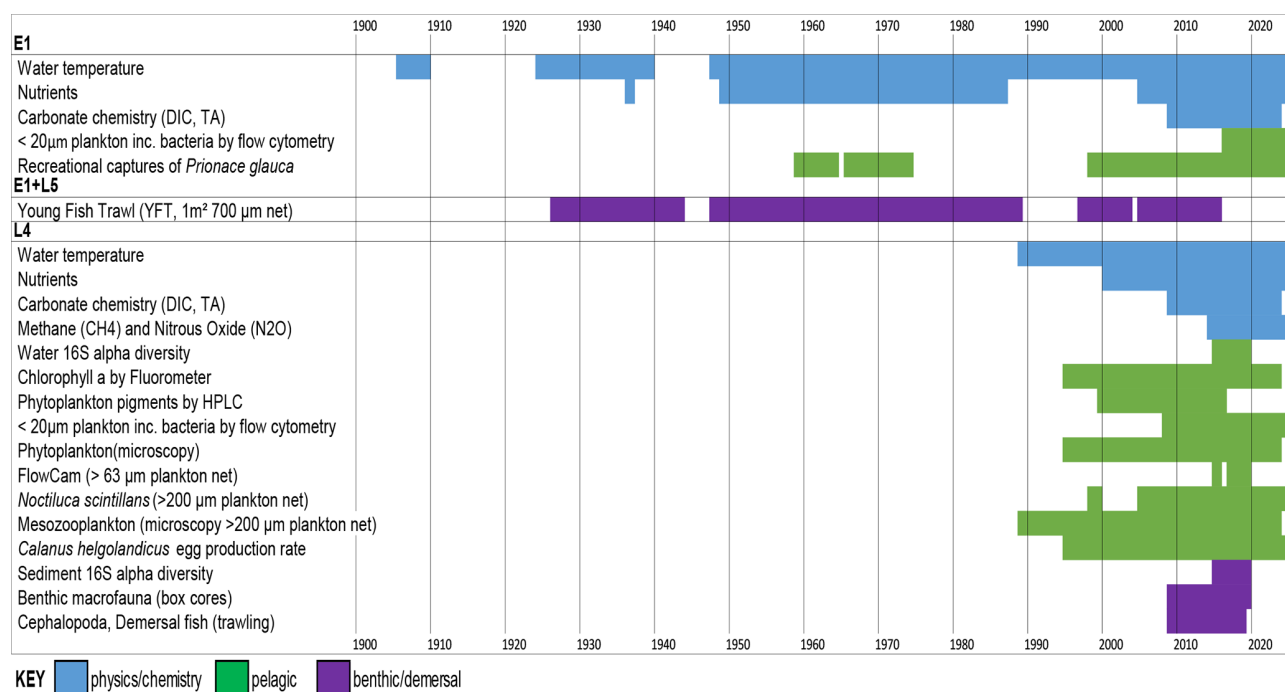


Figure 3. Data combined in this data paper, showing the time span of each.

Table 1. Median cell diameters for plankton groups quantified by flow cytometry.

Group	<i>Synechococcus</i> sp. ^a	Picoeukaryotes ^a	Nanoeukaryotes ^a	Coccolithophores ^b	Cryptophytes ^a	Bacteria ^c
Median diameter (μm) \pm 1 SD	1.72 \pm 0.70	1.83 \pm 0.58	5.40 \pm 2.04	7.68 \pm 0.89	5.48 \pm 1.33	–
Spherical volume (μm^3)	2.66	3.20	82.50	236.87	86.36	–
Carbon per cell (pg)	0.59 ^d	0.70 ^d	18.15 ^d	67.51 ^e	19.00 ^d	0.019

^a Information from Station L4, approximately monthly over an annual cycle from 2013 to 2014 (unpublished). ^b Information from the Celtic Sea, April 2002 (unpublished). Information from ^c Heywood et al. (2006). ^d Carbon conversion factor of 0.22 pg C μm^{-3} (Booth, 1988). ^e Carbon conversion factor 0.285 pg C μm^{-3} (unpublished).

(Tarran and Bruun, 2015). The FlowCam biomass estimates based on 63 μm mesh, full-depth net hauls show the contributions of copepod nauplii and the larger diatoms, dinoflagellates and ciliates, some of which are not statistically quantified in the counts based on samples preserved with Lugol's solution due to their rarity. Conversely, the copepod nauplii are too small to be quantitatively retained by the 200 μm mesozooplankton WP2 net. Rare seasonal profile data of the copepod *Oithona* spp. based on bottle sampling are provided by Cornwell et al. (2020).

The mesozooplankton grazers from the full-depth 200 μm net hauls tend to increase substantially as early as March, before the spring bloom (Atkinson et al., 2015). The peak is typically in the early summer, dominated by copepods, both numerically (Eloire et al., 2010) and in terms of biomass, with a substantial contribution from meroplankton in spring (Highfield et al., 2010). More predatory taxa (often gelatinous or semi-gelatinous forms such as chaetognaths) then become important later in summer. The egg production rate of *Calanus helgolandicus* has, for most of the time series, been highest in the April–June spring bloom months

(Irigoien and Harris, 2003; Maud et al., 2015, 2018), although (as described in Sect. 5.6) this is changing. This copepod species, alongside other zooplankton such as appendicularians (López-Urrutia et al., 2003), decapods (Fileman et al., 2014), bivalve larvae (Lindeque et al., 2015) and *Oithona similis* (Castellani et al., 2016; Cornwell et al., 2018, 2020), has been the focus of a series of detailed studies at L4 (Bonnet et al., 2005; Hirst et al., 2007; Irigoien and Harris, 2003).

While Fig. 4 describes a classic textbook shelf sea production cycle (Kjørboe, 2009), a wide suite of alternative mechanisms have been proposed to drive plankton seasonality (Atkinson et al., 2018). These include the following: the roles of phytoplankton photophysiology (Edwards et al., 2013; Polimene et al., 2014); net heat flux (Smyth et al., 2014); variable temperature dependence of phenology (Mackas et al., 2012; Atkinson et al., 2015); mortality-controlled zooplankton phenology (Irigoien and Harris, 2003; Cornwell et al., 2018); and various predator–prey dynamic models invoking stoichiometry (Polimene et al., 2015), grazing loopholes (Irigoien et al., 2005), grazer traits (Sailley et al., 2015), and the coupling of predator and prey traits (Kenitz et al., 2017).

While these mechanisms are not necessarily mutually exclusive, the high-resolution sampling of the whole food web over multiple years provides a good test bed for numerical and conceptual models of the factors driving seasonality.

5.2 Overall seasonality: L4 benthic system

In contrast to the plankton, the benthic and demersal taxa have more varied seasonal dynamics. Macrofauna biomass is dominated by suspension feeders at L4 with similar biomass for most of the year (Fig. 4) except for low values in early winter. Potentially reflecting seasonal variation in water column food supply, species richness of infauna peaks throughout the summer in surface sediments and is lowest in late autumn. Higher numbers of species are also found in deeper sediment layers during warmer months, with the community seemingly shallowing over winter (Queirós et al., 2019). An assessment of particulate carbon sources to the seabed at L4 (Queirós et al., 2019) also suggested that fauna in shallow sediment layers exhibit strong signals of suspension and deposit feeding reliant on planktonic food sources, with carnivory increasing with sediment depth and reliance on water column food diminishing in tandem. Demersal fish are dominated by gadoids with seasonal minima in both December and March–April. In the benthos, there is no distinguishable seasonal signature to prokaryote diversity, in contrast to the water column, where it is lowest in the summer months.

While the detail of the seasonality of L4's pelagic and benthic component differs, there is strong connectivity between the pelagic and benthic systems, even during stratified periods. This is illustrated in the peaks and troughs of diversity linked to the availability of food sources (see seasonal productivity peaks in Fig. 4e–j; Queirós et al., 2019; Tait et al., 2015; Talbot et al., 2019) and is also reflected by pigment data and stable isotopic signatures of both DIC and particulate organic carbon and nitrogen (Queirós et al., 2019; Tait et al., 2015). Benthic–pelagic connectivity has been found to be seasonally variable in terms of the origin of the suspended and dissolved matter fluctuating between the two ecospheres as well as in terms of the dominant flux directions (Queirós et al., 2015; Rühl et al., 2020; Tait et al., 2015). Strong seasonality is also observed in the dynamics of ecosystem processes mediated by macrofauna in L4 sediments, i.e. bioturbation and bio irrigation (Kristensen et al., 2012), which have strong mediation effects on the rates of biogeochemical processes at the sediment–water interface, such as community respiration, and net carbon sequestration (Queirós et al., 2015, 2019).

Broader analyses of the seafloor time series at L4 have also demonstrated that these dynamics are highly variable on an interannual basis, with the effects of extreme events being particularly important (Rühl et al., 2021). Net vertical flux directions of suspended and particulate matter vary throughout the year, switching in direction and respective importance for the overall flow of matter throughout the system. In summary, the benthic system at L4 has not been as intensively

sampled as the pelagic, but the site still provides an excellent opportunity to better understand benthic–pelagic coupling in a shallow shelf sea.

5.3 Overall seasonality: E1 and L5 in comparison to L4

Figure 5 summarizes the data available for the E1 and L5, which are sites located further offshore than L4. All measurements except those from the Young Fish Trawl and the shark catch data pertain to the E1 site. The Young Fish Trawl data are from site E1 and L5 combined. The shark data are from angling vessels from Looe and within 10 nmi of E1.

The E1 site is more strongly stratified than L4, as evidenced by slightly higher surface temperatures and a larger summer temperature difference between the surface and depth. As the site is further offshore than L4 and receives less riverine nutrient input from the Tamar and Plym rivers, macronutrient concentrations are more severely limiting in summer and, indeed, iron stress has been suggested in some seasons (Schmidt et al., 2020). This is also reflected in the stronger reduction in dissolved inorganic carbon (DIC) during the stratified period, resulting in an average seasonal amplitude of $83 \mu\text{mol kg}^{-1}$ at E1 compared with around $55 \mu\text{mol kg}^{-1}$ at L4. Total alkalinity (TA), in contrast, shows little seasonal cycle at E1 (average seasonal amplitude of $29 \mu\text{mol kg}^{-1}$), compared with a slight increase in spring at L4 (average seasonal amplitude of $40 \mu\text{mol kg}^{-1}$). Both L4 and E1 show a seasonal pattern of seawater CO_2 undersaturation between January and August, followed by supersaturation in September and October, returning to near equilibrium with the atmosphere for the remainder of the year (Kitidis et al., 2012). The subsurface chlorophyll-*a* maxima has been shown to be important for controlling carbon fluxes at these sites as well as for the mixing of freshwater, which is evidenced by the difference between L4 and E1 conditions (Kitidis et al., 2012). The flow cytometry data reflect this (Fig. 5), with an increased contribution of coccolithophores compared with L4, albeit with pronounced interannual variability and large blooms in some years but not others.

Although phytoplankton and zooplankton samples are currently collected at E1, we have not summarized them in this paper because the available time series of data do not cover as long a period as L4. However, a summary of phyto- and zooplankton seasonality at E1 is presented and compared with that of L4 by Djeghri et al. (2018). These authors showed that mesozooplankton biomass at E1 is lower than at L4 and at deeper, offshore sites in the Celtic Sea (Giering et al., 2019). Also, in the context of these Celtic sea stations, Schmidt et al. (2020) examined nutrient dynamics at E1 in relation to pico- and nanoplankton and found that late-season dominance of the picocyanobacteria *Synechococcus* (including intense blooms in some years) tended to follow summers of particularly severe nitrate stress.

Data compiled here from the Young Fish Trawl (Fig. 5d) show strong summer and autumn increases in sardine (here-

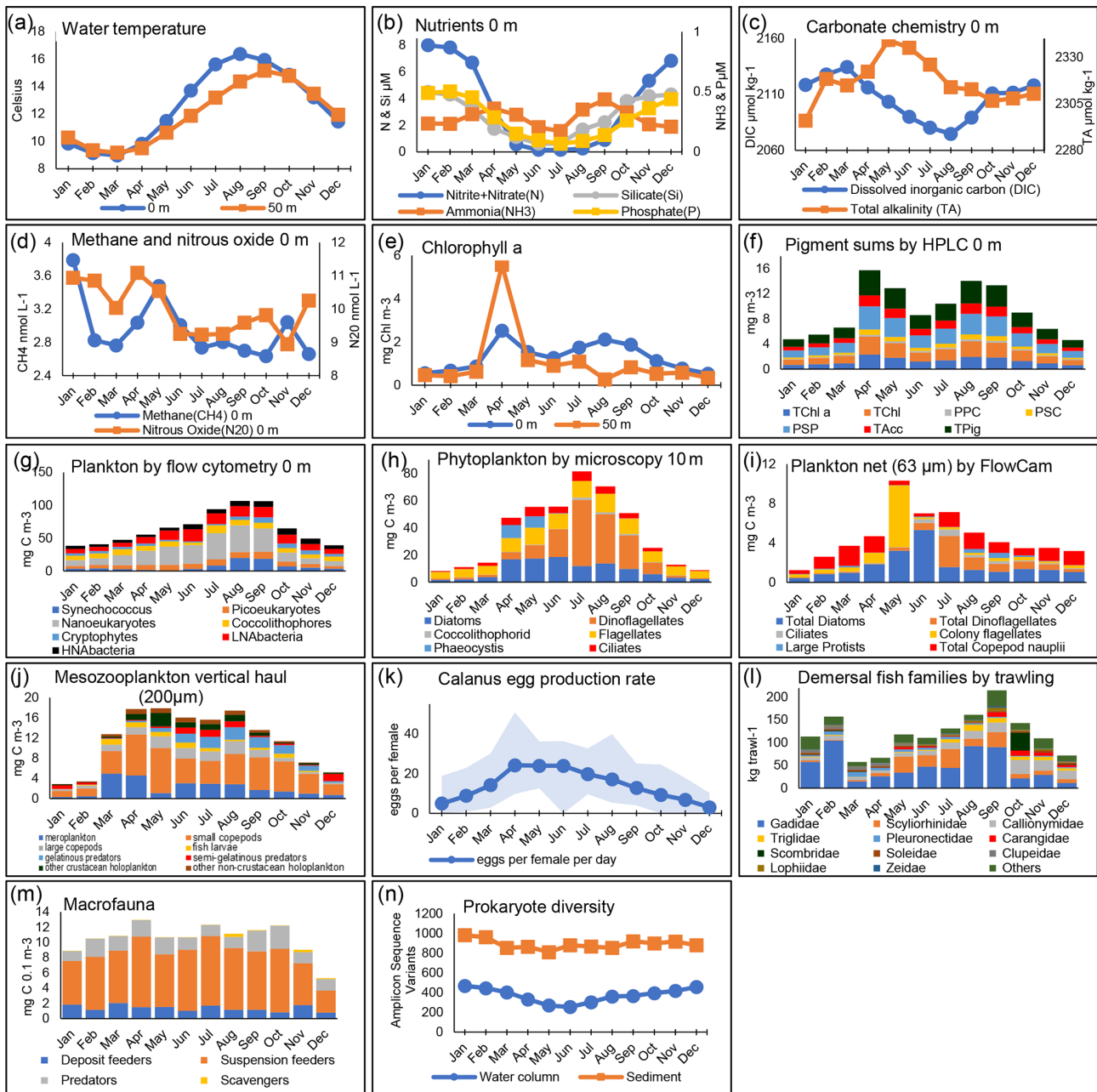


Figure 4. Seasonal patterns at station L4. The panels show monthly mean values calculated across all years of available data, presented for the surface (0 m) unless otherwise stated. For explanations of all data fields, see Table A1. In panel (f), the categories show different pigment sums (see Table A1). Panel (g) presents the biomass of plankton using flow cytometry derived using conversion factors from Table 1. In panel (k), the *Calanus* spp. females are collected from net samples. In panel (l), the fish families plotted here are for the 11 top-ranking groups based on annual mean biomass, with remaining groups (including Cephalopoda) summed here as “others”.

