Mechanisms for a nutrient-conserving carbon pump in a seasonally
stratified, temperate continental shelf sea

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Abstract
Continental shelf seas may have a significant role in oceanic uptake and storage of carbon dioxide (CO$_2$) from the atmosphere, through a ‘continental shelf pump’ mechanism. The northwest European continental shelf, in particular the Celtic Sea (50°N 8°W), was the target of extensive biogeochemical sampling from March 2014 to September 2015, as part of the UK Shelf Sea Biogeochemistry research programme (UK-SSB). Here, we use the UK-SSB carbonate chemistry and macronutrient measurements to investigate the biogeochemical seasonality in this temperate, seasonally stratified system. Following the onset of stratification, near-surface biological primary production during spring and summer removed dissolved inorganic carbon and nutrients, and a fraction of the sinking particulate organic matter was subsequently remineralised beneath the thermocline. Water column inventories of these variables throughout 1.5 seasonal cycles, corrected for air-sea CO$_2$ exchange and sedimentary denitrification and anammox, isolated the combined effect of net community production (NCP) and remineralisation on the inorganic macronutrient inventories. Overall inorganic inventory changes suggested that a significant fraction (>50%) of overall NCP of around 3 mol-C m$^{-2}$ yr$^{-1}$ appeared to be stored within a long-lived organic matter (OM) pool with a lifetime of several months or more. Moreover, transfers into and out of this pool appeared not to be in steady state over the one full seasonal cycle sampled. Accumulation of such a long-lived and potentially C-rich OM pool is suggested to be at least partially responsible for the estimated net air-to-sea CO$_2$ flux of ~1.3 mol-C m$^{-2}$ yr$^{-1}$ at our study site, while providing a mechanism through which a nutrient-conserving continental shelf pump for CO$_2$ could potentially operate in this and other similar regions.
1. Introduction

The ocean is an important buffer for atmospheric carbon dioxide (CO$_2$), naturally storing a large pool of dissolved inorganic carbon (C$_T$), and also absorbing about a quarter of annual anthropogenic CO$_2$ emissions (Le Quéré et al., 2009) and thereby mitigating their climatic impacts (IPCC, 2013). The timescale on which the ocean’s surface mixed layer equilibrates CO$_2$ with the atmosphere varies spatially, but is typically from a few months to a year (Broecker and Peng, 1974; Jones et al., 2014). Transfer from the surface to the deep interior is a much slower rate-limiting step, which is performed by the solubility, soft tissue, carbonate and mixed-layer ‘pumps’ in the open ocean (Volk and Hoffert, 1985; Dall’Olmo et al., 2016). Tsunogai et al. (1999) first proposed the existence of an additional ‘continental shelf pump’ (CSP) that facilitates C$_T$ transfer from shallow coastal surface waters into the interior ocean, and thereby enhances local oceanic CO$_2$ uptake. The CSP operates through autotrophic conversion of C$_T$ into organic matter (OM) in a continental shelf sea, which drives a compensatory net air-to-sea CO$_2$ flux (Chen and Borges, 2009; Chen et al., 2013; Laruelle et al., 2014). In order to prevent the fixed CO$_2$ from being quickly released back into the atmosphere, the carbon-enriched shelf waters need to be transported into the open ocean, beneath the seasonal thermocline (Tsunogai et al., 1999; Thomas et al., 2004). The Celtic Sea is a seasonally stratified, temperate sea that forms part of the northwest European continental shelf (Fig. 1). Previous studies of its surface waters have shown that it acts as a net sink of atmospheric CO$_2$ (Frankignoule and Borges, 2001; Marrec et al., 2015), which is typical for this type of sea (Borges, 2005; Laruelle et al., 2014). There must therefore be a net flux of carbon (C) out of the shelf sea water column, for example through OM export to sediments, or advective exchange with the open ocean. Celtic Sea sediments are unlikely to form an important component of this C sink, as they are predominantly sandy material with relatively low organic matter accumulation (de Haas et al., 2002; Suykens et al., 2011; Diesing et al., 2017), but physical mechanisms that could transport shelf waters into the open ocean have been identified here (Cooper and Vaux, 1949; Shapiro et al., 2003; Ivanov et al., 2004) and in other similar shelf environments (Álvarez-Salgado et al., 2001). In either case, the question arises of how the nutrient supply is sustained that drives the net C uptake. As essential nutrients such as dissolved inorganic nitrogen and soluble reactive phosphorus (DIN and SRP) are also incorporated into OM, but not replenished through air-sea exchange, the net removal of C should be accompanied by a corresponding loss of N and P from the shelf waters (Thomas et al., 2004; Bozec et al., 2006). Further N may be lost from the shelf
waters through denitrification and/or anammox in the shelf sediments (Hydes et al., 2004; Kitidis et al., 2017). Thus, in order to sustain the shelf sea nutrient inventories in the presence of net CO$_2$ uptake from the atmosphere, any incoming nutrient supply would therefore need to be accompanied by a stoichiometric deficit of carbon relative to outgoing waters. The mechanism by which this occurs remains uncertain. One possibility is that riverine inputs provide high-nutrient, low-C waters to the shelf sea. There are also high-nutrient waters in the Irish Sea to the northeast of our study region (Gowen et al., 2002). However, these influences are not thought to be substantial for the Celtic Sea, as on-shelf lateral circulation is relatively slow (Pingree and Le Cann, 1989), and most of the riverine nutrient inputs are quickly removed by biogeochemical processes close to land (Ruiz-Castillo et al., 2017, this issue). In general, riverine nutrient inputs to shelf seas are small relative to inputs at the continental margin, for example through upwelling and internal waves at the shelf break (Wollast, 1998). Also, rivers flowing into the Celtic Sea (e.g. from southern Ireland) typically have high total alkalinity ($A_T$) due to chalk and limestone bedrock in their catchments (McGrath et al., 2015). This high $A_T$ supports high $C_T$ concentrations, so the required C deficit relative to N and P is unlikely to occur in these waters.

Here, we investigate the hypothesis that OM storage coupled with differences in the stoichiometry of OM remineralisation relative to its production could provide the stoichiometric inconsistency required to sustain the nutrient supply in the Celtic Sea, as has been suggested to occur in similar shelf seas (e.g. Álvarez-Salgado et al., 2001; Bozec et al., 2006). This would require the stoichiometric C:N:P ratio for remineralisation to have a lower C coefficient than in the equivalent ratio for net community production (NCP). The shelf nutrient inventory could therefore be sustained through some combination of enhanced remineralisation of non-C nutrients, or the build-up of a stock of C-rich OM that could be transported into the adjacent open ocean prior to complete remineralisation, thereby acting as a nutrient-conserving CO$_2$ sink. Our approach is to investigate the distributions and inventories of $C_T$ and dissolved inorganic nutrients, which record the integrated effects of NCP and OM remineralisation, along with other processes such as air-sea CO$_2$ exchange. We use new seasonally resolved $C_T$ and nutrient observations in this way to estimate the annual magnitude and stoichiometry of NCP and remineralisation at a site in the central Celtic Sea. We subsequently infer seasonal and interannual changes in OM inventories and stoichiometry, and consider mechanisms that could support C export while conserving nutrients, and therefore a CSP.
2. Methods

2.1. Research cruises

Seawater samples were collected during 10 research cruises to the Celtic Sea on the UK research ships RRS *Discovery* and RRS *James Cook*, from March 2014 to August 2015 (Supp. Table S1). All of our datasets are freely available from the British Oceanographic Data Centre.

2.1.1. Marine carbonate system

Samples for \( C_T \) and \( A_T \) were collected via silicone tubing into 250 ml borosilicate glass bottles either from the samplers (Ocean Test Equipment) positioned on a CTD rosette frame, or from the ship’s underway seawater supply, following an internationally established protocol (Dickson et al., 2007). Each bottle was sealed shut with a greased ground glass stopper after introducing a 2.5 ml air headspace and sterilising the sample with 50 µl of saturated mercuric chloride solution. All samples were stored in the dark until analysis.

Measurements of \( C_T \) and \( A_T \) were carried out at the University of Southampton between June 2014 and January 2016, using several different instruments. The results were calibrated using measurements of batches 128, 135, 136, 138, 141, 142, 144, 146 and 148 of seawater certified reference material (CRM) obtained from A. G. Dickson at Scripps Institution of Oceanography, San Diego, USA (Dickson et al., 2003).

Seawater \( C_T \) was measured using either the Versatile INstrument for the Determination of Total inorganic carbon and titration Alkalinity (VINDTA 3C, #024 and #038, Marianda, Germany) or the Dissolved Inorganic Carbon Analyzer AS-C3 (Apollo SciTech Inc., USA). Both of these instruments first acidify a seawater subsample with excess 10% phosphoric acid, to convert all \( C_T \) into aqueous CO\(_2\). Nitrogen gas is then bubbled through to carry the CO\(_2\) to a detector. In the VINDTA 3C, the detection is by coulometric titration (CM5014 CO\(_2\) coulometer, UIC Inc., USA), while the AS-C3 uses infrared absorption (LI-7000 CO\(_2\)/H\(_2\)O Analyzer, LI-COR, USA). Based on CRM measurements throughout the entire UK-SSB programme, the 1σ precision for \( C_T \) was \( \pm 2.6 \) µmol kg\(^{-1}\) for the VINDTA 3C (number of CRM measurements \( n = 547 \)), and \( \pm 4.0 \) µmol kg\(^{-1}\) for the Apollo AS-C3 \( (n = 135) \).

