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Resilience of the Copepod Oithona similis to Climatic Variability: Egg Production, Mortality, and Vertical Habitat Partitioning

Louise Elisabeth Cornwell1*, Elaine S. Fileman1, John T. Bruun2,3, Andrew Garwood Hirst4,5, Glen Adam Tarran1, Helen S. Findlay1, Ceri Lewis6, Timothy James Smyth1, A. J. McEvoy1 and A. Atkinson1

1 Plymouth Marine Laboratory, Plymouth, United Kingdom. 2 College of Engineering, Mathematics and Physical Sciences, University of Exeter, Exeter, United Kingdom. 3 College of Life and Environmental Sciences, University of Exeter, Exeter, United Kingdom. 4 School of Environmental Sciences, University of Liverpool, Liverpool, United Kingdom. 5 Centre for Ocean Life, National Institute for Aquatic Resources, Technical University of Denmark, Lyngby, Denmark. 6 Hatherly Laboratories, College of Life and Environmental Sciences, University of Exeter, Exeter, United Kingdom

There has been an overall decline in copepod populations across the North Atlantic over the past few decades. Reasons for these declines are unclear, and several major species, including the cyclopoid copepod Oithona similis, have maintained stable populations at station L4 in the western English Channel. To identify the factors contributing to this stability, we conducted a 1-year intensive study of O. similis over 2017–2018, a period of high climatic variability. For context, dominant frequency state analysis was applied to the 30-year L4 time series to derive the baseline dynamics of the Oithona spp. population. The Oithona spp. baseline demonstrated stable densities and a bimodal annual cycle. These dynamics, as well as those of reproductive output and phaenological timings, were upheld in 2017–2018, indicating resilience to climatic variability. During 2017–2018, all life stages of O. similis were relatively scarce in the top 2 m of the water column, despite the presence of abundant food. Naupliar stages occurred predominantly around 10 m depth, with subsequent life stages progressively deeper. We suggest this vertical structuring may represent different trade-offs between feeding and mortality risk between ontogenetic stages. To determine the traits that contribute to population stability, we compare O. similis with the large, biomass-dominant copepod, Calanus helgolandicus. Despite having contrasting functional traits, both species have exhibited strong population stability over the time series. Our results provide evidence that mortality plays a major role in maintaining population dynamics.

Keywords: Oithona similis, Western Channel Observatory, vertical distribution, population stability, dominant frequency state analysis

INTRODUCTION

Copepod populations have undergone strong declines across the North Atlantic over the past few decades (Edwards et al., 2016; O’Brien et al., 2017). Although it is well-established that the global oceans are changing at an unprecedented rate (IPCC, 2019), the drivers behind the observed copepod declines are uncertain, and some important species have maintained stable populations.
Small pelagic copepods, such as the cyclopoid *Oithona similis*, are some of the most numerous metazoans on Earth (Turner, 2004), and provide an important link between the microbial food web and higher trophic levels (Nielsen and Sabatini, 1996; Turner, 2004). Furthermore, *O. similis* populations are commonly reproductively active throughout the year (Digby, 1954; McLaren, 1969; Sabatini and Kiorboe, 1994; Franzs and Gonzalez, 1995; Ashjian et al., 2003; Lischka and Hagen, 2005; Dvoretski, 2007; Dvoretsky and Dvoretsky, 2009), making this species a particularly significant ecosystem component in winter, when larger copepod species are scarce (Nielsen and Sabatini, 1996). Therefore, understanding which factors control the population dynamics of this ubiquitous and ecologically significant species is important.

The Western Channel Observatory (WCO) station L4 is a highly seasonal, temperate shelf site. The physical and biological environment at L4 is well-studied, with numerous publications on the plankton community (Elmoire et al., 2010; Highfield et al., 2010; Widdicombe et al., 2010; Atkinson et al., 2015; Tarran and Bruun, 2015; White et al., 2015). The 30-year L4 time series is thus an invaluable resource through which to investigate the impact of environmental variation on zooplankton populations. As seen across the wider North Atlantic (Edwards et al., 2016; O'Brien et al., 2017), there has been a decrease in overall copepod densities at L4 over the past 30 years (Edwards et al., 2020). However, densities and phenological timings of the *O. similis* population at this site have remained relatively stable (Castellani et al., 2016; Cornwell et al., 2018), as they have in the larger calanoid copepod, *Calanus helgolandicus* (Maud et al., 2015; Edwards et al., 2020).