after called by their more locally common name “pilchard”) eggs. The later autumn spawning period has become more dominant in recent years (Coombs et al., 2010). Fish eggs of other species, by contrast, are more abundant in early spring. *Calanus* spp. are not quantitatively sampled by the 700 μm mesh of the Young Fish Trawl, but Fig. 5d shows an increase in mid-summer, which is in line with Fig. 4j, where

the biomass of the large-copepod fraction is dominated by *Calanus* spp. This genus comprises mainly *C. helgolandicus* in this area (Lindeque et al., 2013; Maud et al., 2015, 2018), which are important food for pelagic fish. Success in catching blue sharks (*Prionace glauca*) increases rapidly until late summer.

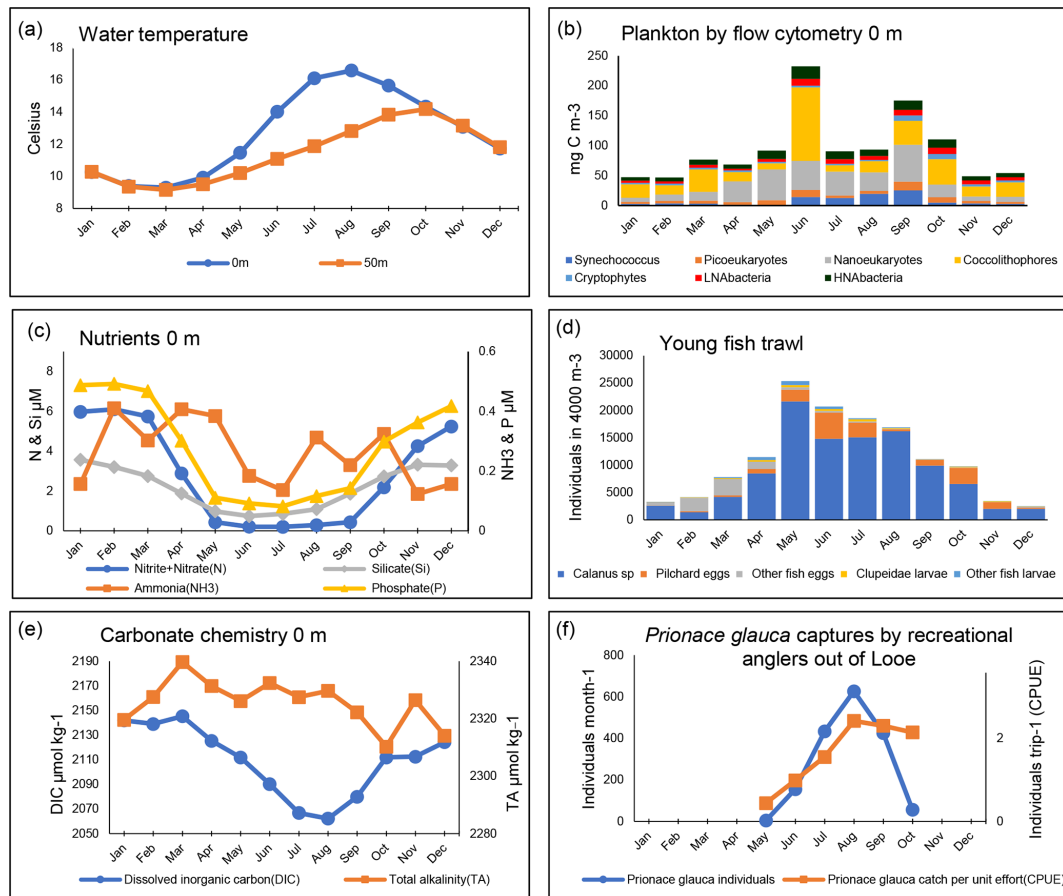


Figure 5. Seasonal patterns at station E1 and L5 (1903–2021): monthly mean values calculated across all years of available data. Illustration shows surface (0 m) data unless stated. For explanations of all data fields, see Table A2. Panel (b) shows the biomass of plankton by flow cytometry derived using conversion factors from Table 1. Panel (d) presents a Young Fish Trawl oblique tow to a depth of 5 m above the seabed.

5.4 Annual time trends: L4

The regular weekly resolution measurements at L4 cover the most recent era of rapid warming (Fig. 2), and both the sampling intensity and the number of planktonic taxa measured allow observation of systematic change and its driving factors. We cannot review all of the literature on change here; therefore, instead, Fig. 6 illustrates some of the key trends.

Because stratification is such a major factor driving seasonality at the WCO, Fig. 6 compares trends separately between the most stratified quarter of the year with the lowest average nutrient concentrations (May–August) and the rest of the year. The temperature increase during the warm, stratified period over the last 30 years is more pronounced than in the other months and is well over 1 °C. The sharp rise in temperature at this time of year coincides with a major decline in nitrate concentrations and DIC, pointing to the effects of enhanced stratification retarding nutrient and carbon supply (Fig. 6a).

Figure 6b compares the trends for surface chlorophyll-*a* concentrations and the biomass-dominant functional groups

of phytoplankton, which together dominate the estimated biomass of cells counted in water samples from 10 m depth preserved in Lugol’s solution, namely, diatoms, dinoflagellates and nanoflagellates (ca. 15 μm). Flagellates (ca. 2–15 μm) are also counted more quantitatively and with full-water-column resolution by flow cytometry (since 2007), and the component fluorescing and non-fluorescing groups (termed nanoflagellates and heterotrophic flagellates, respectively) are major contributors to community dynamics (Atkinson et al., 2021; Tarran and Bruun, 2015). The larger phytoplankton, namely, diatoms, dinoflagellates and the subset of larger nanoflagellates, decline strongly during the summer stratified period, with the chlorophyll-*a* concentration declining by about 50 % overall over 30 years.

In parallel with these summer declines in phytoplankton, there have been declines at the same time of year in the crustacean mesozooplankton functional groups, namely, large and small copepods, other crustacean holoplankton (dominated by *Podon* spp. and *Evadne* spp.) and fish larvae (Fig. 6c). The large-copepod category (defined here as

an adult total body length over 2 mm) is strongly dominated by *Calanus helgolandicus*. From 1988 to around 2015, annual abundances of *C. helgolandicus* were fairly stable, only oscillating about 4-fold between years (Atkinson et al., 2015; Maud et al., 2015). However, in recent years, numbers in summer have started to decline substantially, making it hard to obtain sufficient individuals for egg production experiments. This sudden shift supports the concept of abrupt step changes that “reorganize” assemblages both at this site (Reygondeau et al., 2015) and more widely (Bode, 2023).

Outside of summer, these declines in the crustacean groups were not seen or were not so prevalent in other months, and there was only a weak phenological shift observed at L4 (Atkinson et al., 2015; Uriarte et al., 2021), which does not explain the differential trends between the summer and the rest of the year.

Other taxa, by contrast, have tended to increase at L4. Only a minority of major crustaceans have shown signs of an increase, notably the more carnivorous, late-summer copepod *Centropages typicus* (Corona et al., 2021). The main increases are among meroplankton taxa, fine-mesh filter feeders such as appendicularians (which dominate the “other non-crustacean zooplankton” category) as well as the gelatinous predators (dominated by cnidarians) and semi-gelatinous predators (dominated by chaetognaths). Together, this suggests a shift in the balance of the mesozooplankton, from copepod domination towards a diversity of mero- and holoplankton that are fine-particle feeders, more gelatinous or more carnivorous.

These trends seen at L4 conform to much wider-scale, long-term trends that are coherent across the north-eastern Atlantic and north-western European shelf. They are even broadly similar to those at the Naples Bay monitoring site over a similar timescale (Mazzocchi et al., 2023). As an example, the meroplankton increase is widespread across the north-western European shelf and north-eastern Atlantic (Bedford et al., 2020; Holland et al., 2023), and the trends seen at L4 in summer generally resemble the wider trends seen, particularly to the west of the UK (Schmidt et al., 2020; Holland et al., 2023). As a cause, one recent hypothesis involves a bottom-up mechanism whereby increased summer nutrient stress favours pico-size cells and cyanobacteria such as *Synechococcus* which have a low polyunsaturated fatty acid (PUFA) content and poor nutritional quality. This was suggested to cause a mismatch with the energy demands of crustacean zooplankton grazers at the warmest time of year when their metabolic rates are highest (Schmidt et al., 2020). However, this does not explain the increasing abundance of carnivores or meroplankton, and to fully understand the causes of these trends, time series such as these need to be networked with those from other sites (O’Brien et al., 2017).

5.5 Overall time trends: E1

The E1 site has the longest history of measurements at the WCO and has exemplified progressive technological advances in measuring macronutrients and primary production (Southward et al., 2005). It has also been a testing ground for theories on how climatic variability impacts nutrients and, thus, phytoplankton, cascading up to fish. The “Russell cycle” (Cushing, 1976) was a good example of these progressive ideas which suggested that reduced Atlantic inflows of limiting nutrients (Kemp, 1938) reduced primary production and caused a shift from a herring-dominated ecosystem in the first few decades of last century to one dominated by pilchards in the mid-1930s. Some of these ideas about the mechanism of the Russell cycle have since been revised (Southward et al., 2005); nevertheless, a degree of cyclicity in temperature and nutrients is clear in Fig. 7a, and this is manifested in major cycles of higher trophic levels (Fig. 7b, c).

A major problem when interpreting these long time series is the attribution of cause from correlative-type analyses (Bedford et al., 2020). However, the rapid rise in pilchards after the collapse of the herring fishery may be due to a combination of overfishing and climatic factors (Southward et al., 2005). Similarly, the intensive industrialized pelagic fishing for mackerel and pilchards in the 1970s and its sudden collapse due to overfishing in the 1980s has unknown effects on the trajectories of fish illustrated in Fig. 7c.

5.6 *Calanus* spp. egg production experiments at L4

Although rate measurements have periodically been made at the WCO, such as primary production (Barnes et al., 2015) and grazing (Bautista and Harris, 1992; Fileman et al., 2010), measurements of the *Calanus* spp. egg production rate have been made fairly consistently since 1992 (Fig. 8). This makes it one of the longest zooplankton production time series of its kind (Harris, 2010) and offers valuable insight into both food quantity and quality for grazers and the population dynamics of *Calanus helgolandicus* (Green et al., 1993; Irigoien et al., 2000). While the original weekly data are archived at the British Oceanographic Data Centre (BODC), the monthly values averaged here (i.e. a mean of the weekly average rates) provide a good seasonal and long-term comparison with the respective monthly average water temperature and functional groups of phytoplankton.

A series of publications have used these *Calanus* spp. egg production data and have supplied extra supporting information. Examples include the importance of nutrition in *Calanus* spp. reproduction (Pond et al., 1996); the role of stratification, egg sinking and mortality (Irigoien and Harris, 2003); and the interaction among temperature, food and predators on population dynamics (Bonnet et al., 2005; Cornwell et al., 2018; Maud et al., 2015, 2018). Long-term changes in the phenology and rates of *Calanus helgolandicus*