The \( A_T \) was measured using either the VINDTA 3C or the Total Alkalinity Titrator AS-ALK2 (Apollo SciTech Inc., USA). Both instruments determine \( A_T \) by an open-cell, potentiometric titration of a seawater subsample with 0.1 M hydrochloric acid. The \( A_T \) values from the
VINDTA 3C measurements were recalculated using a modified Gran plot approach (Humphreys, 2015a) as implemented by Calkulate v0.1.2 (freely available from https://github.com/mvdh7/calkulate). The CRM measurements through the entire UK-SSB programme indicated a 1σ precision for $A_T$ of ±2.7 µmol kg$^{-1}$ for the VINDTA 3C ($n = 543$), and ±3.9 µmol kg$^{-1}$ for the Apollo AS-ALK2 ($n = 109$).

We performed a cross-over analysis using XOVER v1.0.1 (Humphreys, 2015b), which is freely available from https://github.com/mvdh7/xover, to assess our $C_T$ and $A_T$ measurement accuracy relative to historical data in GLODAPv2 (Olsen et al., 2016). This used only data from deeper than 250 m (i.e. off the shelf, beneath the base of the thermocline, and a maximum cross-over distance of 200 km). For $C_T$, there was a mean ± standard deviation (SD) offset of 13.6 ± 7.6 µmol kg$^{-1}$ between the combined SSB dataset and nine GLODAPv2 cruises from 1994 to 2008. Despite the large size of this offset, it does not indicate any problem with our measurements, because an ordinary least squares linear regression between the offsets and the sampling date reveal a decrease in offset size at a rate of about 1 µmol kg$^{-1}$ yr$^{-1}$ ($r^2 = 0.66$), corresponding to the anthropogenic $C_T$ increase, and reaching a value of 0 close to the date of the SSB cruises. For $A_T$, the equivalent offset was 6.8 ± 7.1 µmol kg$^{-1}$ for nine cruises from 1983 to 2008, which is close to the minimum adjustment limit of 6 µmol kg$^{-1}$ for this variable in GLODAPv2 (Olsen et al., 2016). In general, cruises closer in time to SSB sampling had a smaller $A_T$ offset. We conclude that our measurements are accurate to within their precision relative to historical observations. Indeed, the internal consistency of data within the GLODAPv2 compilation is reported as ±4 and ±6 µmol kg$^{-1}$ for $C_T$ and $A_T$ respectively (Olsen et al., 2016).

2.1.2. Other variables

Macronutrient samples were collected into 60 ml high density polyethylene (HDPE, Nalgene) bottles that had been aged and acid washed before sampling. The measurements (of nitrate, nitrite, ammonium, and phosphate) were carried out at sea during the UK-SSB cruises, immediately after sample collection, using the Plymouth Marine Laboratory 5-channel Bran and Luebbe AAIII system, following Woodward and Rees (2001). The instrument was calibrated using in-house nutrient standards, and accuracy was monitored by regular comparisons with Nutrient Reference Material obtained from KANSO Technos (Japan). We define dissolved inorganic nitrogen (DIN) as the sum of the concentrations of nitrate, nitrite...
and ammonium, and soluble reactive phosphorus (SRP) as the phosphate concentration. The 1σ precisions were ± 0.13 μmol kg$^{-1}$ and ± 0.0073 μmol kg$^{-1}$ for DIN and SRP respectively.

Water column hydrography was measured by Sea-Bird conductivity-temperature-depth (CTD) sensors attached to the sampling rosette frame, to determine the temperature ($T$ in °C), practical salinity ($S$) and depth for each discrete biogeochemical sample. The sea surface hydrography ($T$ and $S$) was measured continuously by Sea-Bird sensors through the underway seawater supply at a nominal depth of 6 m. The $T$ and $S$ sensor data for both the underway and CTD systems were calibrated using a set of discrete samples collected during each cruise. We consider the 1σ uncertainty in $S$ to be ± 0.002 (Le Menn, 2011). Further information on the sensor measurements and processing can be found in the cruise reports.

### 2.1.3. Unit conversions

The $C_T$ and $A_T$ measurements were determined in units of μmol kg$^{-1}$, while DIN and SRP were reported in mmol m$^{-3}$ under analysis conditions. We therefore converted the latter two into μmol kg$^{-1}$ assuming a constant analysis temperature of 20°C. For the column inventory analysis, we then converted the $C_T$, DIN and SRP measurements from μmol kg$^{-1}$ into mmol m$^{-3}$ using the *in situ* density. All densities for these calculations were determined using the Gibbs-SeaWater (GSW) Oceanographic Toolbox v3.05.5 (McDougall and Barker, 2011).

### 2.2. Central Celtic Sea mooring

A temperature-salinity (TS) chain (a series of sensors positioned vertically along a cable moored to the seafloor) was also deployed on a Cefas SmartBuoy mooring at CCS from March 2014 to July 2015 (Wihsgott et al., 2016). This measured seawater $T$ and $S$ at approximately 2.5 m vertical resolution throughout the water column (i.e. from c. 5 to 145 m) and at 5 minute intervals. A combination of sensor types were used: Star Oddi DST Centi temperature loggers, Star Oddi Starmon mini-underwater temperature recorder and RBRsolo T temperature logger, along with SBE16+ SeaCAT CTD and SBE37 MicroCAT CTD sensors. We calculated mixed layer depth ($z_{ml}$ in m) as the shallowest depth where the potential density increased by 0.02 kg m$^{-3}$ from its value at 10 m.

### 2.3. Auxiliary datasets

We obtained 0.75°×0.75° gridded atmospheric/surface ocean reanalysis data at 3-hourly temporal resolution from the ERA-Interim data product (Dee et al., 2011), which is produced by the European Centre for Medium-Range Weather Forecasts (ECMWF). Data were
downloaded from ECMWF on 1st August 2016 for the following variables, over the period from 1st March 2014 to 31st December 2015: sea surface temperature (SST in °C), mean sea level pressure ($P_{\text{baro}}$ in bar), 10 metre U wind component ($U_{10u}$ in m s$^{-1}$), 10 metre V wind component ($U_{10v}$ in m s$^{-1}$), and 2 metre dewpoint temperature ($T_{dp}$ in °C). We calculated the wind speed at 10 m above the sea surface ($U_{10}$, in m s$^{-1}$) from its components $U_{10u}$ and $U_{10v}$.

2.4. Central Celtic Sea site

Most of the analysis presented here was carried out at CCS, which was the most frequently-occupied location during the UK-SSB sampling campaign. All samples collected within a 12 km radius of 49.4°N, 8.54°W were considered to represent CCS (Fig. 1). This radius was selected such that sufficient data were captured to constrain the interior biogeochemical seasonality, whilst deviating as little as possible from the site itself. We then manually grouped the CCS observational dataset based on the sampling date: we selected a set of date boundaries that minimised the width of each time interval, while ensuring that there were enough data within each interval to determine the vertical profile of each biogeochemical variable, resulting in 15 time intervals (Table 1).

Vertical profiles of biogeochemical variables at each site were typically homogeneous during the winter months, and had a two-layer shape during the stratified summer. As such, it was possible to fit error functions to many variables, in order to objectively determine values for each variable in the deep and surface layers, along with the depth of the divide. We thus generated a fit to all data for each variable at each time interval at each site using an equation of the form:

$$v = v_0 + v_1 \text{erf} \left( \frac{z - z_0}{z_1} \right)$$  \hspace{1cm} (1)

where $v$ and $z$ are the values and depths respectively of the measurements of each variable, $\text{erf}$ is the Gauss error function, and $v_0$, $v_1$, $z_0$ and $z_1$ are coefficients that were adjusted to find the least-squares best fit to these data. These coefficients quantify useful properties: $z_0$ is the depth of the boundary between the surface and deep layers; $z_1$ indicates how sharp or diffuse the boundary is, with a smaller value indicating a transition over a narrower depth range; and the surface and deep layer values of the variable $v$ are given by ($v_0 - v_1$) and ($v_0 + v_1$) respectively (see Supp. Table S2 and Supp. Figs. S1–S4).
2.5. Air-sea CO$_2$ flux

We calculated $p$CO$_2^{sw}$ from the $A_T$, $C_T$, $S$, $T$, pressure, silicate and SRP observations using version 1.1 of the CO$_2$SYS program for MATLAB (van Heuven et al., 2011). We calculated Free scale pH in the same way (Supp. Fig. S5). We used the carbonic acid dissociation constants of Lueker et al. (2000), the bisulfate dissociation constant of Dickson (1990) and the boron:chlorinity ratio of Lee et al. (2010). The uncertainties in the input variables resulted in a 1σ precision of ± 8.1 µatm for calculated $p$CO$_2^{sw}$. Statistical interpolations of these calculated $p$CO$_2^{sw}$ values were limited by low temporal sampling resolution, resulting in key features such as the 2014 spring bloom being partly missed, and therefore inaccurate estimates of the time-integrated air-sea CO$_2$ flux. We therefore created an empirical algorithm to predict the surface $p$CO$_2^{sw}$ time series at CCS from the reanalysis SST data. The algorithm used different equations for different time periods and SST ranges (Table 2) to account for temporal variation in the surface $p$CO$_2^{sw}$-SST relationship resulting from changes in biological production and vertical mixing. The mean ± SD difference between the algorithm-predicted and measured $p$CO$_2^{sw}$ was -4 ± 19 µatm across all 71 measurements above 25 m depth at CCS, which is similar to the uncertainties quoted for published $p$CO$_2^{sw}$ algorithms in this region (e.g. Marrec et al., 2015).