To identify the potential factors behind the *O. similis* population stability at L4, we conducted a 1-year intensive study over 2017–2018. The study period included exceptionally low water temperatures in spring, followed by a rapid rate of warming prior to summer. For context, dominant frequency state analysis (DFSA) was used to derive the baseline *Oithona* spp. dynamic for the 30-year L4 time series. To further explore the population seasonality during 2017–2018, depth-resolved sampling was used to determine the vertical distribution of all *O. similis* life stages, alongside that of potential prey. Our final study, before summarising the contextual measurements made as part of the L4 time series.

**Long-Term Sampling at Station L4**

To provide context for the 2017–2018 study, L4 zooplankton time series data were used to derive the baseline dynamics of the *Oithona* spp. population. L4 time series data were also used to construct the baseline seasonal dynamic for the physical environment. The duration, sampling methodology, and sources of all datasets analysed in our study are provided in Table 1.

Samples for *Oithona* spp. density were collected from two successive 0–50 m vertical hauls with a 200 µm mesh, 57 cm diameter WP2 net (UNESCO, 1968), and fixed in 4% buffered formalin. The category *Oithona* spp. comprises adults of both sexes and late stage copepodites, although the latter are not quantitatively sampled. Data derived from this long-term sampling was used to determine the day of the year at which the *Oithona* spp. population reached 25, 50 and 75% of the total annual cumulative density, indicating the start, middle and end of season, respectively (Mackas et al., 2012). The duration index of the population was derived from the number of days between the 25th and 75th cumulative percentiles. Samples for *O. similis* egg production rate (EPR) were collected from 0–50 m vertical hauls in 2003, and 2011 to 2016. In 2003, samples were collected using a ring net (50 µm mesh, 50 cm diameter) and fixed in 4% buffered formalin. In November 2011 to March 2014 samples were collected using a ring net (63 µm mesh, 57 cm diameter) and fixed in 4% buffered formalin. In June 2015 to December 2016 samples were collected using a ring net (63 µm mesh, 57 cm diameter) and fixed in 2% acid Lugol’s solution.

Water column temperature was determined using a Sea-Bird Scientific SBE 19plus SeaCAT Profiler CTD. Another physical parameter analysed in our study was the surface net heat flux (NHF, W m⁻²) between the atmosphere and the ocean, which incorporates air-sea temperature difference, irradiance, wind speed, and stratification (Smyth et al., 2014). Surface NHF was determined using the following methodology of Smyth et al. (2014). Four processes control air-sea heat flux: shortwave radiation from the sun (QSW), outgoing longwave radiation from the sea surface (QLW), sensible heat transfer resulting from air-sea temperature differences (QSH), and latent heat transfer via evaporation of seawater (QLH). The Woods Hole Oceanographic Institution air-sea exchange MATLAB® tools (Fairall et al., 2003) were used to determine QSW, QLW, QSH and QLH (Pawlowicz et al., 2001), in units of W m⁻². Meteorological parameters were obtained from the European Centre for Medium Range Weather Forecasting (ECMWF) ERA-40 and Operational analyses, extracted for the grid point 50°N, 4°W. These parameters were the following: air temperature (Tₐ, °C), dew point (T_d, °C), wind speed at 10 m (U₁₀, ms⁻¹), cloud fraction (CF, 0: clear; 1: overcast) and atmospheric pressure (P, mb). Sea surface temperature (SST) (T_s, °C), combined with the ECMWF data, was used to run the heat flux model for the period 1988–2018. QSW was calculated as a function of date and position with correction for CF (Reed, 1977); QLW as a function of Tₐ, T_s, T_d and CF using the Berliand bulk formula (Fung et al., 1984). QSH and QLH were calculated as a function of Tₐ, Tₛ, T_d, Tₐ, Tₛ, T_d, Tₐ, Tₛ, T_d, Tₐ, Tₛ, T_d, Tₐ, Tₛ, T_d, Tₐ, Tₛ, T_d, Tₐ, Tₛ, T_d.
TABLE 1 | The duration, sampling methodology, and sources of all datasets analysed in the present study.