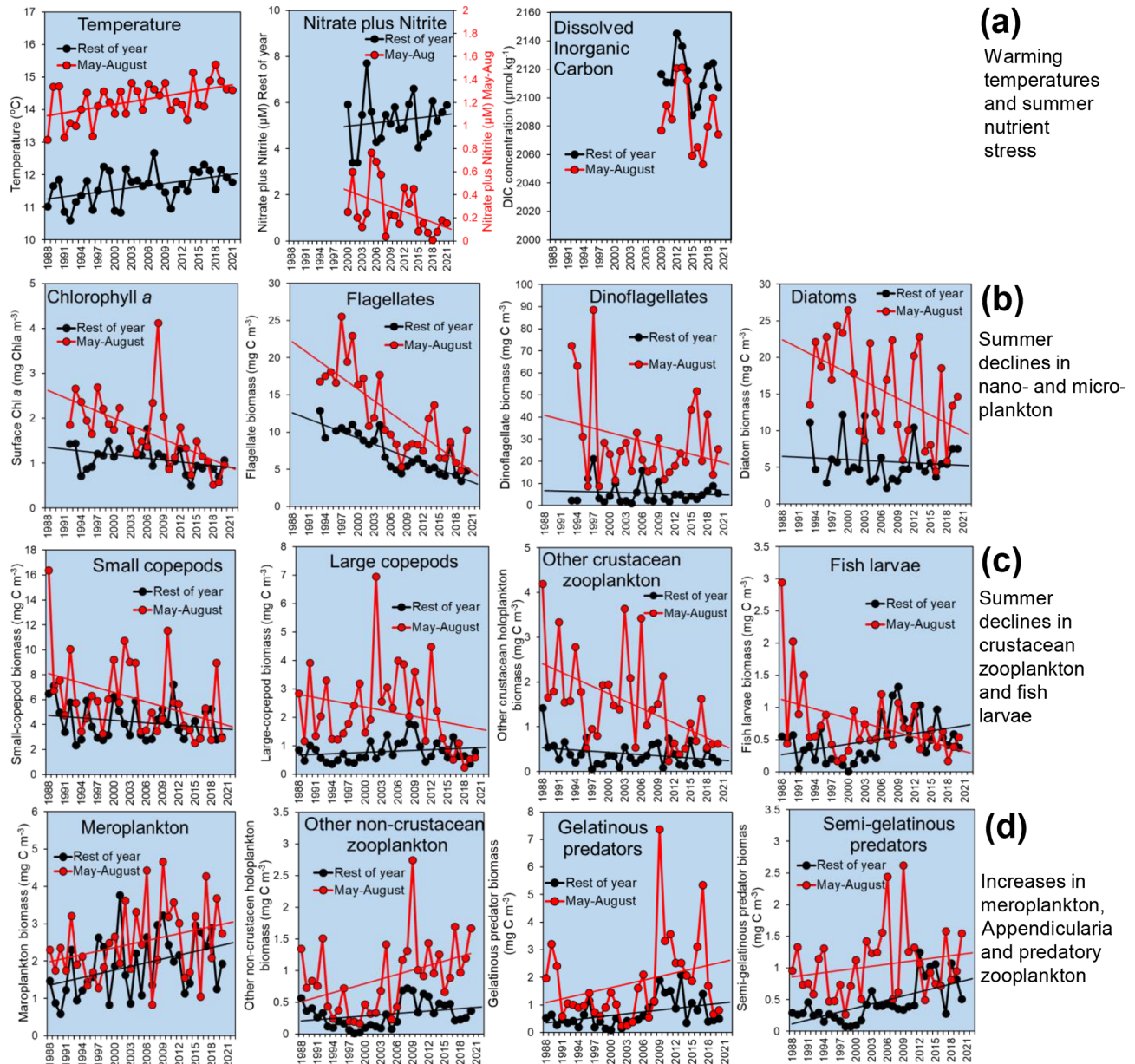


Figure 6. Example plots of L4 time series showing major changes. Each data point represents an annual mean of the monthly values, averaging the main summer stratified period (May–August, red) and the rest of the year (black): **(a)** temperature, surface nitrate and nitrite, and DIC; **(b)** phytoplankton, including surface chlorophyll-*a* (Chl-*a*) concentrations and biomass of the dominant phytoplankton functional groups counted in water samples from 10 m depth preserved in Lugol’s solution; **(c)** mesozooplankton of the “classical food web” that have declined in summer, namely, crustaceans (dominated by Copepoda) and fish larvae; and **(d)** biomass of key taxa that have increased, including meroplankton, other holoplankton (dominated by Appendicularia), semi-gelatinous predators (dominated by Chaetognatha) and gelatinous predators (dominated by Cnidaria). Missing months within these time series have been replaced by overall long-term mean values for the missing month. Trend lines (drawn for data with > 20-year time spans) are illustrative linear regressions and do not necessarily imply statistical significance.

cus egg production have not been studied recently, but these show some interesting patterns (Fig. 8). Until roughly 2006, there was a clear maximum in the egg production per female during the spring bloom months, but this moved later into the summer and autumn months over the following 15 years,

with a general decline in the maximum rates. In the last couple of years, however, there have been suggestions that a pattern of high egg output in spring may be reasserting itself.

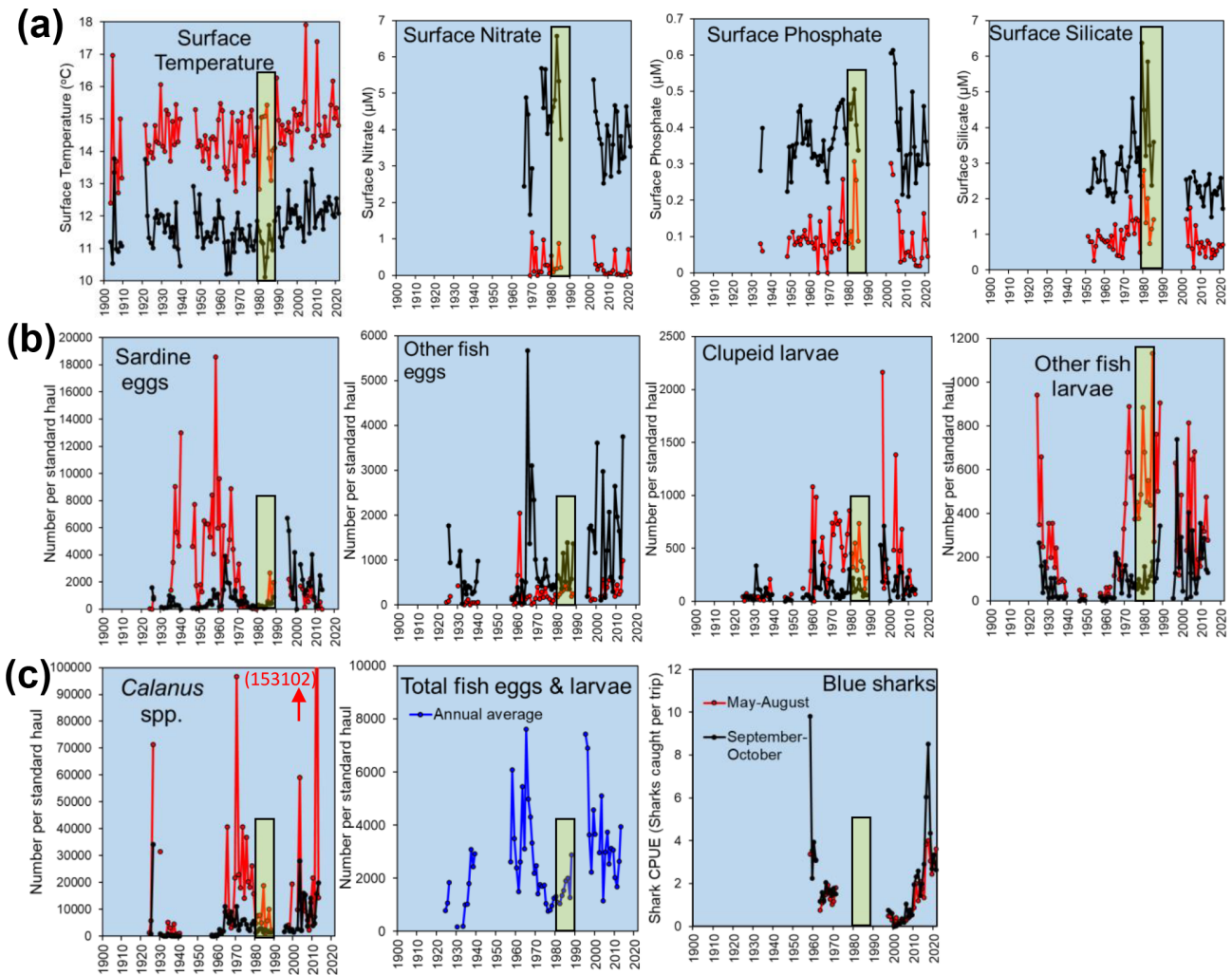


Figure 7. Examples of long time series from stations L5 and E1 from the Western Channel Observatory. Points represent averages of monthly values from the main stratified months (May–August, red) and the rest of the year (black): **(a)** temperature and nutrients; **(b)** catches per “standard haul” of the Young Fish Trawl, showing four categories of fish eggs and larvae with standard haul volumes, standardized to a filtration volume of 4000 m^3 ; and **(c)** respective panels showing the catches of *Calanus* spp. from the Young Fish Trawl, the annual mean values for total fish caught in the Young Fish Trawl (all four categories in panel **(b)** but screened such that records with absent data for any of the four categories were removed) and annual means calculated based on a mean of all available monthly data, and the Blue shark (*Prionace glauca*) catch per unit effort (mean catch per trip from angling boats from Looe fishing within 10 nmi of E1). Yellow bars mark the 1980s for ease of cross-referencing between plots. The 1980s marked major changes, including the onset of rapid warming, the end of a period of intense pelagic trawling off Plymouth and the cessation of funding for many monitoring programmes (including the WCO).

5.7 Carbonate chemistry measurements

Over a decade’s worth of data are now available for the carbonate chemistry at both L4 and E1, which have been used to provide evidence for a number of assessments relating to ocean acidification, including the Convention for the Protection of the Marine Environment of the North-East Atlantic (OSPAR) QSR2023 Ocean Acidification assessment (McGovern et al., 2023) and the recent Marine Climatic Change Impacts Partnership (MCCIP) 2022 Status on Ocean Acidification around the UK and Ireland (Findlay et al., 2022). The data series are one of just two stations’ time series that

record carbonate chemistry parameters in the UK at this frequency and are submitted as part of the UK’s contribution to the UN Sustainable Development Goal Indicator 14.3.1 for ocean acidification.

Over the full time series for L4, there has been an overall decline in both total alkalinity (TA) and dissolved inorganic carbon (DIC), which has resulted in an increase in CO_2 fugacity ($f\text{CO}_2$) of $6.4\ \mu\text{atm yr}^{-1}$ and a decrease in pH of $-0.0126 \pm 0.0022\ \text{yr}^{-1}$. If the 2021 data are excluded, the decrease in pH is slightly slower at $-0.006\ \text{yr}^{-1}$ (Findlay et al., 2022), demonstrating a significant lowering of pH in

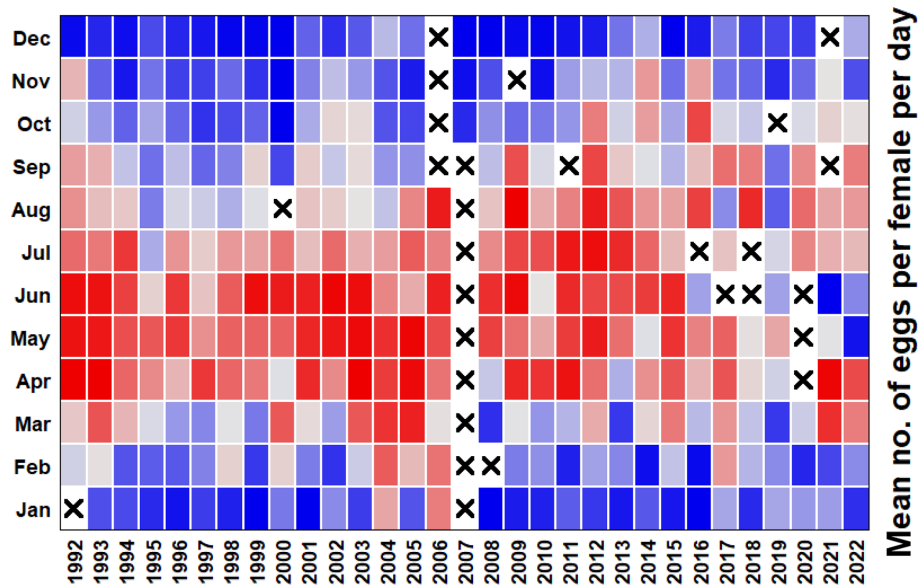


Figure 8. Heatmap of the mean egg production rate of *Calanus helgolandicus*. Pending suitable weather conditions for L4 sampling and enough adult females to incubate, experiments were run weekly: 25 female *Calanus helgolandicus* were incubated for 1 d in egg production chambers, and eggs were then counted. Red cells represent the highest egg production, blue cells represent the lowest egg production, and cells with crosses denote that no data were available.

2021, a result of a decrease in alkalinity and a large reduction in salinity (monthly analysis from same sampling points as carbonate chemistry gives a decline over the time series of -0.01 ± 0.005 psu yr^{-1} ; Gonzalez-Pola et al., 2022). This rate of pH decline is faster than rates observed in the open ocean but is similar to rates found in other near-shore locations off the French coast in the western English Channel and Bay of Biscay (McGovern et al., 2023). Interestingly, the aragonite saturation state shows no significant trend at L4, most likely resulting from the concomitant decline in both DIC and TA as well as the high level of variability in TA caused by organic alkalinity inputs from local rivers. At station E1, there has been a greater decline in DIC, a similar decline in TA and a slightly slower decline in pH (when including 2021 data, this value is -0.008 ± 0.0022 yr^{-1}).

Since autumn 2017, additional water column measurements have been taken at L4, providing a profile view of the carbon dynamics. As a case study, we show the profiles between 2018 and 2020 here, including both 2018 and 2020 (Fig. 9). There is a clear relationship between the in situ density anomaly (σ_t) and both DIC ($\text{DIC} (\mu\text{mol kg}^{-1}) = 38.79\sigma_t + 1086$, $r = 0.7184$, $n = 144$, $p < 0.0001$) and TA ($\text{TA} (\mu\text{mol kg}^{-1}) = 18.36\sigma_t + 1814$, $r = 0.4118$, $n = 144$, $p < 0.0001$). The in situ density anomaly at L4 is primarily driven by temperature, although salinity is important for the dilution of carbonate parameters at this site. The σ_t represents the seasonal cycle of winter mixing followed by stratification through the spring and summer and breakdown of stratification again in the autumn. Both DIC and TA are generally at similar concentrations throughout the water column,

with DIC being reduced in the upper mixed layer during stratification and corresponding to the subsurface chlorophyll blooms (Fig. 9). TA has much higher variability as a result of organic alkalinity contributions.

Data on suspended matter and particulate carbon compounds have also been collected at the WCO during different times over the years. As shown in Rühl et al. (2021), the concentration of particulate organic carbon (POC) at a depth of 10 m at L4, measured between 2013 and 2017, is highly seasonally variable but decreases overall over the 4-year period. It is unclear whether this is part of a more long-term cyclical pattern or a true temporal trend in the data. Variability in the particulate suspended matter concentration in general is less seasonal and does not conform to any clear trend throughout the same time period (Rühl et al., 2021).