We obtained measurements of the atmospheric CO$_2$ dry air mole fraction ($x$CO$_2$) for the Greenhouse Gases Reference Network Site at Mace Head, County Galway, Ireland (Dlugokencky et al., 2015). Geographically, this is the closest such observation site to our study area. We applied a piecewise cubic Hermite interpolating polynomial (PCHIP) function (Fritsch and Carlson, 1980; Kahaner et al., 1988) to their monthly mean data to predict $x$CO$_2$ in parts per million (ppm) for any given date. We estimated the uncertainty in $x$CO$_2$ values interpolated from the monthly means, ± 2.8 ppm, as the standard deviation of the residuals between all of the original individual measurements and the interpolated $x$CO$_2$ values at the same time points. We then converted $x$CO$_2$ into atmospheric CO$_2$ partial pressure ($p$CO$_2^{atm}$) using data from the ERA-Interim reanalysis (Dee et al., 2011) at CCS. First, we estimated the water vapour pressure ($P_v$ in bar) from $T_{dp}$ (Alduchov and Eskridge, 1996; Lawrence, 2005):

$$P_v = 610.94 \exp\left(17.625 T_{dp}/(243.04 + T_{dp})\right) \times 10^{-5} \tag{2}$$

The atmospheric partial pressure of CO$_2$ ($p$CO$_2^{atm}$ in µatm) was then given by:

$$pCO_2^{atm} = xCO_2 \left( P_{baro} - P_v \right) / 1.01325 \tag{3}$$
We calculated the air-sea CO$_2$ flux ($F$, in $\mu$mol m$^{-2}$ hr$^{-1}$) following Wanninkhof (2014), using 3-hourly resolution data from ECMWF, the algorithm-predicted $p$CO$_2^{sw}$ (Table 2), and $p$CO$_2^{atm}$ calculated from $x$CO$_2$ (Eq. 3). The flux equation is:

$$F = k \alpha (pCO_2^{sw} - pCO_2^{atm}) \quad (4)$$

where $\alpha$ is the solubility of CO$_2$, evaluated following Weiss (1974) and converted to units of mol m$^{-3}$ atm$^{-1}$ using the GSW toolbox (McDougall and Barker, 2011), and $k$ is the gas transfer velocity in m hr$^{-1}$:

$$k = 0.00251 U_{10}^2 (660/D)^{0.5} \quad (5)$$

where $D$ is the dimensionless Schmidt number for CO$_2$ at the surface layer seawater temperature (Wanninkhof, 2014). The flux calculations were carried out in MATLAB (MathWorks) using the ‘co2flux’ program available from https://github.com/mvdh7/co2flux.

### 2.6. Column inventories

The column inventory of each biogeochemical variable was determined at each time interval by integrating the fitted profile for the variable (in mmol m$^{-3}$) from the surface to the seafloor depth, taken as the mean seafloor depth across all CCS sampling stations ($Z = 146.9$ m). We then attempted to remove the influence of processes other than NCP and remineralisation on the column inventories. To correct for air-sea gas exchange, we subtracted the air-sea CO$_2$ flux integrated between each time point and winter 2014 ($C_{atm}$) from each time point’s raw $C_T$ inventory. Hydes et al. (2004) suggest a constant denitrification rate of 0.1 mmol-N m$^{-2}$ day$^{-1}$ for the Celtic Sea. We therefore subtracted the product of this rate and the number of days between each time point and winter 2014 from the DIN inventory at each time point to correct for the potential loss of DIN through denitrification ($DIN_{dnf}$). Kitidis et al. (2017) more recently reported a similar mean flux of DIN from sediments into the seawater due to denitrification/anammox processes at a site near CCS. They identified some seasonal variability in this value, but we did not include this effect as it was too small to alter our analysis and conclusions.

### 2.7. Uncertainty analysis

The uncertainties in all of the final results, reported as 95% confidence intervals, were calculated using Monte Carlo simulations. Explicitly, we replicated the column inventory analysis 1000 times, each time adding different random offsets to each individual
measurement. These offsets were normally distributed, with a SD equal to the $1\sigma$ analytical uncertainty for each measurement. The random noise was added both to the measurements of each variable. We report the 95% uncertainty in each column inventory as double the SD of all of its replicates.

2.8. Salinity correlations

We quantified the seasonally changing patterns in $C_T$, DIN and SRP remineralisation across the Celtic Sea by determining their correlations with practical salinity in the deep layer. For these correlations, we used only data satisfying the following conditions: (1) salinity $> 35.2$, to exclude near-coastal waters with nutrient-salinity relationships different from the majority of the shelf; (2a) where seafloor depth $< 200$ m (i.e. on-shelf), sample depth $> 60$ m, to exclude the surface layer where DIN and SRP are reduced to virtually zero in the summer months; and (2b) where seafloor depth $> 200$ m (i.e. off-shelf), $150$ m $<$ sample depth $< 200$ m, again to exclude the surface layer (which was deeper off-shelf) and also deeper waters that were less strongly affected by seasonal remineralisation. The potential density of the off-shelf 150 to 200 m layer throughout the year was similar to the on-shelf winter potential density, supporting our choice of this as the off-shelf endmember for the correlations. Summary statistics for these correlations are provided in Table 3, and scatter plots are show by Supp. Figs. S12–S14.
3. Results and discussion

3.1. Central Celtic Sea site

3.1.1. Air-sea CO\(_2\) flux

Few studies have examined the Celtic Sea inorganic carbon cycle in detail, but those that have were mostly focused on the surface layer and air-sea CO\(_2\) exchange. Frankignoulle and Borges (2001) used a compilation of surface ocean \(p\text{CO}_2\)\text{sw} data to demonstrate that the northwest European continental shelf seas are a net sink for atmospheric CO\(_2\), at a rate between 1.8 and 2.9 mol m\(^{-2}\) yr\(^{-1}\). This result was quantitatively supported by a more recent model study (Wakelin et al., 2012). However, other studies have found smaller net air-sea CO\(_2\) fluxes for this region. Borges et al. (2006) compiled published datasets across the entire northwest European shelf, and Marrec et al. (2015) generated predictive algorithms for Celtic Sea \(p\text{CO}_2\)\text{sw} from variables including sea surface temperature, mixed layer depth and chlorophyll-\(a\) concentration. These studies agree that the Celtic Sea is a net sink for atmospheric CO\(_2\), but they suggest a typical rate a little under 1 mol m\(^{-2}\) yr\(^{-1}\). Our analysis also indicated that the CCS site acted as a net atmospheric CO\(_2\) sink: the mean air-to-sea CO\(_2\) flux at CCS from 21\(^{st}\) March 2014 (i.e. the start of the UK-SSB sampling period) to the same date in 2015 was 1.3 ± 0.3 mol m\(^{-2}\) yr\(^{-1}\) (Fig. 2). The magnitude of this flux is consistent with previous Celtic Sea studies, although it is towards their lower end. A separate analysis of \textit{in situ} CO\(_2\) uptake (photosynthesis) and production (plankton respiration) data also found that CCS was a net atmospheric CO\(_2\) sink, as primary production was typically greater than community respiration (García-Martín et al., 2017, this issue).

The \(\Delta p\text{CO}_2\) was vertically uniform and close to atmospheric equilibrium during March-April. Its seasonal pattern was similar to \(C_T\), with CO\(_2\) undersaturation in the surface layer during the spring-summer. The surface \(p\text{CO}_2\)\text{sw} variability was dominated by biological \(C_T\) uptake and release, rather than seasonal seawater heating and cooling, which would have been expected to induce an increase in \(p\text{CO}_2\)\text{sw} (and therefore \(\Delta p\text{CO}_2\)) during the warmer summer months. Indeed, the amplitude of the SST seasonal cycle was about 10°C. Alone, this would have driven a winter-summer \(p\text{CO}_2\)\text{sw} increase of about 200 μatm (Takahashi et al., 2009), whereas we observed a c. 85 μatm decline during this period (Fig. 2a). Other factors that could affect the \(\Delta p\text{CO}_2\) cycle are atmospheric \(p\text{CO}_2\), and seawater \(A_T\). The amplitude of seasonal atmospheric \(p\text{CO}_2\) cycle was an order of magnitude smaller than that of \(p\text{CO}_2\)\text{sw}, while \(A_T\) was not noticeably affected by seasonal stratification, remaining relatively
homogeneous both laterally and vertically throughout the Celtic Sea for the duration of the SSB sampling campaign (Supp. Fig. S8). The seasonal variability of atmospheric $p$CO$_2$ and $A_T$ therefore did not significantly influence that of $\Delta$pCO$_2$.