<table>
<thead>
<tr>
<th>Time series</th>
<th>Years analysed</th>
<th>Sampling method</th>
<th>Data source</th>
</tr>
</thead>
<tbody>
<tr>
<td>Oithona spp. Density</td>
<td>March 1988–December 2017</td>
<td>Two successive 0–50 m vertical hauls, 200 µm mesh, 57 cm diameter, WP2 net (UNESCO, 1968), fixed in 4% buffered formalin</td>
<td>Western Channel Observatory</td>
</tr>
<tr>
<td>Oithona similis egg production rate</td>
<td>January 2003–December 2003</td>
<td>Ring net (50 µm mesh, 50 cm diameter), 0–50 m, fixed in 4% buffered formalin</td>
<td>Cornwell et al., 2018</td>
</tr>
<tr>
<td></td>
<td>November 2011–March 2014</td>
<td>Ring net (63 µm mesh, 57 cm diameter), 0–50 m, fixed in 4% buffered formalin</td>
<td></td>
</tr>
<tr>
<td></td>
<td>June 2015–December 2016</td>
<td>Ring net (63 µm mesh, 50 cm diameter), 0–50 m, fixed in 2% acid Lugol’s solution</td>
<td></td>
</tr>
<tr>
<td>Depth integrated temperature</td>
<td>January 2002–December 2017</td>
<td>Sea-Bird Scientific SBE 19plus SeaCAT Profiler CTD</td>
<td>Western Channel Observatory</td>
</tr>
<tr>
<td>Surface net heat flux</td>
<td>March 1988–December 2017</td>
<td>Meteorological parameters obtained from the European Centre for Medium Range Weather Forecasting (ECMWF) operational and ERA-40 datasets</td>
<td>Centre for Environmental Data Analysis</td>
</tr>
</tbody>
</table>

Access to the most recently updated versions of the Western Channel Observatory (WCO) time series data is available from Plymouth Marine Laboratory upon request (http://www.westernchannelobservatory.org.uk/).

CF, P, and U₁₀. The sum of all four components results in surface NHF, with the sign convention of positive NHF being heat flux into the water column.

**Plankton Sampling During 2017–2018**

Sampling was conducted at mid-morning on a near-weekly basis from 18th September 2017 to 15th October 2018. To determine the vertical distribution of *O. similis* life stages, 10 L Niskin bottles were attached to a CTD rosette, and fired at depths of 2, 10, 25 and 50 m. At each depth, 10 L of water was collected and transferred from the Niskin bottle into a carboy via silicone tubing attached to the Niskin bottle valve. Triplicate seawater samples for picoplankton and small nanoplankton (≤12 µm) were collected at each depth from the same, or a subsequent, CTD cast. These samples were transferred from the Niskin bottle via silicone tubing into 250 mL polycarbonate bottles, and stored in a cool box. All samples were transported to the laboratory within 3 h of collection, and analysed immediately.

**Plankton Analysis**

In the laboratory, the contents of each carboy were gently decanted into 5 L beakers to ensure the sample was well-mixed. Subsequently, 250 mL sub-samples were taken for each depth and fixed in 2% acid Lugol’s solution for analysis of the composition and biomass of large nanoplankton (>12 µm) and microplankton. The remainder of the 10 L sample was concentrated down to 50–200 mL by reverse filtration using a 20 µm mesh, and fixed in 2% acid Lugol’s solution for analysis of *O. similis* density. These concentrated samples were settled for ~24 h, after which the top layer was gently removed via syringe, leaving the bottom 50 mL to be settled for a further ~15 h into a 3 mL Hydrobios® counting chamber (Utermöhl, 1958). Following this, adult males and females, copepodites, nauplii and eggs of *O. similis* were enumerated under an Olympus IMT-2 inverted microscope at 40x magnification. Egg sacs of *O. similis* were easily identified, being relatively uniform and elongated compared to the irregular clusters of eggs produced by other sac-spawning species at L4. Both detached and attached egg sacs were counted. Egg densities were calculated as the product of the number of egg sacs and the mean number of eggs per sac. Nauplii were identified following Gibbons and Ogilvie (1933) and Lovegrove (1956). Early life stages were difficult to distinguish from those of the congener *O. nana*. However, as *O. nana* comprises just ~7% of total *Oithona* spp. density at L4 (Cornwell et al., 2018), all early life stages of the genus were assumed to belong to *O. similis*. Nauplii and copepodites were counted separately but were not staged. For each sampling event, the median count of total *O. similis* across all life stages and depths was 1101 individuals. The weighted mean depth (WMD) for each life stage sampled from the Niskin bottles was calculated as:

\[
WMD = \frac{\sum (D_i \times \text{depth}_i)}{\sum D_i}
\]

Where \(D_i\) is density (ind m\(^{-3}\)) of a specific life stage at depth \(i\).