6 Wider context

6.1 Numerical modelling

Long-term time series like the ones reported here have been paramount in shaping our understanding of biogeochemical cycling and plankton dynamics (Benway et al., 2019). Not only have they provided the necessary data consistency to generate hypotheses to progress our understanding of marine ecosystems, but they have also been critical to the advancement of our capacity to model the complex interactions between environmental and plankton dynamics. The breadth of ecosystem components that are measured routinely at the WCO has enabled a form of digital hypothesis testing using

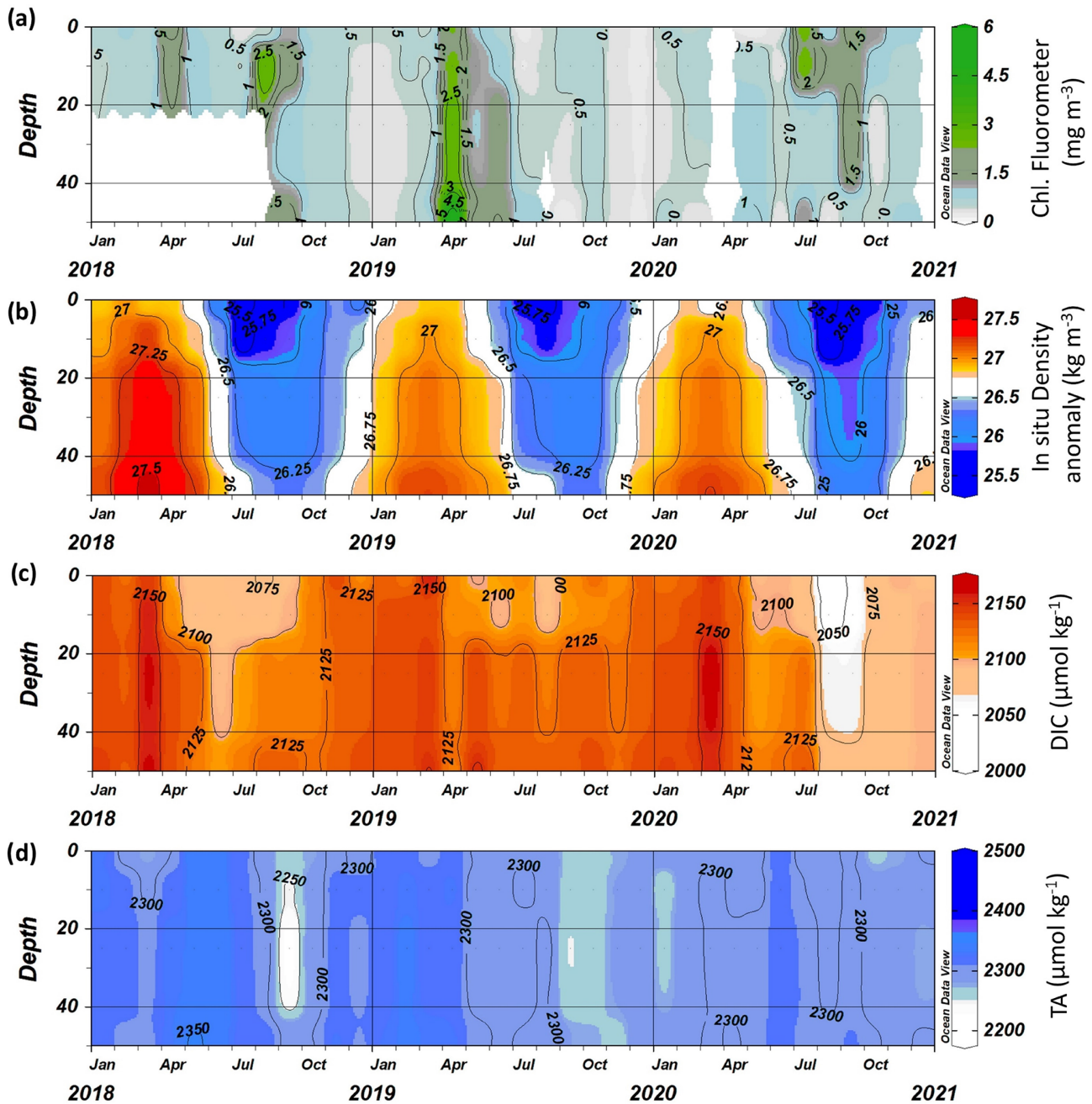


Figure 9. Depth profiles of (a) chlorophyll, (b) density, (c) dissolved inorganic carbon (DIC) and (d) total alkalinity (TA) at station L4 between 2018 and 2021.

biogeochemical and plankton models (Polimene et al., 2014) comparable to the more traditional approach to hypothesis testing through experimental work under controlled laboratory conditions. The WCO time series have contributed to a broad range of developments of the European Regional Seas Ecosystem Model (ERSEM) originating from testable hypotheses. These include the photophysiological control of plankton succession (Atkinson et al., 2018), the role of food quality on plankton blooms (Polimene et al., 2015), the bac-

terial carbon pump (Polimene et al., 2017) or the role of mixoplankton in plankton succession dynamics (Leles et al., 2021). Models can also represent a key source of information (a concept generally referred to as data augmentation) for the interpretation of time series. For example, operational models, such as the Western Channel Observatory Operational Forecast (WCOOF) model (Torres and Uncles, 2011), can be used to reconstruct back trajectories of plankton samples to explain community variations or assist in the evaluation of

carbon sequestration estimates (Queirós et al., 2023). Models like WCOOF can also be used to interpolate environmental conditions to explain observed plankton shifts (e.g. rapid changes to weak stratification not captured by the time series' sampling frequency) or to interpolate sparse measurements (Sims et al., 2022). Ultimately, models can also inform and optimize observational approaches, e.g. primary production estimation from oxygen/argon ratios and oxygen isotopes.

6.2 The WCO contribution to wider observing networks to report on ocean health

Marine time series such as those provided by the WCO form an important component to a series of wider networks for reporting on pelagic and benthic ecosystem status, and these networks span a range of scales. At the smallest scale of the south-western UK, the Western Channel Observatory observations form important contributions to the annual Southwest Marine Ecosystems Annual Reports (Smyth, 2022; Atkinson et al., 2022). At the UK scale, WCO data inform on regional-scale trends in plankton as part of the Marine Climate Change Impacts Partnership (Edwards et al., 2020; Findlay et al., 2022) and the plankton data contribute to Indicator C5 within the UK's 25 Year Environment Plan. At a slightly wider scale (north-western European shelf and north-eastern Atlantic), the WCO data form part of the policy reporting to meet statutory UK policy obligations under the UK Marine Strategy and OSPAR, for example in relation to carbonate chemistry (McGovern et al., 2023) or pelagic habitats (Ostle et al., 2021; Holland et al., 2023).

One advantage of the WCO plankton data is that they are both relatively complete in terms of taxonomic resolution and that they span multiple decades, which has enabled their use as a test bed dataset for developing indicators (McQuatters-Gollop et al., 2019), to examine how representative single sampling stations are of wider areas (Ostle et al., 2017) and to develop indicators that include the full suite of plankton, including major groups such as gelatinous species and picoplankton, which are not included as indicators from other longer-term monitoring programmes (such as the Continuous Plankton Recorder). At wider ocean basin scales, the Western Channel Observatory data contribute to a series of reporting networks, for example oceanography, through ICES Report on Ocean Climate (IROC; Gonzalez-Pola et al., 2022) publications, or plankton, through the International Group of Marine Ecological Time series (IGMETS; O'Brien et al., 2017). Because the WCO spans a small area, building these wider networks of time series is a vital tool to understand the spatial–temporal imprint of climate change amid other, more acute and localized stressors (Ratnarajah et al., 2023).

6.3 The future: melding new technological developments with existing long time series

Most of the longer time series of data that we provide here have been collected with traditional techniques that require direct collection of samples, their transport to the laboratory, and subsequent expert chemical or taxonomic analysis. These methods are expensive and time-consuming; for this reason, time series worldwide are under threat from funding cuts and loss of expert taxonomists (Vucetich et al., 2020). Concurrent with this, new techniques are being developed for time series, for example, underway autonomous vehicles or remote collection of data with acoustic methods or data from moorings are processed by automated particle imaging using machine learning particle classification. Likewise, there is much interest in the development of molecular (e.g. eDNA) approaches to observing plankton.

Some of these newer “big-data” approaches are now being developed at the WCO. The NERC-funded “Automated in situ Plankton Imaging and Classification System” (APICS) project represents the first in situ co-deployment of an Imaging FlowCytobot (McLANE Research Laboratories, Inc.) and a plankton imager (Pi-10; Plankton Analytics Ltd.) in the world. APICS will generate abundance and diversity data for organisms spanning 3–4 orders of magnitude in size, i.e. 5 µm–20 mm, on hourly timescales, which will allow a ca. 100-fold increase in phyto- and zooplankton sampling frequency at station L4 in 2024. APICS will allow critical plankton data to be collected at the same temporal resolution as physical and chemical variables. The establishment of an in situ imaging time series of plankton and particles at station L4 will also facilitate the collection of highly temporally resolved pelagic suspended particulate matter/particulate organic carbon (POC) concentration data, using image-based POC estimation methods that are currently being refined (Giering et al., 2020; Saskia Rühl, unpublished data). Likewise, molecular metabarcoding approaches are being developed and tested alongside traditional techniques (Lindque et al., 2013; Parry et al., 2021), with eDNA time series being run alongside conventional sampling (Karen Tait, unpublished data).

The WCO's rich background of contextual data, relative ease of accessibility and concurrent exposure to large wave amplitudes from the south-western weather systems also make it an ideal testing ground for new technology, and this development is currently highly active, with Smart Sound Plymouth (<https://www.smartsoundplymouth.co.uk>, last access: 29 November 2023) providing ambitious new directions in this area.

A common conception of funders and policy-makers is that new moored and autonomous instrumentation will provide a substitute for traditional monitoring that involves the collection of samples and analysis by skilled humans in a land-based laboratory. This may seem like an attractive way of reducing a whole suite of costs, including those for staff

time, training of taxonomic skills, and ship time and fuel, as well as the carbon footprint. These new methods, however, produce fundamentally different types of data to traditional approaches. This presents difficulties when melding the data together. This is a key detail, because the detection of climate change responses usually requires multiple decades of data collected in consistent fashion to have sufficient statistical power to detect change. Instead, the new approaches provide novel insights, often at much higher temporal and spatial resolution than traditional methods, better suited to capturing delicate organisms (Cross et al., 2015), vertical structure (Cornwell et al., 2020) or revealing the “hidden” diversity of assemblages through molecular metabarcoding (Lindeque et al., 2013; Parry et al., 2021). This is complementary, rather than alternative, information with respect to ongoing monitoring and provides fresh views on how these ecosystems function. These new technologies provide far more data than can be processed manually, and traditional methods are currently essential for ground-truthing the new data. We hope that sustained observations such as the WCO will embrace the strengths of both traditional and new approaches in the following decades.

7 Data availability

The full data and metadata are stored in a reputable UK repository known as DASSH (The Data Archive for Marine Species and Habitats) at the Marine Biological Association, Plymouth: <https://www.dassh.ac.uk> (last access: 29 November 2023). The data are available via the following link: <https://doi.org/10.17031/645110fb81749> (McEvoy and Atkinson, 2023). Upon using this particular version, we kindly request that both McEvoy and Atkinson (2023) and this paper in *Earth System Science Data* are cited. This paper gives a full description of the methods and, via citations, due credit to the authors who contributed the datasets. Citing the present paper when the data are used also allows standard literature searches to reveal data usage, and this provides valuable evidence to warrant continued funding of the WCO.

In future, we aim to produce more WCO data papers with updated, persistently identified (using DOIs) time series, corrections of any errors and extended data fields. Importantly, some of the older (pre-1988) datasets’ time series and metadata held by the MBA were not available for this data paper. We anticipate that later persistently identified versions of the data will be able to include more complete historical data as well as their metadata.

8 Potential uses and limitations of these data

The dataset that we provide here has both strengths and limitations. Its main strength is that it combines, for the first time, data that span from oceanography to sharks and from microbial diversity up to benthic macrofauna and fish. We hope

that this is particularly valuable for education purposes – for example, student projects in which the student can spend less time trying to hunt down scattered datasets and melding them together and more time analysing datasets. We have also, if possible, presented the data in units of mass as well as abundance, as this is particularly amenable to carbon budgets (Queirós et al., 2019), biogeochemical studies (Barnes et al., 2015) or models (Kenitz et al., 2017; Polimene et al., 2014). This study also represents the first attempt to put benthic and pelagic datasets together, and we hope that this helps to make the WCO a natural laboratory to study benthic–pelagic coupling. Another advantage of this summary version of the dataset is that it spans over 100 years, with over 3 decades of high-quality data from multiple trophic levels. Because the WCO has witnessed substantial warming and broadly responded in a manner similar to the wider north-western European shelf (Bedford et al., 2020; Schmidt et al., 2020), our summary dataset provides a test bed to study the mechanisms that control seasonality and climate change response across multiple trophic levels.

Our data summary also has a series of key limitations, the first one being its taxonomic resolution. To make our database manageable in size, we have condensed a large species lists (over 400 planktonic taxa alone) into a just a few dozen functional groups. Users wishing to estimate diversity changes or responses of individual key species will need to source the original datasets via the points of contact listed on the WCO data catalogue: https://westernchannelobservatory.org.uk/pelagic_TS.php (last access: 29 November 2023). Likewise, those studying short-term dynamics or “events”, for instance phenology shifts, bloom dynamics or extreme weather, may prefer to access the individual time points, which are typically weekly at L4. Despite this proviso and recommendations that < 20 d resolution are needed to reveal phenology shifts (Henson et al., 2018), long-term studies of such phenomena have tended to take the pragmatic approach by averaging irregularly spaced time points into monthly blocks to improve precision, data coverage and to fill data gaps (Atkinson et al., 2021; Barton et al., 2020; Edwards and Richardson, 2004; Fanjul et al., 2018; Uriarte et al., 2021).

A final three requests to users, in addition to correct citation, are as follows: (1) please let us know if you find any errors, (2) please suggest any improvements for the next version, and (3) please let us know if you want to incorporate this dataset into a wider data networking or databasing initiative. This is to ensure that data do not become separated from metadata and that old, outdated legacy versions of the data do not linger on data portals.

Appendix A

Table A1. L4 metadata for WCO monthly time series (1988–2022). The information includes sampling and analysis protocols as well as data coverage and links with respect to the availability of the full datasets. Column numbers reference the data sheet available from <https://doi.org/10.17031/645110fb81749> (McEvoy and Atkinson, 2023).