The algorithm that we created to predict surface $p$CO$_2^{sw}$ (Table 2) produced an annual range of about 85 µatm, similar to previous studies (e.g. Marrec et al., 2015). However, there were differences in the seasonal pattern. Firstly, we found that the $p$CO$_2^{sw}$ had two distinct minima during 2014, first in May-June and then a weaker minimum in November-December associated with an autumn phytoplankton bloom (Wihsgott et al., 2017, this issue). Our 2015 data end too early in the seasonal cycle to determine whether these dual minima were repeated. Secondly, Marrec et al. (2015) projected that $\Delta$pCO$_2$ should be positive for approximately a third of each year, during the winter months, reaching values up to about 30 µatm. However, neither our algorithm nor our observations indicate such consistently high $\Delta$pCO$_2$ values. We did not collect any carbonate chemistry samples between November and March, so we cannot be certain which is correct. However, the observational $p$CO$_2^{sw}$ data from SOCAT (Bakker et al., 2016) used by Marrec et al. (2015) to test their algorithm were in atmospheric equilibrium during this season, rather than supersaturated as predicted, for about half of the years that they considered. This winter supersaturation may therefore be an intermittent phenomenon, but our dataset is unable to identify whether it occurred between 2014 and 2015.

The most uncertain part of our air-sea CO$_2$ flux algorithm was between autumn 2014 and winter 2015, due to a lack of observations during that period. Comparison of the pattern of $p$CO$_2^{sw}$ (Fig. 2a) with other studies (e.g. Marrec et al., 2015) and our understanding of marine carbonate system drivers suggests that the $p$CO$_2^{sw}$ predicted by the algorithm for this period is likely to underestimate the true value. Consequently, we consider the air to sea CO$_2$ flux calculated for this period as a maximum estimate.

3.1.2. Inorganic inventory changes

Vertical profiles of $C_T$, DIN, SRP and related hydrographic variables were constrained at two roughly equivalent time points both in 2014 and 2015, referred to as ‘winter’ (i.e. March-April, pre-spring bloom) and ‘summer’ (August). Between these, there was a single ‘late spring’ profile in June 2014, and a series of ‘in-bloom’ profiles (April-May) captured the progression of the 2015 spring bloom at a relatively high temporal resolution, followed by an ‘early spring’ profile in May 2015. There was an ‘autumn’ (November) profile in 2014 only.
Direct comparisons between the two years can therefore be drawn from comparing winter and summer, while the additional points can be used to more finely resolve the seasonal changes.

The TS chain measurements clearly illustrate the seasonal cycle of surface warming and stratification followed by winter mixing (Fig. 3a). The CCS site was vertically mixed during the winter, but physical and biogeochemical variables exhibited a two-layer stratified structure during the spring and summer. Most variables were vertically homogeneous in winter each year, after which $C_T$, DIN and SRP declined strongly in the surface layer (i.e. shallower than 40-50 m) through the spring and summer (Fig. 3b-d), through drawdown associated with NCP. DIN reached virtually zero (i.e. beneath the detection limit of ~0.02 mmol m$^{-3}$), while SRP declined to a minimum of about 0.05 mmol m$^{-3}$, consistent with previous studies that have identified DIN as the limiting nutrient in this region (e.g. Davis et al., 2014). The concentrations of these variables increased in the deep layer over the same period, consistent with remineralisation of sinking OM. Stratification began to break down towards the autumn, and the water column was reset to a vertically homogeneous state by the following winter.

Stratification had a less coherent effect on $S$ (as measured by the TS chain), but here we saw other forms of variability (Fig. 3e). In particular, there was a notable increase in deep layer $S$ (i.e. below about 50 m) from August to December 2014. This suggests that open ocean waters increased in influence relative to on-shelf waters during this time, as no in situ processes can modify deep layer $S$ to this extent. The DIN and SRP increased disproportionately in the deep layer during the same period as the aforementioned high $S$ anomaly (Fig. 3c), suggesting that increasing nutrient concentrations relative to $C_T$ at this time may have been associated with the advective flux that elevated $S$, rather than in situ OM remineralisation.

Deep layer salinity (i.e. beneath c. 50 m) increased from spring 2014 through to December 2014 (Fig. 3e) by an amount that cannot be accounted for by any in situ process. The DIN and SRP also increased disproportionately in this deep layer during the same period (Fig. 3c-d), suggesting that increasing nutrient concentrations at this time may have been associated with an advective flux that elevated $S$, rather than in situ OM remineralisation. In support of this, Ruiz et al. (2017, this issue) describe a shelf-wide estuarine-type circulation that advects saline, nutrient-rich waters on to the shelf during the stratified summer months. They estimate
that 70-80% of the increase in deep layer DIN observed at CCS since the fully mixed winter state could be supplied by this physical transport mechanism.

### 3.1.3. Production and remineralisation

During the first half of each year, when stratification was increasing, changes in $C_T$, DIN and SRP profiles between successive time points can reveal the depth ranges over which NCP and net remineralisation occurred (e.g. Fig. 4). For intervals when surface layer concentrations declined, the average NCP rate can therefore be estimated by integrating the change in $C_T$ only over the depth range where $C_T$ declined and correcting the result for air-sea CO$_2$ exchange, which can be assumed to affect only the surface layer when the water column is stratified. Similarly, deep layer increases in $C_T$ can be attributed to remineralisation. The same procedure can be carried out for DIN and SRP, with the remineralisation calculation for DIN corrected for denitrification and/or anammox (Hydes et al., 2004; Kitidis et al., 2017), and no gas exchange correction (Table 4). The main spring bloom in 2014 resulted in a mean NCP of 35.9 ± 2.3 mmol-C m$^{-2}$ d$^{-1}$ from winter to late spring, which is consistent with an NCP estimate for the seasonally-stratified North Sea of 16-46 mmol m$^{-2}$ d$^{-1}$ for this season (Bozec et al., 2006). This represents a total of 3.08 ± 0.20 mol-C m$^{-2}$ $C_T$ conversion into OM since the mixed winter state, integrated over the period between our winter and late spring 2014 time points (i.e. 86 days; Table 1). The equivalent DIN and SRP drawdowns were 342 ± 7 mmol m$^{-2}$ d$^{-1}$ (Fig. 4a) and 20.74 ± 0.36 mmol m$^{-2}$ d$^{-1}$ respectively, resulting in an overall C:N:P for NCP of about 117:13:0:0.79. Corresponding remineralisation of $C_T$, DIN and SRP within the subsurface layer was 0.78 ± 0.50 mol m$^{-2}$, 85 ± 73 mmol m$^{-2}$ and 8 ± 99 mmol m$^{-2}$, with C:N:P therefore 117:12.7:1.24. This indicates that 20–40% of the NCP was remineralised within the sub-surface (presumably following sinking of the corresponding OM), while 60–80% must have remained within some OM pool, assuming negligible net advection (Pingree and Le Cann, 1989; Brown et al., 2003). The spring bloom period was followed by much weaker NCP and net water column remineralisation from late spring to summer, and then vertical mixing into the autumn and winter (Fig. 4b-c), during which period the decline in deep layer $C_T$ was due to mixing with low-$C_T$ surface waters, not NCP.

The spring bloom of the following year was sampled at higher temporal resolution, during cruise DY029 in April 2015. From winter to early spring 2015 (i.e. 61 days; Table 1), the mean NCP determined from $C_T$ profiles and air-sea CO$_2$ exchange was 30.5 ± 2.3 mmol-C m$^{-2}$ d$^{-1}$, comparable to the previous year, estimates from nearby similar regions (Bozec et al.,
(Seguro et al., 2017, this issue). This NCP integrated from the winter to early spring 2015 time points (i.e. 139 days; Table 1) represented 1.86 ± 0.14 mol-C m⁻² of OM production, compared with 3.08 ± 0.20 mol-C m⁻² for the roughly equivalent period (winter to late spring) in 2014. However, further NCP from early spring to summer in 2015 brought the total NCP since winter to 3.30 ± 0.13 mol-C m⁻², similar to the total value for the part of 2014 with positive NCP. The corresponding DIN- and SRP-derived values (from winter to summer 2015) were 343.4 ± 3.9 and 20.86 ± 0.22 mmol m⁻² respectively, resulting in a C:N:P for NCP of about 117:12.2:0.74, similar to the previous year (i.e. 117:13.0:0.79), but with lower N and P coefficients in 2014, or equivalently more C-rich OM production relative to typically assumed (Redfield, 1934; Redfield et al., 1963) and measured (Anderson and Sarmiento, 1994) ratios. Therefore the total amount of OM generated through NCP appeared to be similar during the spring bloom each year, although this may have occurred more rapidly during 2014. Consequently, although the winter and summer time points were at a very similar time for both years, it may be more appropriate to compare ‘winter to late spring 2014’ with ‘winter to summer 2015’. Remineralisation from winter to summer 2015 occurred at C:N:P of 117:11.7:0.88, and represented 24–35% of NCP, in both ways similar to 2014. Both the production and remineralisation of OM were C-rich relative to the likely limiting nutrient, with C:N ranging from 9 to 10 compared with typical values of 6.6 to 7.3 (Redfield, 1934; Redfield et al., 1963; Anderson and Sarmiento, 1994).

### 3.1.4. Total column inventories

The preceding section’s approach of separating surface NCP from deep remineralisation was less useful from summer onwards, once stratification had begun to break down. At this time, both surface and deep layer Cتراث and nutrient inventories were influenced by vertical mixing, along with remineralisation throughout the water column. It is therefore more appropriate to consider changes in the total inventories integrated vertically across the entire water column. Remineralisation of both Cتراث and nutrients continued after summer 2014, first with apparent excess DIN generation until autumn 2014, and then excess Cتراث until winter 2015 (Fig. 5a-b). We note that the excess N period coincides with the previously-noted apparent advective flux of high salinity to CCS, high DIN waters in the deep layer, so this may have driven the unexpected and transient high DIN, rather than remineralisation (Fig. 5c). However, this high S water had disappeared by winter 2015, due to vertical mixing and shut-down of the summer
estuarine-type circulation (Ruiz-Castillo et al., 2017, this issue). The overall trajectory from summer 2014 to winter 2015 is therefore probably representative of net remineralisation, presumably ongoing from some long-lived OM pool.

From late spring 2014 to winter 2015, remineralisation returned a further 2.17 ± 0.43 mol-C m⁻² C_T, 169 ± 50 mmol-N m⁻² DIN and 3.27 ± 0.88 mmol-P m⁻² SRP to the water column, corresponding to a C:N:P ratio of 117:9.1:0.2, with substantially lower N and P coefficients and hence higher C:N and C:P than the NCP stoichiometry. Adding the additional apparent late year remineralisation to that experienced in the deep layer from winter to late spring 2014 resulted in an overall remineralisation C:N:P of 117:10.0:0.5. This corresponded to 96 ± 22 %, 72 ± 26 % and 56 ± 479 % of the OM production of C_T, DIN and SRP respectively in the 2014 spring bloom being remineralised by winter 2015. In other words, almost all of the C_T that was biologically fixed in the 2014 spring bloom was remineralised by winter 2015, but only about three-quarters and half of the fixed DIN and SRP respectively were remineralised over the same period, while noting that the uncertainty in the SRP value is too great for us to speculate about it further. As we previously described (Section 3.1.1), the air-sea CO₂ flux value that was used to correct these inventories was likely an overestimate from autumn 2014 to winter 2015. Therefore the C_T result is a minimum estimate, and it is possible that more C_T was remineralised than was produced by NCP that year at CCS.

These total inventories indicate that the shelf was not operating in steady state from winter 2014 to 2015. The winter inventories of DIN and SRP were 10.3 ± 1.4 % and 10.6 ± 1.1 % lower respectively in 2015 compared to 2014, while that of C_T was 0.36 ± 0.18 % higher (Fig. 5a). The equal (or slightly greater) OM formation during the 2015 spring-summer relative to 2014 was therefore unexpected, because lower nutrient concentrations at the end of the winter mixed state should have supported less NCP. As NCP was the ultimate driver of CO₂ uptake, we would also have expected a lower air-to-sea CO₂ flux following these lower winter nutrient concentrations. This did not occur, because the nutricline was about 10-15 m deeper in 2015 (Fig. 6a), resulting in a greater total N_T conversion into OM despite its lower initial concentration, and matching the behaviour of the thermocline (Fig. 6b). There does not appear to be an obvious mechanism that would drive a deeper nutricline at times of lower surface nutrient concentrations, which would lead summer NCP to be generally consistent across different years. Rather, it appears to be a coincidence that these phenomena balanced each other out for the two years that we observed.
3.1.5. Organic matter stoichiometry

Overall mass balance dictates that changes in the OM composition should mirror the changes in the inorganic inventories (corrected for gas exchange and denitrification/anammox). In Fig. 5b, trajectories between adjacent sampling points that trend towards the bottom right (low $C_T$, high DIN or SRP) thus indicate the on-shelf generation of relatively C-rich OM, while trajectories that tend towards the top left (high $C_T$, low DIN or SRP) indicated more N-rich (and/or P-rich) OM generation, relative to an assumed ‘typical’ C:N:P of 117:16:1 (Anderson and Sarmiento, 1994). The high resolution samples from during the 2015 spring bloom therefore suggest that NCP may have begun with N- and P-rich OM formation, before switching to more C-rich production once surface layer DIN concentrations fell close to zero (Fig. 5b). As mentioned above, the overall change from spring to summer 2015 was similar to the C:N:P uptake stoichiometry observed for the same period in 2014. However, from early spring to summer 2015 we saw the opposite pattern than during the similar period in 2014: the OM became more C-rich. This change to seawater $C_T$ loss with no nutrient loss (as opposed to loss of both to OM following a constant stoichiometry) coincided approximately with CCS surface DIN concentrations approaching zero. We suggest that the $C_T$ loss was therefore a result of continued NCP and OM formation, but that this was using regenerated DIN within the surface layer as its N source rather than drawing from the inorganic pool, which was by then locked away by stratification in the deep layer, as has been observed in the North Sea (Bozec et al., 2006). Although this process should therefore have generated an increasingly C-rich OM pool, presumably associated with more rapid N than C remineralisation from the OM pool, it was not sufficient to return to the original winter 2014 C:N:P stoichiometry. These predicted changes in the OM composition are broadly in agreement with the pattern observed by direct OM stoichiometric measurements at CCS (Davis et al., 2017, this issue), although our approach is not able to distinguish between the dissolved and particulate OM pools.

3.2. Celtic Sea transect

3.2.1. Stratification across the Celtic Sea

In addition to the CCS site, we sampled a transect across the Celtic Sea during each UK-SSB cruise. Analysis of these measurements adds a third dimension (i.e. distance from the open ocean) to our so far two-dimensional analysis (i.e. depth and time), highlighting some additional caveats, sources of uncertainty and broader context. Hartman et al. (2017, this
issue) describe surface layer UK-SSB measurements from across the entire northwest European continental shelf, thus providing further context.

The pattern of winter mixing followed by surface layer uptake of $C_T$, DIN and SRP and their increase in the deep layer was consistent across the entire Celtic Sea transect (Fig. 7). However, the deep layer experienced greater increases in these variables (and others such as $\Delta pCO_2$) further away from the open ocean, although this was reduced for DIN by denitrification/anammox in the seafloor sediments (Hydes et al., 2004; Kitidis et al., 2017). The depth of the surface layer (as defined by the nutricline, 40-50 m) remained relatively constant across the transect while the seafloor became shallower towards the land, so the greater deep layer concentrations could simply indicate that the same amount of NCP had been remineralised, but spread over a narrower deep layer, consistent with the limiting factor for total remineralisation within the deep layer being some function of the OM supply. In addition, the increasing deep layer $C_T$ and nutrients away from the open ocean could reflect the fact that deep waters further from the open ocean had experienced a longer transit time across the shelf within the estuarine-type circulation (Ruiz-Castillo et al., 2017, this issue), and thus greater accumulation of sinking organic matter from the surface layer, in a manner analogous to how nutrient concentrations tend to increase along isopycnals away from the surface in the open ocean (Takahashi et al., 1985).

The increasing strength of biogeochemical stratification moving away from the open ocean has important implications for interpreting our results from CCS, which lay towards the open ocean end of the Celtic Sea transect. For example, the surface layer air-sea $CO_2$ disequilibrium during the stratified period increased away from the open ocean. We would therefore expect the Celtic Sea as a whole to exhibit a greater net air-sea $CO_2$ flux than that calculated for the CCS site, which was relatively close to the open ocean end of the transect. This is consistent with our CCS air-sea $CO_2$ flux estimate falling towards the lower end of previously published results (Section 3.1.1). A second example concerns potential advective fluxes within the Celtic Sea, as discussed in the following section.

3.2.2. Biogeochemical relationships with salinity

In the Celtic Sea, $S$ acts as a tracer of the relative influence of open ocean and shelf waters, particularly in the stratified deep layer where no other processes can modify it. Throughout the year, $S$ decreases moving on to the shelf, ultimately as a result of dilution by riverine inputs. During the winter, $S$ was vertically homogeneous at each point across the shelf, while
the isohalines were tilted away from vertical towards the open ocean in the surface during the summer (i.e. increasing $S$ with depth, Fig. 7h). This suggests that there was likely a net transport towards the open ocean in the surface layer, and towards the land at depth at CCS during the summer. During the winter, while $C_T$ and SRP were laterally homogeneous, DIN was correlated with salinity (Table 3) and declined away from the open ocean (Figs. 7d and 8b). There were no relationships between $S$ and any of $C_T$, DIN or SRP in the surface layer during the summer months, when NCP was the dominant control on these biogeochemical variables. However, correlations could be identified in the deep layer throughout the seasonal cycle, which were variable through time, but fairly consistent with each other (Fig. 8, Table 3). These relationships with $S$ indicated that $C_T$ and SRP were enriched on the shelf relative to the open ocean during the summer months, and returned close to open-ocean values during the winter, consistent with a seasonal cycle of remineralisation and vertical mixing. Although the DIN-$S$ relationship had a similar pattern to those for $C_T$ and SRP, there was a significant offset from the values predicted from the SRP relationship and an assumed N:P stoichiometry of 16:1 (Fig. 8b). This offset was positive, with a mean value of 7.5 ± 2.1 mmol m$^{-3}$ per unit practical salinity, indicating that the shelf waters were virtually always depleted in DIN relative to the open ocean, likely as the result of denitrification and/or anammox in the shelf sediments (Hydes et al., 2004; Kitidis et al., 2017).