Data type and column headings	Sampling and analysis method	Data coverage and availability of full data
L4: water temperature		
L4_Temp_0m_degC L4_Temp_10m_degC L4_Temp_25m_degC L4_Temp_50m_degC Columns 3–6	<p>Measurements were taken weekly (conditions permitting).</p> <p>From March 1988 to April 1993, the surface temperature (0 m) was measured using a mercury thermometer in a stainless-steel bucket of freshly collected seawater.</p> <p>From May 1993 to December 2001, a PML conductivity–temperature–depth (CTD) instrument was also concurrently used in addition to the bucket method.</p> <p>In January 2002, this PML CTD was replaced with a SeaBird SBE19+ CTD.</p> <p>Bucket temperatures were adjusted to CTD equivalents using a regression equation for parallel determinations. For surface values, we obtained a value for each sampling week based on this adjusted bucket temperature if only this was available. If both bucket and CTD data were available, we used the CTD temperature. We then derived arithmetic mean temperatures for each month.</p>	<p>From March 1988 to December 2021, monthly data were derived from 1491 sampling points.</p> <p>From May 1993 onwards, a CTD was used that provided 1142, 1140 and 977 weekly time points for 10, 25 and 50 m, respectively.</p> <p>Data are available from https://www.westernchannelobservatory.org.uk/data.php (last access: 29 November 2023).</p>
L4: nutrients		
L4_Nitrite_0m_μM L4_Nitrite_10m_μM L4_Nitrite_25m_μM L4_Nitrite_50m_μM L4_Nitrite+Nitrate_0m_μM L4_Nitrite+Nitrate_10m_μM L4_Nitrite+Nitrate_25m_μM L4_Nitrite+Nitrate_50m_μM L4_Ammonia_0m_μM L4_Ammonia_10m_μM L4_Ammonia_25m_μM L4_Ammonia_50m_μM L4_Silicate_0m_μM L4_Silicate_10m_μM L4_Silicate_25m_μM L4_Silicate_50m_μM L4_Phosphate_0m_μM L4_Phosphate_10m_μM L4_Phosphate_25m_μM L4_Phosphate_50m_μM Columns 7–26	<p>Measurements were taken weekly (conditions permitting).</p> <p>Samples were returned (in the cool and dark) to the laboratory in Plymouth as soon as possible.</p> <p>Triplicate samples were analysed using 0.2 μm Millipore Fluoropore filtered and non-filtered water.</p> <p>The analyser used was a five-channel BRAN+LUEBBE segmented flow system.</p> <p>The methodology was standardized according to PML protocols.</p> <p>Since 2007, samples have been analysed as soon as possible after collection. Prior to this, samples were frozen and analysed in batches.</p> <p>Due to the storage method utilized, concentrations of ammonia should be treated with care. It is more appropriate to consider trends than accurate concentrations.</p> <p>Quality control procedures were carried out using KANSO certified reference material.</p> <p>Scientists participated in the Quality Assurance of Information for Marine Environmental Monitoring In Europe (QUASIMEME) programme. This summary dataset provides a mean value of all available determinations within any given calendar month.</p> <p>In the original dataset, the symbol “<” refers to concentrations below the detection limit; these have been assigned a value of zero before averaging.</p>	<p>Surface (0 m) data were collected from January 2000 to December 2021.</p> <p>Profiles for 10, 25 and 50 m were collected from January 2012 to December 2021.</p> <p>Full data lists include individual replicate measurements from the weekly resolution sampling.</p> <p>Publicly accessible nutrient data were accessed on 14 July 2022 at https://www.westernchannelobservatory.org.uk/data.php.</p>

Table A1. Continued.

Data type and column headings	Sampling and analysis method	Data coverage and availability of full data
L4: carbonate chemistry DIC (dissolved inorganic carbon) and TA (total alkalinity)		
L4_DIC_0m_μmol kg ⁻¹ L4_DIC_10m_μmol kg ⁻¹ L4_DIC_25m_μmol kg ⁻¹ L4_DIC_50m_μmol kg ⁻¹ L4_TA_0m_μmol kg ⁻¹ L4_TA_10m_μmol kg ⁻¹ L4_TA_25m_μmol kg ⁻¹ L4_TA_50m_μmol kg ⁻¹ Columns 27–34	<p>Measurements were taken weekly (conditions permitting).</p> <p>Borosilicate glass bottles with ground glass stoppers were used to collect seawater from the Niskin bottles. Sample bottles were rinsed, filled and poisoned with mercuric chloride according to standard procedures detailed in Dickson et al. (2007).</p> <p>Samples were returned to PML for analysis. DIC was measured using a dissolved inorganic carbon analyser (Model AS-C3, Apollo SciTech). The analyser adds a strong acid (10 % H₃PO₄ plus 10 % NaCl solution) that causes carbon species within the seawater to be converted to CO₂ gas; this gas is purged from the sample by pure nitrogen (N₂) carrier gas and is dried and cooled to reduce water vapour. The concentration of the dried CO₂ gas is measured with a LI-COR LI-7000 CO₂ analyser. The total amount of CO₂ is quantified as the integrated area under the concentration–time curve and converted to DIC using a standard curve created by analysing known concentrations of the certified reference materials (CRMs, Dickson CO₂). A measurement volume of 0.75 mL was used, with up to five measurements made from each sample. Values outside a 0.1 % range were excluded from the final result.</p> <p>Duplicate measurements provided an estimate of measurement error of < 0.1 %. DIC was corrected for the addition of mercuric chloride.</p> <p>TA was measured with the open-cell potentiometric titration method (Dickson et al., 2007) on 12 mL sample volumes using an automated titrator (alkalinity titrator model AS-ALK2, Apollo SciTech). Calibration was carried out using CRMs (Dickson CO₂). Duplicate measurements were made for each sample, and the estimate of measurement error was < 0.5 %. TA was corrected for the addition of mercuric chloride.</p>	<p>Surface (0 m) and 50 m measurements were made from October 2008 to December 2020, and 10 and 25 m measurements were made from September 2017 to December 2020.</p> <p>Data are available from the British Oceanographic Data Centre (BODC) at https://doi.org/10.5285/1ec0cae5-071d-16e1-e053-6c86abc07d47 (Cummings et al., 2015) and https://www.westernchannelobservatory.org.uk/C_chem.php (last access: 29 November 2023).</p>

Table A1. Continued.

Data type and column headings	Sampling and analysis method	Data coverage and availability of full data
L4: methane (CH ₄) and nitrous oxide (N ₂ O) concentrations		
L4_Ch4_0m nmol L ⁻¹ L4_Ch4_10m nmol L ⁻¹ L4_Ch4_25m nmol L ⁻¹ L4_Ch4_50m nmol L ⁻¹ L4_N2O_0m nmol L ⁻¹ L4_N2O_10m nmol L ⁻¹ L4_N2O_25m nmol L ⁻¹ L4_N2O_50m nmol L ⁻¹ Columns 35–42	<p>Borosilicate glass bottles with ground glass stoppers were used to collect seawater from the Niskin bottles for methane and nitrous oxide analysis; both gases were determined from the same bottle.</p> <p>Prior to all depths being collected in 2019, samples were collected in triplicate. Sample bottles were rinsed, filled and poisoned with mercuric chloride according to standard procedures detailed in Dickson et al. (2007). Samples were returned to PML for analysis.</p> <p>All samples were analysed within 3 months of collection.</p> <p>Samples were placed into a water bath at 25 °C, and the temperature was equilibrated for a minimum of 1 h before analysis.</p> <p>Samples were analysed by single-phase equilibration gas chromatography using a flame ionization detector for CH₄ and an electron capture detector for N₂O, similar to the procedure described by Upstill-Goddard et al. (1996). Samples were calibrated against three certified (±5 %) reference standards (Air Products and Chemicals, Inc.) that are traceable to NOAA WMO-N2O-X2006A. Concentrations in seawater at equilibration temperature (~ 25 °C) and salinity were calculated from the solubility tables of Weiss and Price (1980).</p>	<p>Surface N₂O coverage is from 2011, whereas CH₄ coverage is from 2013.</p> <p>All four depths were sampled from 2019.</p> <p>Data are available from https://www.westernchannelobservatory.org.uk/data.php (last access: 29 November 2023).</p>
L4: water 16S alpha diversity		
L4_water_prokaryote_diversity_S_0m_16s SEQ (sequencing) L4_water_prokaryote_diversity_Pielou_0m_16s SEQ L4_water_prokaryote_diversity_Shannon_0m_16s SEQ Columns 43–45	<p>Measurements were taken weekly (conditions permitting).</p> <p>On each sampling date, 5 L of seawater was collected from the surface and filtered immediately (on board) through a 0.22 µm Sterivex cartridge (Millipore).</p> <p>This was then stored at –80 °C at PML before further processing.</p> <p>Nucleic acids were extracted using the Qiagen AllPrep DNA/RNA Mini Kit. The Sterivex barrel was first filled with Buffer RLT lysis buffer and heated to 65 °C for 30 min. DNA and RNA were then extracted from the lysate following the manufacturer's instructions. DNA samples were used for microbiome analyses by sequencing of 16S rRNA genes using the 515F (GTGYCAGCMGCCGCGGTAA) and 806R (GGACTACNVGGGTWTCTAAT) Earth Microbiome V4 PCR primers. Sequencing was performed on the MiSeq personal sequencer (Illumina, San Diego, CA, USA) using the V2 500 reagent kit by commercial contract (NU-OMICS, UK). Demultiplexed paired-end FASTQ files were analysed using QIIME2 and amplicon sequence variants (ASVs) generated using DADA2. For each sample, the number of ASVs, Pielou evenness and Shannon diversity were calculated.</p>	<p>Data span from February 2012 to November 2019.</p> <p>Data are available from Karen Tait (PML): https://www.westernchannelobservatory.org.uk/data.php (last access: 29 November 2023).</p>

Table A1. Continued.

Data type and column headings	Sampling and analysis method	Data coverage and availability of full data
L4: fluorometer-derived chlorophyll- <i>a</i> concentrations		
L4_Ch1_0m_Fluorom_mgChl m-3 L4_Ch1_10m_Fluorom_mgChl m-3 L4_Ch1_25m_Fluorom_mgChl m-3 L4_Ch1_50m_Fluorom_mgChl m-3 Columns 46–49	Measurements were taken weekly (conditions permitting). Triplicate 100 mL water samples were filtered onto 25 mm GFF filters. Samples were extracted overnight at 4 °C and analysed on a Turner fluorometer, according to the procedure outlined in Welschmeyer (1994).	Surface (0 m) and 10 m measurements span from February 1992 to 2020 and comprise 1110 and 568 weekly resolution samples, respectively. All depths were sampled from 2018. Publicly accessible nutrient data are available from https://www.westernchannelobservatory.org.uk/data.php (last access: 29 November 2023).
L4: pigment sums generated from primary pigment data, determined by high-performance liquid chromatography (HPLC)		
L4_[TChl a]_0m_HPLC_mg m-3 L4_[TChl]_0m_HPLC_mg m-3 L4_[PPC]_0m_HPLC_mg m-3 L4_[PSC]_0m_HPLC_mg m-3 L4_[PSP]_0m_HPLC_mg m-3 L4_[TAcc]_0m_HPLC_mg m-3 L4_[TPig]_0m_HPLC_mg m-3 L4_[TChl a]_10m_HPLC_mg m-3 L4_[TChl]_10m_HPLC_mg m-3 L4_[PPC]_10m_HPLC_mg m-3 L4_[PSC]_10m_HPLC_mg m-3 L4_[PSP]_10m_HPLC_mg m-3 L4_[TAcc]_10m_HPLC_mg m-3 L4_[TPig]_10m_HPLC_mg m-3 L4_[TChl a]_25m_HPLC_mg m-3 L4_[TChl]_25m_HPLC_mg m-3 L4_[PPC]_25m_HPLC_mg m-3 L4_[PSC]_25m_HPLC_mg m-3 L4_[PSP]_25m_HPLC_mg m-3 L4_[TAcc]_25m_HPLC_mg m-3 L4_[TPig]_25m_HPLC_mg m-3 L4_[TChl a]_50m_HPLC_mg m-3 L4_[TChl]_50m_HPLC_mg m-3 L4_[PPC]_50m_HPLC_mg m-3 L4_[PSC]_50m_HPLC_mg m-3 L4_[PSP]_50m_HPLC_mg m-3 L4_[TAcc]_50m_HPLC_mg m-3 L4_[TPig]_50m_HPLC_mg m-3 Columns 50–77	Measurements were taken weekly (conditions permitting). Parameter names (if shortened versions are used in column titles) are as follows: [TChl <i>a</i>] is total chlorophyll <i>a</i> = [Chlide <i>a</i>] + [DVChl <i>a</i>] + [Chl <i>a</i>]; [TChl] is total chlorophyll = [TChl <i>a</i>] + [TChl <i>b</i>] + [TChl <i>c</i>]; [PPC] is photoprotective carotenoids = [Allo] + [Diad] + [Diato] + [Zea] + [Caro]; [PSC] is photosynthetic carotenoids = [But] + [Fuco] + [Hex-fuco] + [Perid]; [PSP] is photosynthetic pigments = [PSC] + [TChl]; [TAcc] is total accessory pigments = [PPC] + [PSC] + [TChl <i>b</i>] + [TChl <i>c</i>]; and [TPig] is total pigments = [TAcc] + [TChl <i>a</i>]. Total chlorophyll <i>a</i> = chlorophyllide <i>a</i> + divinyl chlorophyll <i>a</i> + chlorophyll <i>a</i> ; this parameter may be underestimated if chlorophyllide <i>a</i> is not quantified. Total chlorophyll <i>b</i> = chlorophyll <i>b</i> + divinyl chlorophyll <i>b</i> . Divinyl chlorophyll <i>b</i> co-elutes with chlorophyll <i>b</i> under the HPLC conditions used to generate these data, so these values were not quantified separately. Divinyl chlorophyll <i>b</i> is not expected to be present in UK waters. Total chlorophyll <i>c</i> = chlorophyll <i>c</i> 1 + chlorophyll <i>c</i> 2 + chlorophyll <i>c</i> 3. Carotenes = $\beta\epsilon$ -carotene + $\beta\beta$ -carotene. Alloxanthin is quantified by both HPLC methods used to generate L4 pigment data. 19'-Butanoyloxyfucoxanthin is quantified by both HPLC methods used to generate L4 pigment data. Diadinoxanthin is quantified by both HPLC methods used to generate L4 pigment data. Diatoxanthin is quantified by both HPLC methods used to generate L4 pigment data. Fucoxanthin is quantified by both HPLC methods used to generate L4 pigment data. 19'-Hexanoyloxyfucoxanthin is quantified by both HPLC methods used to generate L4 pigment data. Data generated using the Barlow HPLC method (1999–2011) may include prasinoxanthin (when present).	