The preceding interpretation of the changing $C_T$ and nutrient vertical profiles at CCS mostly ignored the possibility of advective influences. However, given the changing deep layer $C_T$ and nutrient concentrations in the stratified period, advection of water from another part of the shelf could generate an apparent increase or decrease in remineralisation at CCS. As well as directly changing the inorganic inventories, this could also supply OM to CCS that had originally formed elsewhere on the shelf, particularly given our evidence that a significant fraction of NCP (>50%) remained within the OM pool, as opposed to the deep layer remineralised pool, over seasonal timescales (Fig. 5). Significant advection of a cross-shelf OM gradient would hence introduce uncertainties into our one-dimensional estimates of the fraction of NCP that had been remineralised. Indeed, production and advection of a long-lived OM pool could provide a source for the possible excess of remineralisation over NCP that we observed from winter 2014 to winter 2015 at CCS. It is also important to note that we cannot determine how CCS could have been affected by any advective fluxes of material perpendicular to the Celtic Sea transect that we sampled, which could have a different relationship with $S$ than observed along the UK-SSB transect. However, in general we do not...
expect this to have significantly affected our results, because previous studies have found that water exchange between the Celtic Sea and adjacent open ocean is relatively slow (Holt et al., 2009; Huthnance et al., 2009), as is net lateral circulation on the shelf itself (Pingree and Le Cann, 1989), with water taking about 2 years to reach the central Celtic Sea after entering the shelf from the open ocean (Hydes et al., 2004).

The Celtic Sea transect dataset also indicates that the increase in $C_T$ observed at CCS from winter 2014 to winter 2015, and the simultaneous declines in DIN and SRP, were not a result of spatial variability. As these biogeochemical variables appeared to diverge from their open ocean values moving into the shelf sea (Fig. 8), apparent changes at a static site such as CCS could be caused by advective reorganisation of the water on the shelf, with a greater or lesser open ocean influence at CCS in different years. If this were the case, then the apparent nutrient loss that we observed could simply be an artefact of the spatial variability, rather than a temporal biogeochemical trend. However, it appears that the changes in $C_T$, DIN and SRP that we observed at CCS over this time period were consistent throughout most of the range observed across the Celtic Sea transect (Fig. 9). This therefore supports our earlier interpretation that the seasonal biogeochemical changes observed at CCS were representative of processes operating across the wider Celtic Sea.

The shelf $A_T$ was relatively homogeneous both spatially and temporally throughout the UK-SSB sampling campaign (Fig. 7i; Supp. Fig. S8), with a weakly positive relationship with salinity. At CCS, we observed no low $A_T$ excursions from the $A_T$-salinity relationship that could be evidence for pelagic calcification, for example by coccolithophores (Suykens et al., 2010). There was therefore no need to include a correction for this effect in calculating the water column inventories of $C_T$ at CCS through the UK-SSB sampling period.

3.2.3. Nutrient-conserving mechanism

Our estimated net oceanic CO$_2$ uptake of >1 mol-C m$^{-2}$ yr$^{-1}$ within the Celtic Sea suggests that an active CSP operates within this system. Given the preceding presented data and analysis, we suggest two mechanisms through which such a CSP could operate in a nutrient-conserving manner, i.e. without a significant external nutrient-rich source from rivers, and without the shelf becoming depleted in these nutrients over time, as follows.

Firstly, we saw an increase in vertically-integrated $C_T$ at CCS from winter 2014 to winter 2015, along with a simultaneous decline in DIN and SRP (Fig. 5a). The initial winter 2014
values of $C_T$, DIN and SRP were relatively close to the base of the thermocline in the adjacent open ocean, suggesting that there may recently have been relatively strong water exchange across the shelf break, which did not occur again during our sampling period. If the shelf sea were to follow a cycle of increasing $C_T$ while DIN and SRP declined between these irregular ‘flushing’ events, then the exchange of shelf waters with the open ocean would constitute a net export of $C_T$ while DIN and SRP were replenished.

Secondly, as recently been identified in a similar environment in the nearby North Sea (Johnson et al., 2013), it appears that a substantial fraction (>50%) of the OM produced during the spring bloom each year was only slowly remineralised over many months (Fig. 5). Indeed, evidence that deep layer advective transport could have accounted for 70-80% of the DIN increase at CCS from winter to summer each year (Ruiz-Castillo et al., 2017, this issue) suggests that our calculated fraction of OM that was not remineralised is a minimum estimate, so the long-lived OM pool could be greater still. Moreover, the non-steady state situation that we observed over the 1.5 sampled years further suggests that some fraction of this (semi-)refractory OM pool was potentially not remineralised within the same year and hence may play a role in redistributing C and nutrients within the Celtic Sea itself (Fig. 7). We therefore speculate that this long-lived OM pool could also function to export C from the shelf sea into the open ocean, for example during exchange with off-shelf waters with a lower OM concentration.

Future synthesis of the inorganic measurements presented here along with OM concentration and stoichiometry datasets collected during the UK-SSB research programme in the context of a system which likely operates on a multi-annual timescale will be required to assess the capacity of these possible mechanisms to sustain a nutrient-conserving CSP in temperate, seasonally stratified continental shelf seas like the Celtic Sea.
4. Conclusions

The Celtic Sea experiences seasonal physical and biogeochemical stratification with low $C_T$ and nutrient concentrations driving a net air to sea CO$_2$ flux, while remineralisation increases these concentrations in the deep layer. Repeated observations at the CCS site suggest that the inventories of DIN and SRP declined by about 10% from winter 2014 to winter 2015, while $C_T$ increased. Transect data indicate that these temporal changes observed at CCS were consistent across much of the Celtic Sea, rather than being an artefact of advection and spatial variability. Lower availability of nutrients for NCP in 2015 was compensated by a deeper nutricline, so there was no reduction in either the total amount of NCP or air-sea CO$_2$ exchange during the subsequent spring bloom. Analysis of stoichiometry during nutrient uptake for NCP and remineralisation indicated the production of C-rich OM. However, some fraction of the OM generated on the shelf appears to not have been remineralised within the year it was created, suggested that a long-lived stock of OM could accumulate in the shelf waters. This, and the overall changes in inorganic inventories that we observed, could provide alternative mechanisms through which a nutrient-conserving CSP could operate, both in the Celtic Sea and in other seasonally stratified, temperate continental shelf seas.
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7. Tables

Table 1. Summary of the time intervals used in the CCS site analysis.

| Name of time interval | Year | Interval code | Sampling date range (dd/mm/yyyy) | Cruise(s) |
|-----------------------|------|---------------|-----------------------------------|-----------|
| Winter 2014           | W14  | 26/03/2014 – 27/03/2014 | DY008                             |
| Late Spring 2014      | lSp14| 19/06/2014 – 22/06/2014 | JC105                             |
| Summer 2014           | Su14 | 05/08/2014 – 23/08/2014 | DY026A, DY026B                    |
| Autumn 2014           | A14  | 10/11/2014 – 29/11/2014 | DY018                             |
| Winter 2015           | W14  | 22/03/2015     | DY021                             |
| In-bloom 1 2015       | IB1  | 03/04/2015 – 06/04/2015 | DY029                             |
| In-bloom 2 2015       | IB2  | 11/04/2015 – 12/04/2015 | DY029                             |
| In-bloom 3 2015       | IB3  | 15/04/2015     | DY029                             |
| In-bloom 4 2015       | IB4  | 16/04/2015     | DY029                             |
| In-bloom 5 2015       | IB5  | 20/04/2015     | DY029                             |
| In-bloom 6 2015       | IB6  | 21/04/2015     | DY029                             |
| In-bloom 7 2015       | IB7  | 25/04/2015 – 26/04/2015 | DY029                             |
| In-bloom 8 2015       | IB8  | 28/04/2015     | DY029                             |
| Early Spring 2015     | eSp15| 21/05/2015 – 23/05/2015 | DY030                             |
| Summer 2015           | Su15 | 24/07/2015 – 23/08/2015 | DY033, DY034                      |
Table 2. Details of the empirical algorithm used to predict surface $p\text{CO}_2^{sw}$ at CCS.

| Day of year (DoY) range | SST range / °C        | $p\text{CO}_2^{sw}$ / µatm |
|-------------------------|------------------------|-------------------------------|
| 80 < DoY < 232          | SST < 10.5             | $1.547 \times 10^{-3} \exp(SST) + 372.7$ |
| 80 < DoY < 232          | 10.5 ≤ SST ≤ 11.0      | $324.5 + 6.596 \times 10^{24} \exp[-5.000(SST)]$ |
| 80 < DoY < 232          | 11.0 < SST             | $-0.557(SST)^2 + 23.57(SST) + 141.1$ |
| DoY ≤ 80 or 233 ≤ DoY   | All                    | $-0.2479(SST)^3 + 10.846(SST)^2 - 151.42(SST) + 1050.3$ |
Table 3. Correlation statistics between biogeochemical variables and salinity. Columns headed dX/dS contain gradient ± its standard error in μmol kg\(^{-1}\) (Figure 8); N is the number of observations used in each regression.

| Cruise  | Interval code | C\(_T\)   | DIN       | SRP       |
|---------|---------------|-----------|-----------|-----------|
|         |               | dC\(_T\)/dS | r\(^2\)  | p  | N  | dDIN/dS | r\(^2\) | p  | N  | dSRP/dS | r\(^2\) | p  | N  |
| DY008   | W14           | -10.6 ± 8.9 | 0.11 | 0.26 | 13 | 7.6 ± 0.6 | 0.86 | 0.00 | 25 | -0.23 ± 0.09 | 0.21 | 0.02 | 25 |
| JC105   | lSp14         | -109.6 ± 10.0 | 0.86 | 0.00 | 21 | 1.0 ± 0.8 | 0.07 | 0.24 | 21 | -0.46 ± 0.06 | 0.75 | 0.00 | 21 |
| DY026*  | Su14          | -60.6 ± 10.8 | 0.78 | 0.00 | 11 | 4.1 ± 1.2 | 0.36 | 0.00 | 23 | -0.29 ± 0.06 | 0.47 | 0.00 | 26 |
| DY018   | A14           | -99.5 ± 10.6 | 0.49 | 0.00 | 92 | -0.2 ± 1.1 | 0.00 | 0.89 | 183 | -0.46 ± 0.07 | 0.18 | 0.00 | 202 |
| DY021   | W15           | -19.6 ± 9.1 | 0.28 | 0.05 | 14 | 5.9 ± 0.7 | 0.59 | 0.00 | 47 | 0.01 ± 0.06 | 0.00 | 0.87 | 47 |
| DY029   | IB1–IB8       | -39.5 ± 7.3 | 0.54 | 0.00 | 27 | 7.4 ± 0.4 | 0.65 | 0.00 | 192 | 0.11 ± 0.02 | 0.12 | 0.00 | 202 |
| DY030   | eS15          | -26.7 ± 13.4 | 0.50 | 0.12 | 6  | 9.5 ± 1.2 | 0.85 | 0.00 | 13 | 0.32 ± 0.10 | 0.48 | 0.01 | 13 |
| DY033   | Su15          | -63.1 ± 8.1 | 0.68 | 0.00 | 31 | 4.2 ± 0.4 | 0.42 | 0.00 | 182 | -0.21 ± 0.02 | 0.30 | 0.00 | 186 |
| DY034   | Su15          | -120.9 ± 24.0 | 0.78 | 0.00 | 9  | 0.4 ± 1.8 | 0.00 | 0.84 | 44 | -0.51 ± 0.12 | 0.31 | 0.00 | 44 |

*DY026 includes all data from both DY026A and DY026B.
| Start | End  | Type | C | N | P |
|-------|------|------|---|---|---|
| W14   | lSp14| NCP  |   |   |   |
| W15   | eSp15| NCP  |   |   |   |
| W15   | Su15 | NCP  |   |   |   |
8. Figures

Figure 1. The UK-SSB sampling locations in the Celtic Sea. The pink circle encloses the data considered to represent the Central Celtic Sea site (i.e. within 12 km of 49.4°N, 8.54°W), while the yellow lines indicate the transect route for Fig. 7 and enclose the data included in those plots; the points labelled A, B and C correspond to the equivalent points on Fig. 7. The 200 m bathymetry contour, traditionally used to define the edge of the shelf, is shown as a black line. Topography data are from the GEBCO_2014 30 arc-second grid (version 20150318, http://www.gebco.net).
Figure 2. Time-series of air-sea CO₂ disequilibrium and flux at CCS. (a) Atmospheric (red) and sea surface (blue) $pCO₂$ at CCS. The black points show the actual measurements of surface $pCO₂^{sw}$, while the blue line shows our algorithm-predicted interpolation of these data (Table 2) that was used to calculate $ΔpCO₂$ and then the air-sea CO₂ flux. (b) The sea to air CO₂ flux. The shaded areas show the 95% confidence intervals.
Figure 3. Hovmöller diagrams of CCS water column (a) temperature, (b) dissolved inorganic carbon, (c) dissolved inorganic nitrogen (i.e. nitrate + nitrate + ammonium), (d) soluble reactive phosphorus, (e) practical salinity, and (f) air-sea $\text{CO}_2$ disequilibrium. Where shown, sampling locations are indicated as black points. The cycle of winter mixing and summer stratification is clear for most variables, while $S$, DIN and SRP show an anomalous increase in the deep layer during late 2014, indicating an influx of open ocean waters.
Figure 4. Changes in the DIN profile at CCS for selected time intervals. In each panel, the earlier profile is shown as a dashed line, while the later one is solid. The blue shaded areas therefore indicate decreases in DIN, while orange shows increases. (a) Primary production in the surface layer drew down DIN, while remineralisation at depth increased the DIN concentration. (b) The onset of vertical mixing eroded the surface-deep DIN gradient, overlaid by a remineralisation-driven increase in the total DIN inventory. (c) The water column returned to a vertically homogeneous state following further vertical mixing.
Figure 5. Water column inventories at CCS throughout the SSB study period. The inventories have been divided by the water depth, so the values shown are the vertical mean at each time point. Downwards/upwards pointing triangles represent 2014/2015 data respectively, and the grey diagonal lines indicate a C:N:P stoichiometry of 117:16:1 (Anderson and Sarmiento, 1994). W = winter; eSp = early spring; lSp = late spring; Su = summer; A = autumn; 14/15 indicates the year. (a) Raw inventories of $C_T$ and $N_T$ measured at CCS, coloured by the sampling date. (b) Here, the $C_T$ inventory is corrected for air-sea gas exchange relative to the first sampling point, while the DINis corrected for denitrification/anammox. The colour shows the surface layer DIN concentration. (c) Relative changes in DIN (corrected for denitrification/anammox) and SRP, coloured by mean salinity at CCS at each time point. Higher salinity values indicate the influence of open ocean waters.
Figure 6. Winter to summer changes in (a) DIN and (b) seawater temperature for 2014 (blue) and 2015 (red). In (a), the dashed lines show the lowest possible ΔDIN, i.e. its value if the entire winter DIN inventory had been converted into OM. All available DIN in the surface layer was converted to OM in both years. This represented a greater change in DIN in 2014 due to the higher initial DIN concentration, but the deeper nutricline in 2015 led to greater NCP overall. Remineralisation increased the deep layer DIN concentration by the same amount in both years, but this represented a smaller absolute DIN addition in 2015 owing to the deeper nutricline.
Figure 7. Transects across the Celtic Sea for selected variables and UK-SSB cruises. In each panel, the area above the horizontal dotted line in the left plot (shelf break) is the same as that to the left of the vertical dotted line in the right plot (on the shelf), and the black points indicate where samples were collected. Points A, B, CCS and C are located on Fig. 1. (a) and (b) show $C_T$ in winter and summer 2014 respectively, while (c) shows $\Delta pCO_2$ in summer 2014. (d), (e) and (f) show the evolution of DIN from spring through summer to autumn 2014. (g) and (h) illustrate changes in salinity between winter and summer 2015, while (i) shows $A_T$ in summer 2014. A full set of transects for every variable and UK-SSB cruise is provided in Supp. Figs. S6–S11.
Figure 8. Time-series of regression slopes between $S$ and (a) $C_T$, (b) DIN, and (c) SRP.

Negative values indicate that the on-shelf deep layer is enriched in the nutrient relative to the open ocean, while positive values indicate on-shelf depletion. Error bars show the standard error in the gradient at each point (Table 3). In (b), the predicted relationship based on dSRP/dS is also shown, considered to represent the component driven by OM remineralisation. The offset between this and the solid line (mean ± SD difference = 7.5 ± 2.1 mmol m$^{-3}$) was likely driven by denitrification/anammox in the Celtic Sea; note that this offset value is reported per unit practical salinity (which is itself dimensionless).
Figure 9. Changes in (a) $C_T$, (b) DIN and (c) SRP, across the entire Celtic Sea transect from winter 2014 (blue) to winter 2015 (red). The × symbols show the mean conditions at CCS each winter. Salinity values increase towards the open ocean. The increase in $C_T$ observed at CCS from winter 2014 to winter 2015, and corresponding declines in DIN and SRP, appear to be consistent across much of the Celtic Sea, rather than being a feature of spatial variability local to CCS.
9. Supplementary information

9.1. Supplementary tables

Table S1. Summary of the UK-SSB research cruises.

| Cruise code | Cruise sampling dates (dd/mm/yy) | Cruise Principle Investigator | $C_T/A_T$ sample collectors | No. of $C_T/A_T$ samples |
|-------------|---------------------------------|-------------------------------|-----------------------------|--------------------------|
| DY008       | 21/03—04/04/14                  | H. A. Ruhl                    | L. Darroch                  | 45                       |
| JC105       | 16/06—22/06/14                  | J. E. Hopkins                 | J. Fox, R. Houlding, P. Nelson | 105                     |
| DY026A      | 04/08—13/08/14                  | R. Sanders                    | R. Sims                     | 60                       |
| DY026B      | 17/08—22/08/14                  | D. Sivyer                     | R. Sims                     | 16                       |
| DY018       | 10/11—01/12/14                  | J. Sharples                   | L. Darroch, J. E. Hopkins   | 301                      |
| DY021       | 03/03—24/03/15                  | E. M. S. Woodward             | N. Hicks                    | 47                       |
| DY029       | 02/04—28/04/15                  | A. Poulton                    | A. Poulton, K. Mayers       | 214                      |
| DY030       | 05/05—23/05/15                  | G. Fones                      | R. Sims                     | 72                       |
| DY033       | 13/07—01/08/15                  | C. M. Moore                   | R. Sims                     | 201                      |
| DY034       | 07/08—31/08/15                  | H. A. Ruhl                    |                             | 52                       |