Table A1. Continued.

Data type and column headings	Sampling and analysis method	Data coverage and availability of full data
	<p>Peridinin is quantified by both HPLC methods used to generate L4 pigment data.</p> <p>Zeaxanthin is quantified by both HPLC methods used to generate L4 pigment data.</p> <p>Chlorophyll <i>a</i> includes allomers and epimers and is quantified by both HPLC methods used to generate L4 pigment data.</p> <p>Divinyl chlorophyll <i>a</i> is quantified by both HPLC methods used to generate L4 pigment data.</p> <p>Chlorophyllide <i>a</i> was quantified in 2002, 2004–2005 and May 2011 onwards.</p> <p>With respect to chlorophyll <i>b</i> and divinyl chlorophyll <i>b</i>, divinyl chlorophyll <i>b</i> co-elutes with chlorophyll <i>b</i> under the HPLC conditions used to generate these data; thus, these values were not quantified separately. Divinyl chlorophyll <i>b</i> is not expected to be present in UK waters.</p> <p>Chlorophyll <i>c</i>1 is quantified separately from chlorophyll <i>c</i>2 from May 2011 onwards.</p> <p>Chlorophyll <i>c</i>2 includes chlorophyll <i>c</i>1 for data from 1999 to April 2011.</p> <p>Chlorophyll <i>c</i>3 is quantified by both HPLC methods used to generate L4 pigment data.</p> <p>$\beta\epsilon$-Carotene (alpha-carotene) is quantified separately from $\beta\beta$-carotene from May 2011 onwards.</p> <p>$\beta\beta$-Carotene (beta-carotene) includes $\beta\epsilon$-carotene for data from 1999 to April 2011.</p> <p>The Barlow HPLC method reference was sourced from Barlow et al. (1997).</p> <p>The column used was a MOS-2 Hypersil column (100 × 4.6 mm, 3 μm particle size).</p> <p>A flow rate of 1 mL min⁻¹ was utilized.</p> <p>The mobile phase was employed, as per Barlow et al. (1997).</p> <p>The extraction solvent and volume were as follows: 90 % acetone and 2 mL.</p> <p>An internal standard was used; <i>trans</i>-β-Apo-8'-carotenal was employed until 2008.</p> <p>The disruption method and time were as follows: sonication (probe) and 35 s.</p> <p>The soak time was 1 h.</p> <p>The clarification procedure was centrifugation.</p> <p>The injection procedure and volume were as follows: autosampler mixes sample with ammonium acetate (1 M) in 50/50 ratio by volume and injects 50 μL.</p> <p>The calibration procedure was a single point technique.</p> <p>Standards were sourced from DHI, Denmark.</p> <p>The absorption coefficients used were those provided with the standards from DHI.</p> <p>The expected capability of the method was not recorded.</p> <p>Quality assurance protocols were as follows: up to 20 samples were analysed per day, so the maximum time of samples in the autosampler was 24 h. The autosampler was maintained at 4 °C.</p> <p>The Zapata HPLC method reference was sourced from Zapata et al. (2000).</p>	

Table A1. Continued.

Data type and column headings	Sampling and analysis method	Data coverage and availability of full data
	<p>The column used was a Waters Symmetry C8 column (150 × 2.1 mm, 3.5 μm particle size). A flow rate of 200 μL min⁻¹ was utilized.</p> <p>The mobile phase was employed, as described by Zapata et al. (2000).</p> <p>The extraction solvent and volume were as follows: 90 % acetone and 2 mL.</p> <p>No internal standard was used.</p> <p>The disruption method and time used were as follows: sonication (probe) and 35 s.</p> <p>The soak time was 1 h.</p> <p>The clarification procedure was centrifugation and filtration (0.45 μm Teflon syringe filter).</p> <p>The injection procedure and volume were as follows: autosampler mixes 200 μL sample and 80 μL water in a vial, and a total of 25 μL of this mixture is injected (actual injection volume of sample is 17.86 μL).</p> <p>The calibration procedure was a multipoint technique: three solutions bracketing the limit of quantification (LOQ) and three bracketing the expected sample concentration.</p> <p>Standards were sourced from DHI, Denmark.</p> <p>The absorption coefficients used were those provided with standards from DHI.</p> <p>The expected capability of the method was as follows: average precision and accuracy for chlorophyll <i>a</i> (standards) was 1.44 % and 2.01 %, respectively.</p> <p>Quality assurance protocols were as follows: first run of the day was discarded. A sample of mixed pigments was run prior to any samples to check retention times and the resolution of critical pairs.</p> <p>Three samples of chlorophyll standard were analysed with each sample set to check that the response factor is within 5 % of the calibration value.</p> <p>Up to 20 samples are analysed per day, so the maximum time of samples in the autosampler was 24 h. The autosampler was maintained at 4 °C.</p> <p>Pipette accuracy was determined daily by weighing.</p>	<p>Surface coverage is from March 1999 to December 2014, whereas coverage for 10, 25 and 50 m depths is from 2009 onwards (some gaps in data) until 2014.</p> <p>Source data are available from https://www.westernchannelobservatory.org.uk/data.php (last access: 29 November 2023).</p>

Table A1. Continued.

Data type and column headings	Sampling and analysis method	Data coverage and availability of full data
L4: < 20 µm plankton abundance profiles measured by flow cytometry		
L4_Syn_0m_FCM_cells mL-1 L4_Picoeuk_0m_FCM_cells mL-1 L4_Nanoeuk_0m_FCM_cells mL-1 L4_Cocco_0m_FCM_cells mL-1 L4_Crypto_0m_FCM_cells mL-1 L4_HNan_0m_FCM_cells mL-1 L4_HNAbacteria_0m_FCM_cells mL-1 L4_LNAbacteria_0m_FCM_cells mL-1 L4_Syn_10m_FCM_cells mL-1 L4_Picoeuk_10m_FCM_cells mL-1 L4_Nanoeuk_10m_FCM_cells mL-1 L4_Cocco_10m_FCM_cells mL-1 L4_Crypto_10m_FCM_cells mL-1 L4_HNan_10m_FCM_cells mL-1 L4_HNAbacteria_10m_FCM_cells mL-1 L4_LNAbacteria_10m_FCM_cells mL-1 L4_Syn_25m_FCM_cells mL-1 L4_Picoeuk_25m_FCM_cells mL-1 L4_Nanoeuk_25m_FCM_cells mL-1 L4_Cocco_25m_FCM_cells mL-1 L4_Crypto_25m_FCM_cells mL-1 L4_HNan_25m_FCM_cells mL-1 L4_HNAbacteria_25m_FCM_cells mL-1 L4_LNAbacteria_25m_FCM_cells mL-1 L4_Syn_50m_FCM_cells mL-1 L4_Picoeuk_50m_FCM_cells mL-1 L4_Nanoeuk_50m_FCM_cells mL-1 L4_Cocco_50m_FCM_cells mL-1 L4_Crypto_50m_FCM_cells mL-1 L4_HNan_50m_FCM_cells mL-1 L4_HNAbacteria_50m_FCM_cells mL-1 L4_LNAbacteria_50m_FCM_cells mL-1	Measurements were taken weekly (conditions permitting). Samples were analysed in triplicate (phytoplankton and bacteria) or duplicate (heterotrophic nanoflagellates). Vertical profiles of the mean abundance of groups of microbial plankton, presented as cells per millilitre, were measured using flow cytometry (BD Accuri C6 flow cytometer). The groups quantified were divided into phytoplankton and heterotrophs. Phytoplankton groups quantified were as follows: Syn – <i>Synechococcus</i> sp. (cyanobacteria); Picoeuk – picoeukaryotes (smaller than 3 µm); Crypto – cryptophytes; Cocco – coccolithophores; and Nanoeuk – nanoeukaryotes (not already mentioned; 2–20 µm). Heterotrophs quantified were as follows: HNan – heterotrophic nanoflagellates; HNAbacteria – heterotrophic bacteria with relatively high nucleic acid content; and LNAbacteria – heterotrophic bacteria with relatively low nucleic acid content.	Data span from April 2007 to December 2021. Source data are available from https://www.westernchannelobservatory.org.uk/data.php (last access: 29 November 2023).
Columns 78–109		

Table A1. Continued.

Data type and column headings	Sampling and analysis method	Data coverage and availability of full data
L4: microscopy analysis of phytoplankton preserved in Lugol's solution and formalin		
L4_Diatoms_10m_microscopy_cells mL-1	Measurements were taken weekly (conditions permitting).	Data for a single depth (10 m) are available from October 1992 to December 2020, except for gaps in sampling between October 1994 and May 1995 and in December 2011.
L4_Dinoflagellates_10m_microscopy_cells mL-1	Paired 200 mL water samples collected from 10 m depth using a Niskin bottle attached to the CTD were immediately fixed in (1) acid Lugol's iodine (for all taxa except coccolithophores) and (2) neutral formaldehyde for coccolithophores.	Data are available from the British Oceanographic Data Centre (BODC):
L4_Coccolithophores_10m_microscopy_cells mL-1		http://doi.org/10.5285/c9386b5c-b459-782f-e053-6c86abc0d129
L4_Flagellates_10m_microscopy_cells mL-1		(Widdicombe and Harbour, 2021) and https://www.westernchannelobservatory.org.uk/data.php (last access: 29 November 2023).
L4_Phaeocystis_10m_microscopy_cells mL-1	Subsamples were analysed by light microscopy using the settlement technique (Utermohl, 1958), identified to the species level where possible, and organized into six functional groups.	
L4_Ciliates_10m_microscopy_cells mL-1		
L4_Diatoms_10m_microscopy_mgC m-3	The mean cell dimensions of each taxa were used to calculate species-specific biovolumes which were converted to carbon biomass using the equations of Menden-Deuer and Lessard (2000).	
L4_Dinoflagellates_10m_microscopy_mgC m-3		
L4_Coccolithophorid_10m_microscopy_mgC m-3	Abundance data are presented as cells per millilitre and biomass as milligrams of carbon per cubic metre.	
L4_Flagellates_10m_microscopy_mgC m-3		
L4_Phaeocystis_10m_microscopy_mgC m-3	Note that sample collection was via a deck hose in 2005. This caused damage to the fragile ciliates; hence, the count is much lower for that year.	
L4_Ciliates_10m_microscopy_mgC m-3		
Columns 110-121		

Table A1. Continued.

Data type and column headings	Sampling and analysis method	Data coverage and availability of full data
L4: FlowCam analysis of 63 μm mesh plankton net hauls (50–0 m)		
L4_Total_Diatoms_FlowCam_mgC m-3	Measurements were taken weekly (conditions permitting).	Data spanning from September 2012 to December 2013 are from 43 time points, whereas
L4_Total_Dinoflagellates_FlowCam_mgC m-3	Water samples were collected from a 0–50 m vertical haul using a 63 μm mesh WP2-style net (UNESCO, 1968, p. 153–157). There was a mesh change in July 2019: from 63 to 50 μm .	those from June 2015 to December 2019 are from 163 time points.
L4_Ciliates_FlowCam_mgC m-3	Prior to analysis, samples were prescreened using a 300 μm mesh. However, net samples collected between June 2015 and May 2016 were prescreened using a 200 μm mesh.	Abundance data are also available for meroplankton taxa, but these have not been converted to biomass to date.
L4_Colony flagellates_FlowCam_mgC m-3	Samples were analysed live, if possible, using a FlowCam VS IV model fitted with a 300 μm flow cell.	Source data are available from https://www.westernchannelobservatory.org.uk/data.php (last access: 29 November 2023).
L4_Large_Protists_FlowCam_mgC m-3	Analysis was carried out using $\times 4$ magnification and auto-image mode.	
L4_Total_Copepod nauplii_FlowCam_mgC m-3	Classification of the acquired images was carried out using VisualSpreadsheet (2012–2016) and EcoTaxa (2017–2019). Taxa were then assigned to six broad functional groups.	
Columns 122–127	The mean cell dimensions of each taxa were used to calculate species-specific biovolumes which were converted to carbon biomass using suitable carbon conversion equations. Biomass is presented as milligrams of carbon per cubic metre.	
	For diatoms, dinoflagellates (excluding <i>Noctiluca</i> and <i>Neoceratium</i> spp.) and ciliates, morphological information and shape assignment was used to calculate biovolume (Álvarez et al., 2012, their Table 1).	
	For <i>Noctiluca</i> and for <i>Neoceratium</i> spp., mean cell volumes were taken from Widdicombe et al. (2010). For all other dinoflagellates, diatoms and ciliates, cell biovolumes were converted to carbon biomass using the equations of Menden-Deuer and Lessard (2000).	
	For large protists mostly Radiolaria, the carbon conversion in Michaels et al. (1995) was used. Colonial flagellates were converted to carbon according to Børsheim and Bratbak (1987). Biomass of copepod nauplii was calculated using the equations of Uye et al. (1996).	

Table A1. Continued.

Data type and column headings	Sampling and analysis method	Data coverage and availability of full data
L4: <i>Noctiluca scintillans</i> microscopy analysis of WP2 net hauls (50–0 m)		
L4_Noctiluca_scintillans_WP2net_no.m-3 L4_Noctiluca_scintillans_WP2net_mcgC.m-3 Columns 128–129	<p>Measurements were taken weekly (conditions permitting).</p> <p>Two vertical hauls (50–0 m) were taken using 200 µm WP2 nets (UNESCO, 1968, p. 153–157).</p> <p>Both replicates' samples were analysed by subsampling, enumerated and identified, currently using an Olympus SZX16 stereo microscope fitted with a SZX2-ILLT LED transmitted-light illuminator stand.</p> <p>Source data represent weekly average abundance across the two replicates and were converted to number per cubic metre.</p> <p>Monthly abundance represents an arithmetic mean value from between one and five visits in any given month and on a weekly basis.</p> <p>Biomass calculations were derived from abundance data using a conversion factor of 0.020375 µg C per cell using the equations of Menden-Deuer and Lessard (2000).</p> <p>Be aware that zeros are present from 2009 onwards where there is confidence in the data. A zero represents “looked for but not present in the sample analysed”. Data prior to this are less certain, so zeros have been omitted.</p>	<p>Data are available from July 1997 to 2021 and are given in McEvoy et al. (2022a). Zooplankton abundance time series from net hauls at site L4 off Plymouth, UK, between 1988 and 2021.</p> <p>Data are available from the British Oceanographic Data Centre (BODC): https://doi.org/10.5285/e785f2f7-05d5-2f47-e053-6c86abc08bee (McEvoy et al., 2022a) and https://www.westernchannelobservatory.org.uk/data.php (last access: 29 November 2023).</p>

Table A1. Continued.

Data type and column headings	Sampling and analysis method	Data coverage and availability of full data
L4: zooplankton microscopy analysis of WP2 net hauls (50–0 m)		
L4_meroplankton_WP2net_no.m-3 L4_small_copepods_WP2net_no.m-3 L4_large_copepods_WP2net_no.m-3 L4_fish_larvae_WP2net_no.m-3 L4_gelatinous_predators_WP2net_no.m-3 L4_semi-gelatinous_predators_WP2net_no.m-3 L4_other_crustacean_holoplankton_WP2net_no.m-3 L4_other_non_crustacean_holoplankton_WP2net_no.m-3 L4_meroplankton_WP2net_mgCm-3 L4_small_copepods_WP2net_mgCm-3 L4_large_copepods_WP2net_mgCm-3 L4_fish_larvae_WP2net_mgCm-3 L4_gelatinous_predators_WP2net_mgCm-3 L4_semi-gelatinous_predators_WP2net_mgCm-3 L4_other_crustacean_holoplankton_WP2net_mgCm-3 L4_other_non_crustacean_holoplankton_WP2net_mgCm-3 Columns 130–145	<p>Measurements were taken weekly (conditions permitting). Two vertical hauls (50–0 m) were taken using 200 µm WP2 nets (UNESCO, 1968).</p> <p>Both replicates' samples were analysed by subsampling, enumerated and identified, currently using an Olympus SZX16 stereo microscope fitted with a SZX2-ILLT LED transmitted-light illuminator stand. More details are provided in Atkinson et al. (2015).</p> <p>Source data comprise the average abundance of the taxa that have been consistently identified since 1988. These source data are weekly averages across the two replicates, converted to number per cubic metre, and estimated biomass.</p> <p>Data presented here have been aggregated into functional groups broadly based on the life-forms for policy reporting in Ostle et al. (2021). There has, however, been a few further subdivisions to better reflect trophic mode. These functional group allocations are coded, and number-to-biomass conversion factors are provided within the “trait” header bar data from the source dataset.</p> <p>Therefore there are eight functional groups based partly on size, taxonomy and trophic mode, with separate columns for abundance and estimated biomass. Because biomass is a derived property, often with different conversion factors between the four seasons (see data source DOI), it is best to use numerical abundance data for population dynamics studies and biomass data for models, carbon budgets etc.</p> <p>As previously stated, the groups comprise the whole of the consistently identified zooplankton; therefore, adding them will give a good estimate of total metazoan zooplankton with the exception of ctenophores (see below).</p> <p>Meroplankton comprise all 14 taxa with code no. 38. They are numerically dominated by Cirripedia larvae. The biomass is strongly dominated by the larvae of Cirripedia, Decapoda and Polychaeta as well as Gammaridea amphipods. Fish and cnidarians are excluded, some of whom are meroplanktonic, but are all pooled within the “Fish larvae” and “Gelatinous predators” life-forms instead.</p> <p>Small copepods, excluding nauplii, are grouped under code no. 36 with the addition of the uncoded harpacticoid copepods. They are species with a total adult body length under 2 mm. They comprise 20 taxa dominated numerically by the <i>Oithona</i>, <i>Oncaea</i>, <i>Paracalanus</i> and <i>Pseudocalanus</i> genera and in biomass by <i>Pseudocalanus</i>, <i>Temora</i> and <i>Paracalanus</i>.</p> <p>Large copepods comprise eight taxa with code no. 36 and are species with a total body length of 2 mm or greater. Their numbers and biomass are strongly dominated by <i>Calanus helgolandicus</i>.</p> <p>Fish larvae, following the life-forms group for plankton reporting in Ostle et al. (2021), include eggs and larvae pooled, but the fish eggs have been omitted here to better describe the abundance of actively carnivorous groups.</p> <p>Gelatinous predators comprise cnidarians only, and they are dominated in terms of numbers and biomass by siphonophores. A notable taxon not included is ctenophores, due to potential inconsistency in counting in early years and due to preservation issues, which we are in the process of resolving.</p> <p>Semi-gelatinous predators comprise chaetognaths and <i>Tomopteris</i> spp., with numbers and biomass strongly dominated by the former.</p> <p>Other crustacean holoplankton are the remaining groups of crustacean holoplankton not covered above, namely, <i>Evadne</i> spp., <i>Podon</i> spp., Hyperiididae amphipods, mysids, and the various nauplii to adult stages of euphausiids. They are strongly dominated numerically and in terms of biomass by the Cladocera (miscoded as non-crustaceans in the source file).</p> <p>Other non-crustacean holoplankton are the remaining groups of crustacean holoplankton not covered above, namely, appendicularians, <i>Limacina</i> spp., doliolids and <i>Clione</i> spp. They are strongly dominated numerically and in terms of biomass by appendicularians.</p>	<p>Data span from March 1988 to December 2020 and were derived from 1452 sampling time points with a weekly resolution.</p> <p>Monthly mean data are available for each intervening month except August 2000 and typically represent an arithmetic mean value across between one and five weekly visits in any given month.</p> <p>Data are given in McEvoy et al. (2022b).</p> <p>Data are available from the British Oceanographic Data Centre (BODC): https://doi.org/10.5285/d7fb6ce3-7bc9-307b-e053-6c86abc0671b and https://www.westernchannelobservatory.org.uk/l4_zooplankton.php (last access: 29 November 2023).</p>

Table A1. Continued.

Data type and column headings	Sampling and analysis method	Data coverage and availability of full data
L4: <i>Calanus helgolandicus</i> weekly egg production using females from the western English Channel site L4		
L4_Calanus_eggs_watercolumn_expt_eggs per female per day Column 146	Measurements were taken weekly (conditions permitting). A live sample was collected and returned in the cool and dark to the laboratory in Plymouth as soon as possible. The sample was gently poured through a 200 µm mesh sieve. <i>Calanus</i> sp. females in healthy condition were picked out gently using stork-billed forceps under a microscope as quickly as possible. Five replicates, each containing five female <i>Calanus</i> sp., were incubated in the dark in filtered seawater for 24 h. Each beaker contained an egg collector. Temperature followed ambient conditions at the L4 surface. The eggs produced were collected and counted. Females were identified for species. Eggs were retained for hatching success.	The data span from February 1992 to November 2021 (where the availability of <i>Calanus</i> females allows). Data represent the mean number of eggs per female per day. Data are given in McEvoy et al. (2022c) and are available from https://doi.org/10.5285/e28496a4-0c72-0e7a-e053-6c86abc0d7c7 and https://www.westernchannelobservatory.org.uk/calanus_egg_production.php (last access: 29 November 2023).
L4: sediment 16S alpha diversity		
L4_sediment_prokaryote_diversity_S_0m_16S SEQ L4_sediment_prokaryote_diversity_Pielou_0m_16s SEQ L4_sediment_prokaryote_diversity_Shannon_0m_16s SEQ Columns 147–149	Sediments were collected using a box corer, and the uppermost 0–1 cm was carefully sampled by scraping into a sterile 2 mL tube. Eight replicate samples were taken for each sampling time, and four of those were employed for DNA extraction using 0.5 g sediment and Qiagen's DNeasy PowerSoil Kit according to the manufacturer's instructions. The 16S rRNA genes were sequenced using the Earth Microbiome V4 515F (GTGYCAGCMGCCGCG-GTAA) and 806R (GGACTACNVGGGTWCTAAT) PCR primers. Sequencing was performed on the MiSeq personal sequencer (Illumina, San Diego, CA, USA) using the V2 500 reagent kit by commercial contract (NU OMICS, UK). Demultiplexed paired-end FASTQ files were analysed using QIIME2 and amplicon sequence variants (ASVs) generated using DADA2. For each sampling occasion, the mean number of ASVs for the four replicates was calculated (S), along with the Pielou evenness and Shannon diversity.	Data span from February 2012 to 2019. In 2012 and from 2014 to 2019, the aim was to sample monthly when possible. In 2013, samples were collected in February and September only. Data are available from Karen Tait (PML): https://www.westernchannelobservatory.org.uk/data.php (last access: 29 November 2023).
L4: benthic fauna from box cores		
L4_Macrofaunal_Deposit_Feeders_50m_0.1m3_Box_Core_Average abundance of individual taxa per month L4_Macrofaunal_Suspension_Feeders_50m_0.1m3_Box_Core_Average abundance of individual taxa per month L4_Macrofaunal_Predators_50m_0.1m3_Box_Core_Average abundance of individual taxa per month L4_Macrofaunal_Scavengers_50m_0.1m3_Box_Core_Average abundance of individual taxa per month L4_Macrofaunal_Deposit_Feeders_50m_0.1m3_Box_Core_Average biomass of individual taxa per month L4_Macrofaunal_Suspension_Feeders_50m_0.1m3_Box_Core_Average biomass of individual taxa per month L4_Macrofaunal_Predators_50m_0.1m3_Box_Core_Average biomass of individual taxa per month L4_Macrofaunal_Scavengers_50m_0.1m3_Box_Core_Average biomass of individual taxa per month Columns 150–157	Measurements were taken monthly (conditions permitting). Four or five replicate 0.1 m ³ box cores of sediment were collected from 50 m depth. All sediment collected was sieved over a 0.5 mm mesh, and retained fauna was preserved in 10 % formaldehyde solution. Source taxa were identified and counted using stereo and compound microscopy to the species level or the lowest possible taxonomic resolution. The abundance and blotted wet weight (0.00000 g) per taxa were recorded per 0.1 m ³ box core sample. From four principle feeding traits, based on information primarily gathered from the BIOTIC database, one unique principle trait was assigned per taxa: calculated using algorithms based upon body composition, maximum length and body mass (https://www.marlin.ac.uk/biotic/ (last access: 6 December 2023)).	July 2008 to July 2019. Abundance and biomass data from 65 time points are presented as monthly averages per corresponding feeding trait: suspension feeders, deposit feeders, scavengers and carnivores. Mesher and McNeill (2022). Data are available from the British Oceanographic Data Centre (BODC): https://doi.org/10.5285/d9f44202-b0d4-646c-e053-6c86abc018c6 and https://www.westernchannelobservatory.org.uk/data.php (last access: 29 November 2023).

Table A1. Continued.

Data type and column headings	Sampling and analysis method	Data coverage and availability of full data
L4: Cephalopoda and demersal fish families by trawling		
L4_Cephalopoda_abundance_50-60m_Standard_Haul_individuals.trawl-1 L4_Bothidae_abundance_50-60m_Standard_Haul_individuals.trawl-1 L4_Soleidae_abundance_50-60m_Standard_Haul_individuals.trawl-1 L4_Callionymidae_abundance_50-60m_Standard_Haul_individuals.trawl-1 L4_Caproidae_abundance_50-60m_Standard_Haul_individuals.trawl-1 L4_Cepolidae_abundance_50-60m_Standard_Haul_individuals.trawl-1 L4_Triglidae_abundance_50-60m_Standard_Haul_individuals.trawl-1 L4_Clupeidae_abundance_50-60m_Standard_Haul_individuals.trawl-1 L4_Engraulidae_abundance_50-60m_Standard_Haul_individuals.trawl-1 L4_Pleuronectidae_abundance_50-60m_Standard_Haul_individuals.trawl-1 L4_Gadidae_abundance_50-60m_Standard_Haul_individuals.trawl-1 L4_Merlucciidae_abundance_50-60m_Standard_Haul_individuals.trawl-1 L4_Mullidae_abundance_50-60m_Standard_Haul_individuals.trawl-1 L4_Carangidae_abundance_50-60m_Standard_Haul_individuals.trawl-1 L4_Zeidae_abundance_50-60m_Standard_Haul_individuals.trawl-1 L4_Gobiidae_abundance_50-60m_Standard_Haul_individuals.trawl-1 L4_Scombridae_abundance_50-60m_Standard_Haul_individuals.trawl-1 L4_Scyllorhinidae_abundance_50-60m_Standard_Haul_individuals.trawl-1 L4_Scopthalmidae_abundance_50-60m_Standard_Haul_individuals.trawl-1 L4_Cephalopoda_biomass_50-60m_Standard_Haul_g.trawl-1 L4_Bothidae_biomass_50-60m_Standard_Haul_g.trawl-1 L4_Soleidae_biomass_50-60m_Standard_Haul_g.trawl-1 L4_Callionymidae_biomass_50-60m_Standard_Haul_g.trawl-1 L4_Cepolidae_biomass_50-60m_Standard_Haul_g.trawl-1 L4_Triglidae_biomass_50-60m_Standard_Haul_g.trawl-1 L4_Clupeidae_biomass_50-60m_Standard_Haul_g.trawl-1 L4_Engraulidae_biomass_50-60m_Standard_Haul_g.trawl-1 L4_Pleuronectidae_biomass_50-60m_Standard_Haul_g.trawl-1 L4_Gadidae_biomass_50-60m_Standard_Haul_g.trawl-1 L4_Merlucciidae_biomass_50-60m_Standard_Haul_g.trawl-1 L4_Mullidae_biomass_50-60m_Standard_Haul_g.trawl-1 L4_Carangidae_biomass_50-60m_Standard_Haul_g.trawl-1 L4_Zeidae_biomass_50-60m_Standard_Haul_g.trawl-1 L4_Scombridae_biomass_50-60m_Standard_Haul_g.trawl-1 L4_Scyllorhinidae_biomass_50-60m_Standard_Haul_g.trawl-1 L4_Lophiidae_biomass_50-60m_Standard_Haul_g.trawl-1 L4_Triakidae_biomass_50-60m_Standard_Haul_g.trawl-1 L4_Scopthalmidae_biomass_50-60m_Standard_Haul_g.trawl-1 L4_Rajidae_biomass_50-60m_Standard_Haul_g.trawl-1 L4_Lotidae_biomass_50-60m_Standard_Haul_g.trawl-1 L4_Moronidae_biomass_50-60m_Standard_Haul_g.trawl-1 L4_Congridae_biomass_50-60m_Standard_Haul_g.trawl-1 L4_Squalidae_biomass_50-60m_Standard_Haul_g.trawl-1 Columns 158–200	Standard hauls were collected using a large otter trawl (2008–June 2014), a Channel Hunter box trawl (July 2014–March 2015) deployed from Plymouth Quest, and a modified Channel Hunter box trawl (April 2015–September 2018) deployed from MBA Sepia. The trawl duration was approximately 40 min. Only trawls from 50 to 60 m were included. Individuals were identified to the species level, measured (mm) and weighed (g) on board. If a species was abundant, a subsample was weighed and total biomass was extrapolated. The abundances and biomass are reported at the family level, and only families comprising at least a 1 % contribution in at least 1 month are included.	Data span from April 2008 to September 2018. Between one and seven trawls were collected per month sampled (total of 282 and average of 2.88). Source data for 2015–2018 are available from the Data Archive for Seabed Species (DASSH): https://doi.org/10.17031/1802 (Brittain, 2021).