* Cruise codes beginning with DY were on the RRS Discovery, while JC stands for the RRS James Cook. * Cruise sampling dates indicate the time period for collection of samples used in this study, not necessarily the entire duration of the cruises.
Table S2. Coefficients fitted to Eq. (1) at CCS for each time interval (cf. Table 1 and Supp. Figs. S1–S4) for the variables DIC, DIN, DIP, and practical salinity. Units for \( v_0 \) and \( v_1 \) are the same as for the relevant variable (i.e. mmol m\(^{-3}\) for \( C_T \), DIN and SRP, and dimensionless for practical salinity), while \( z_0 \) and \( z_1 \) are in m.

| Interval code | \( C_T \) | DIN | SRP | Practical salinity |
|---------------|----------|-----|-----|--------------------|
|               | \( v_0 \) | \( v_1 \) | \( z_0 \) | \( z_1 \) | \( v_0 \) | \( v_1 \) | \( z_0 \) | \( z_1 \) | \( v_0 \) | \( v_1 \) | \( z_0 \) | \( z_1 \) |
| W14           | 2199.3   | 0    | 0    | 0                   | 8.22   | 0    | 0    | 0    | 35.310 | 0    | 0    | 0    |
| ISp14         | 2168.9   | 38.70| 40.0 | 9.64               | 4.64   | 4.391| 43.8 | 6.80 |
| Su14          | 2173.2   | 42.87| 36.2 | 8.36               | 4.68   | 4.542| 39.9 | 6.48 |
| A14           | 2190.3   | 27.88| 54.4 | 9.99               | 6.48   | 3.808| 53.3 | 9.24 |
| W15           | 2207.1   | -1.08| 80.0 | 12.38              | 7.34   | 0.056| 28.0 | 0.69 |
| IB1           | 2206.6   | 2.04 | 70.7 | 0.68               | 6.56   | 0.529| 52.9 | 15.83|
| IB2           | 2203.6   | 7.86 | 35.8 | 13.93              | 5.37   | 1.465| 34.3 | 17.85|
| IB3           | 2192.8   | 15.97| 26.8 | 19.54              | 3.21   | 3.549| 19.7 | 26.09|
| IB4           | 2197.8   | 12.38| 36.5 | 15.30              | 4.37   | 2.385| 35.8 | 18.15|
| IB5           | 2195.1   | 13.27| 42.3 | 7.71               | 4.70   | 2.027| 41.7 | 8.91 |
| IB6           | 2195.5   | 11.54| 30.2 | 13.78              | 4.06   | 2.897| 32.4 | 8.04 |
| IB7           | 2191.3   | 17.68| 35.9 | 9.39               | 3.92   | 2.944| 35.2 | 10.53|
| IB8           | 2186.6   | 21.58| 37.7 | 3.21               | 4.03   | 3.247| 37.2 | 4.89 |
| eSp15         | 2186.4   | 21.92| 35.1 | 4.04               | 4.17   | 3.634| 35.6 | 5.34 |
| Su15          | 2180.1   | 37.27| 44.1 | 12.17              | 4.20   | 4.122| 48.6 | 7.78 |

| Interval code | \( v_0 \) | \( v_1 \) | \( z_0 \) | \( z_1 \) | \( v_0 \) | \( v_1 \) | \( z_0 \) | \( z_1 \) |
|---------------|----------|----------|----------|----------|----------|----------|----------|----------|
| W14           | 0.59     | 0        | 0        | 0        | 35.310   | 0        | 0        | 0        |
| ISp14         | 0.37     | 0.303    | 42.1     | 10.80    | 35.324   | -0.0101  | 18.8     | 2.33     |
| Su14          | 0.36     | 0.302    | 38.8     | 7.01     | 35.395   | 0.0429   | 21.8     | 17.17    |
| A14           | 0.45     | 0.248    | 54.2     | 9.41     | 35.437   | 0.0444   | 58.1     | 2.45     |
| W15           | 0.49     | 0.040    | 0.4      | 29.42    | 35.356   | 0.0096   | 15.0     | 3.33     |
| IB1           | 0.50     | 0.025    | 51.6     | 12.65    | 35.343   | 0.0222   | 54.8     | 14.72    |
| IB2           | 0.43     | 0.089    | 36.5     | 14.87    | 35.338   | -0.0251  | 23.7     | 52.69    |
| IB3           | 0.27     | 0.250    | 17.4     | 30.97    | 35.326   | -0.0097  | 43.7     | 18.90    |
| IB4           | 0.34     | 0.146    | 37.0     | 20.29    | 35.321   | -0.0106  | 31.8     | 6.99     |
| IB5           | 0.38     | 0.143    | 42.1     | 8.16     | 35.326   | -0.0174  | 42.4     | 6.72     |
|     | 0.34 | 0.189 | 33.1 | 8.33 | 35.322 | -0.0168 | 34.9 | 8.36 |
|-----|------|-------|------|------|--------|----------|------|------|
| Ib6 | 0.33 | 0.184 | 36.5 | 11.04| 35.312 | -0.0058 | 42.3 | 0.90 |
| Ib7 | 0.34 | 36.363 | 0.16 | 5.02 | 35.311 | 0.0041  | 35.3 | 2.01 |
| Ib8 | 0.35 | 35.486 | 0.2  | 6.03 | 35.322 | 0.0218  | 35.9 | 7.20 |
| eSp15 | 0.34 | 48.082 | 0.27 | 9.08 | 35.398 | -0.0266 | 35.1 | 0.03 |

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9.2. Supplementary figures

Figure S1. Dissolved inorganic carbon ($C_T$) profiles at each time point at CCS (Table 1). The grey circles show the entire CCS dataset, while the blue circles highlight the data at each time point. The blue line shows the final vertical profile used for inventory calculations (Supp. Table S2).
Figure S2. Dissolved inorganic nitrogen (DIN, nitrate + nitrite + ammonium) profiles at each time point at CCS (Table 1). The grey circles show the entire CCS dataset, while the blue circles show the data at each time point. The blue line shows the final vertical profile used for inventory calculations (Supp. Table S2).
Figure S3. Soluble reactive phosphorus (SRP) profiles at each time point at CCS (Table 1). The grey circles show the entire CCS dataset, while the blue circles show the data at each time point. The blue line shows the final vertical profile used for inventory calculations (Supp. Table S2).
Figure S4. Practical salinity profiles at each time point at CCS (Table 1). The grey circles show the entire CCS dataset, while the blue circles show the data at each time point. The blue line shows the final vertical profile used for inventory calculations (Supp. Table S2).
Figure S5. Seawater pH on the Free scale at each time point at CCS (Table 1), calculated from $A_T$ and $C_T$ measurements (van Heuven et al., 2011). The grey circles show the entire CCS dataset, while the blue circles show the data at each time point.
Figure S6. Transects of dissolved inorganic carbon ($C_I$) measured across the Celtic Sea for all of the SSB cruises. In each panel, the area above the horizontal dotted line in the left plot is the same as that to the left of the vertical dotted line in the right plot. Black points indicate the sample locations. The geographical locations of points A, B, CCS and C are shown by Fig. 1.
Figure S7. Transects of $\Delta pCO_2$, calculated from $C_T$ and $A_T$ measurements, across the Celtic Sea for each SSB cruise. The layout is as described for Fig. 7.
Figure S8. Transects of total alkalinity ($A_T$) across the Celtic Sea for all of the UK-SSB cruises. In each panel, the area above the horizontal dotted line in the left plot is the same as that to the left of the vertical dotted line in the right plot. Black circles indicate sample locations.
Figure S9. Transects of dissolved inorganic nitrogen (DIN, nitrate + nitrite + ammonium) across the Celtic Sea for all of the UK-SSB cruises. In each panel, the area above the horizontal dotted line in the left plot is the same as that to the left of the vertical dotted line in the right plot. Black circles indicate sample locations.
Figure S10. Transects of soluble reactive phosphorus (SRP) across the Celtic Sea for all of the UK-SSB cruises. In each panel, the area above the horizontal dotted line in the left plot is the same as that to the left of the vertical dotted line in the right plot. Black circles indicate sample locations.
Figure S11. Transects of practical salinity across the Celtic Sea for all of the UK-SSB cruises.

In each panel, the area above the horizontal dotted line in the left plot is the same as that to the left of the vertical dotted line in the right plot. Black circles indicate sample locations.
Figure S12. Relationship between dissolved inorganic carbon ($C_T$) and practical salinity for each UK-SSB cruise. The points used to generate each regression are shown in blue (Section 2.8), and the slope is indicated towards the top right of each panel (see also Fig. 7 and Table 3).
Figure S13. Relationship between dissolved inorganic nitrogen (DIN, nitrate + nitrite + ammonium) and practical salinity for each UK-SSB cruise. The points used to generate each regression are shown in blue (Section 2.8), and the slope is indicated towards the top right of each panel (see also Fig. 7 and Table 3).
Figure S14. Relationship between soluble reactive phosphorus (SRP) and practical salinity for each UK-SSB cruise. The points used to generate each regression are shown in blue (Section 2.8), and the slope is indicated towards the top right of each panel (see also Fig. 7 and Table 3).