Table A2. E1 metadata for WCO monthly time series (1903–2021). The information includes sampling and analysis protocols as well as data coverage and links with respect to the availability of full datasets. Column numbers reference the data sheet available from <https://doi.org/10.17031/645110fb81749> (McEvoy and Atkinson, 2023).

Data type and column headings	Sampling and analysis method	Data coverage and availability of full data
E1: water temperature		
E1_Temp_0m_DegC E1_Temp_10m_DegC E1_Temp_20m_DegC E1_Temp_30m_DegC E1_Temp_40m_DegC E1_Temp_50m_DegC E1_Temp_60m_DegC E1_Temp_70m_DegC Columns 3–10	For the early period of the E1 time series, reversing thermometers were used. Values are derived from Niskin bottles and CTD measurements except for the period from December 1985 to April 2002, during which time no in situ sampling was undertaken and satellite sea surface temperature data pertaining to the middle of each month were used instead. When multiple sampling time points existed for a calendar month, we used the arithmetic mean value. Post-2002, a SeaBird SBE19+ was used.	Data begin in 1903, and no data are available from 1910 to 1920 nor from 1939 to 1945. Surface data are most extensive. For each depth, the number of sampling time points was 1146, 954, 892, 609, 740, 908, 262 and 815, respectively. The source dataset was produced for the ICES Report on Ocean Climate by Tim Smyth: https://ocean.ices.dk/core/iroc (last access: 29 November 2023) and https://www.westernchannelobservatory.org.uk/data.php (last access: 29 November 2023).
E1: nutrients		
E1_Nitrite_0m_μm E1_Nitrite_10m_μm E1_Nitrite_20m_μm E1_Nitrite_30m_μm E1_Nitrite_40m_μm E1_Nitrite_60m_μm E1_Nitrite+Nitrate_0m_μm E1_Nitrite+Nitrate_10m_μm E1_Nitrite+Nitrate_20m_μm E1_Nitrite+Nitrate_30m_μm E1_Nitrite+Nitrate_40m_μm E1_Nitrite+Nitrate_60m_μm E1_Ammonia_0m_μm E1_Ammonia_10m_μm E1_Ammonia_20m_μm E1_Ammonia_30m_μm E1_Ammonia_40m_μm E1_Ammonia_60m_μm E1_Silicate_0m_μm E1_Silicate_10m_μm E1_Silicate_20m_μm E1_Silicate_30m_μm E1_Silicate_40m_μm E1_Silicate_60m_μm E1_Phosphate_0m_μm E1_Phosphate_10m_μm E1_Phosphate_20m_μm E1_Phosphate_30m_μm E1_Phosphate_40m_μm E1_Phosphate_60m_μm Columns 11–40	Measurements were taken fortnightly (conditions permitting). Data from 2002 were collected and processed as follows: samples were returned in the cool and dark to the laboratory in Plymouth; samples were stored for 2–3 h before the return for analysis and were sometimes frozen; triplicate samples were analysed using 0.2 μm Millipore Fluoropore filtered and non-filtered water; the analyser was a five-channel Bran + Luebbe segmented flow system; the methodology was standardized according to PML protocols; due to the storage method, concentrations of ammonia should be treated with care (more appropriate to consider trends rather than accurate concentrations); quality control procedures were carried out using KANSO certified reference material; and scientists participated in the QUASIMEME programme. Data from last century were collected as follows: data were obtained from the link on the data page of the Western Channel Observatory website and extracted from the NOWESP (North West European Shelf Program) database for the period from 1934 to 1987. Source data include profile data from 0 to 80 m. These monthly data use depths of 0, 10 and 20 m because these are compatible with post-2002 records. The Nitrite + Nitrate column header describes post-2002 records. It is unclear if the last century values refer strictly to nitrate only or Nitrite + Nitrate. This summary dataset provides a mean value of all available determinations within any given calendar month. In the original dataset, the symbol “<” refers to concentrations below the detection limit. These have been assigned a value of zero before averaging.	Data from January 1934 comprise a few records for phosphate. In April 1948, records begin again for phosphate. In January 1951, records begin for silicate. In January 1966, records begin again for Nitrite + Nitrate. From January 1986 to December 2001, no data are available. From January 2002 to October 2021, all parameters are covered. The full data list comprises individual replicate measurements from the weekly resolution sampling. Publicly accessible nutrient data are available from https://www.westernchannelobservatory.org.uk/data.php (last access: 14 July 2022).

Table A2. Continued.

Data type and column headings	Sampling and analysis method	Data coverage and availability of full data
E1: carbonate chemistry DIC (dissolved inorganic carbon) and TA (total alkalinity)		
E1_DIC_0m_micromol kg-1 E1_DIC_60m_micromol kg-1 E1_TA_0m_micromol kg-1 E1_TA_60m_micromol kg-1 Columns 41–44	<p>Measurements were taken fortnightly (conditions permitting).</p> <p>Borosilicate glass bottles with ground glass stoppers were used to collect seawater from the Niskin bottles. Sample bottles were rinsed, filled and poisoned with mercuric chloride according to standard procedures detailed in Dickson et al. (2007).</p> <p>Samples were returned to PML for analysis.</p> <p>DIC was measured using a dissolved inorganic carbon analyser (Model AS-C3, Apollo SciTech). The analyser adds a strong acid (10 % H₃PO₄ plus 10 % NaCl solution), causing carbon species within the seawater to be converted to CO₂ gas that is then purged from the sample by pure nitrogen (N₂) carrier gas and is dried and cooled to reduce water vapour. The concentration of the dried CO₂ gas is measured with a LI-COR LI-7000 CO₂ analyser. The total amount of CO₂ is quantified as the integrated area under the concentration–time curve and converted to DIC using a standard curve created by analysing known concentrations of CRMs (Dickson CO₂). A measurement volume of 0.75 mL was used, with up to five measurements made from each sample. Values outside a 0.1 % range were excluded from the final result.</p> <p>Duplicate measurements provided an estimate of measurement error < 0.1 %. DIC was corrected for the addition of mercuric chloride.</p> <p>TA was measured with the open-cell potentiometric titration method (Dickson et al., 2007) on 12 mL sample volumes using an automated titrator (alkalinity titrator model AS-ALK2, Apollo SciTech). Calibration was carried out using CRMs (Dickson CO₂). Duplicate measurements were made for each sample, and the estimate of measurement error was < 0.5 %. TA was corrected for the addition of mercuric chloride.</p>	<p>Surface (0 m) and 60 m depth coverage measurements were made from October 2008 to December 2020.</p> <p>Data are available from the British Oceanographic Data Centre (BODC):</p> <p>https://doi.org/10.5285/50bb1181-960e-58b4-e053-6c86abc0e44f (Findlay et al., 2017) and</p> <p>https://www.westernchannelobservatory.org.uk/C_chem.php (last access: 29 November 2023).</p>

Table A2. Continued.

Data type and column headings	Sampling and analysis method	Data coverage and availability of full data
E1: < 20 µm plankton abundance profiles measured by flow cytometry		
E1_Syn_0m_FCM_cells mL-1 E1_Picoeuk_0m_FCM_cells mL-1 E1_Nanoeuk_0m_FCM_cells mL-1 E1_Cocco_0m_FCM_cells mL-1 E1_Crypto_0m_FCM_cells mL-1 E1_HNAbacteria_0m_FCM_cells mL-1 E1_LNAbacteria_0m_FCM_cells mL-1 E1_Syn_10m_FCM_cells mL-1 E1_Picoeuk_10m_FCM_cells mL-1 E1_Nanoeuk_10m_FCM_cells mL-1 E1_Cocco_10m_FCM_cells mL-1 E1_Crypto_10m_FCM_cells mL-1 E1_HNAbacteria_10m_FCM_cells mL-1 E1_LNAbacteria_10m_FCM_cells mL-1 E1_Syn_20m_FCM_cells mL-1 E1_Picoeuk_20m_FCM_cells mL-1 E1_Nanoeuk_20m_FCM_cells mL-1 E1_Cocco_20m_FCM_cells mL-1 E1_Crypto_20m_FCM_cells mL-1 E1_HNAbacteria_20m_FCM_cells mL-1 E1_LNAbacteria_20m_FCM_cells mL-1 E1_Syn_30m_FCM_cells mL-1 E1_Picoeuk_30m_FCM_cells mL-1 E1_Nanoeuk_30m_FCM_cells mL-1 E1_Cocco_30m_FCM_cells mL-1 E1_Crypto_30m_FCM_cells mL-1 E1_HNAbacteria_30m_FCM_cells mL-1 E1_LNAbacteria_30m_FCM_cells mL-1 E1_Syn_40m_FCM_cells mL-1 E1_Picoeuk_40m_FCM_cells mL-1 E1_Nanoeuk_40m_FCM_cells mL-1 E1_Cocco_40m_FCM_cells mL-1 E1_Crypto_40m_FCM_cells mL-1 E1_HNAbacteria_40m_FCM_cells mL-1 E1_LNAbacteria_40m_FCM_cells mL-1 E1_Syn_60m_FCM_cells mL-1 E1_Picoeuk_60m_FCM_cells mL-1 E1_Nanoeuk_60m_FCM_cells mL-1 E1_Cocco_60m_FCM_cells mL-1 E1_Crypto_60m_FCM_cells mL-1 E1_HNAbacteria_60m_FCM_cells mL-1 E1_LNAbacteria_60m_FCM_cells mL-1 Columns 45–86	Measurements were taken fortnightly (conditions permitting). Most samples were analysed in triplicate (phytoplankton and bacteria) for the surface (0 m), whereas single samples were used for all other depths. Vertical profiles of the mean abundance of groups of microbial plankton, expressed as cells per millilitre, were measured using flow cytometry (BD Accuri C6 flow cytometer). The groups quantified are divided into phytoplankton and heterotrophs. The phytoplankton groups quantified are as follows: Syn – <i>Synechococcus</i> sp. (cyanobacteria); Picoeuk – picoeukaryotes (smaller than 3 µm); Crypto – cryptophytes; Cocco – coccolithophores; and Nanoeuk – nanoeukaryotes not already mentioned (2–20 µm). The heterotrophs quantified are as follows: HNAbacteria – heterotrophic bacteria with relatively high nucleic acid content and LNAbacteria – heterotrophic bacteria with relatively low nucleic acid content.	Data span from March 2014 to October 2021. The source dataset is available from https://www.westernchannelobservatory.org.uk/data.php (last access: 29 November 2023).
E1 + L5: combined Young Fish Trawl (YFT)		
E1+L5_Calanus sp_YFT_No.4000m-3 E1+L5_Pilchard eggs_YFT_No.4000m-3 E1+L5_Other fish eggs_YFT_No.4000m-3 E1+L5_Clupeidae larvae_YFT_No.4000m-3 E1+L5_Other fish larvae_No.4000m-3 Columns 87–91	Although the net design and methods of deployment have changed on several occasions, care has been taken to ensure that sampling characteristics have not altered appreciably. The 1 m ² Young Fish Trawl (YFT) fitted with a 700 µm knitted mesh was hauled for 20 min in an oblique profile to an ideal depth of ~ 5 m above the seabed (Ostle et al., 2021). The samples were preserved in 4 % buffered formalin and analysed as soon as possible after collection using a WILD M5 binocular microscope. The volume of filtered water was calculated using flow data recorded by a flowmeter fitted across the net mouth. Results are standardized to the number of individuals per 4000 m ³ in order to mitigate historical changes in sampling gear and deployment. A comprehensive summary of these macroplankton sampling methods and analysis is given in Southward et al. (2005). Note that the reader should be aware of zero values within this dataset; generally, these are true zeros, but this is not necessarily the case in all instances. This is being checked and will be addressed in future versions of the dataset.	Data span from 1924 to 1940, from 1945 to 1987 and from 2001 to 2013. Source data are available from the Data Archive for Seabed Species (DASSH): https://doi.org/10.17031/1636 (Marine Biological Association, 2019).

Table A2. Continued.

Data type and column headings	Sampling and analysis method	Data coverage and availability of full data
E1: recreational captures of blue shark (<i>Prionace glauca</i>)		
E1_ <i>Prionace glauca</i> captures_recreational anglers out of Looe_individuals E1_ <i>Prionace glauca</i> catch per unit effort_recreational anglers out of Looe_captures.trip ⁻¹ Columns 92–93	The Pat Smith database is a collaboration between the Shark Angling Club of Great Britain (SACGB) and the Sportfishing Club of the British Isles (SCBI). It is a collation of information records kept by the SACGB. Recreational angling trips stem from the port of Looe, Cornwall, within a 10 nmi radius of E1. The data presented here are for years when monthly log book information is currently available. The data record of 64 287 captures comprises 32 906 trips from the port in 200 monthly periods between 1958 and 2021. Since 1998, all captures have been released. Data presented are the total number of captures in a given month, and the average catch per unit effort (as captures per trip).	Data span from 1958 to 1971 and from 1997 to 2021. Annual data are available for all years (1953–2022) from Simon Thomas https://www.researchgate.net/publication/372251839 (last access: 29 November 2023).

Author contributions. AJM and AA co-ordinated the project. AJM compiled the monthly database from the component datasets. AA, RLA, RB, IB, ESF, HSF, CLM, AJM, CO, PJS, TJS, GAT, KT, ST and CEW submitted the individual datasets. AJM and AA prepared the manuscript with contributions from HSF, GAT, CEW, SR, RT, AMQ, RLA, RB, ESF, TJS, KT and CO. All authors are either current or previous producers of the component datasets.

Competing interests. The contact author has declared that none of the authors has any competing interests.

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