To assess food availability in the water column, the 250 mL sub-samples taken to measure the composition and biomass of the large nanoplankton and microplankton were settled for ~60 h, after which the top 225 mL was gently removed via syringe. The remaining 25 mL was then analysed using FlowCam VS-IVc (Fluid Imaging Technologies, Inc.). The FlowCam was fitted with a 10x objective and 100 µm flow cell. Images were collected using AutoImage Mode at 20 frames sec\(^{-1}\). For each sampling event, the median cell count across all depths was 28382 cells. To avoid clogging, samples were passed through a 100 µm mesh before transfer into the sample funnel. Thus, our calculation of prey biomass excludes cells greater than 100 µm. Cells were classified manually using VisualSpreadsheet® (V 4.1.95) software, and categorised as pennate diatoms, centric
diatoms, large flagellates, round-shaped dinoflagellates, ellipsoid-shaped dinoflagellates, and ciliates. Chain forming diatoms were counted per chain. FlowCam data for cell volume, derived from area based diameter, were used to calculate carbon biomass for each category from the conversion equations of Menden-Deuer and Lessard (2000). As samples for FlowCam analysis were fixed with 2% acid Lugol’s solution, some cells may have shrunk or become swollen. However, biomass estimates for fixed species assemblages based on live or fixed cell volumes have been shown not to differ significantly (Menden-Deuer et al., 2001), thus no corrections were applied.

Samples for picoplankton and small nanoplanckton (≤12 µm) were analysed live on an Accuri C6 flow cytometer (for details see Tarran and Bruun, 2015). Autotrophic picoeukaryotes (<3 µm) and nanoeukaryotes (2–12 µm), coccolithophores (5–8 µm), cryptophytes (4–10 µm), ‘Phaeocystis-like’ cells (5–6 µm), and ‘dinoflagellate-like’ cells (<20 µm) were grouped as ‘autotrophic flagellates,’ with heterotrophic nanoeukaryotes (2–12 µm) grouped separately. The group ‘heterotrophic bacteria’ consisted of high and low nucleic acid-containing bacteria. The final group consisted of autotrophic cyanobacteria Synechococcus. Cell volumes were generated by gravity filtering seawater samples through a series of polycarbonate membrane filters from 18 µm down to 0.2 µm. The filtrates were then analysed on the flow cytometer, and compared against a sample of unfiltered seawater. For each group, cell counts were used to calculate the percentage of cells remaining in the sample after filtration relative to the unfiltered sample. These percentages were plotted against filter pore size to create a series of sigmoid curves. Median cell size per group was estimated from the filter pore size at which 50% of cells were retained by the filter. From median cell size, cell volumes were calculated assuming the volume of a sphere as an approximation of cell shape. A carbon conversion factor of 0.22 pgC µm⁻³ (Heywood et al., 2006), was applied to autotrophic flagellates (excluding coccolithophores and ‘dinoflagellate-like’ cells), heterotrophic nanoeukaryotes and Synechococcus. Coccolithophore cell density was converted to carbon biomass using a carbon conversion factor of 0.285 pgC µm⁻³ (Tarran et al., 2006) to account for the higher carbon associated with coccoliths. Carbon biomass of ‘dinoflagellate-like’ cells was calculated from the carbon to volume relationship of Menden-Deuer and Lessard (2000). The carbon biomass of heterotrophic bacteria was calculated using a carbon conversion factor of 20 fgC cell⁻¹ (Lee and Fuhrman, 1987).

Dominant Frequency State Analysis

The physical and biological systems at L4 are strongly influenced by the annual cyclic motion of the Earth and by decadal processes (Smyth et al., 2014). DFSA is a highly accurate and efficient estimation signal analysis approach designed to resolve these cyclic Earth-system processes (Smyth et al., 2014; Barnes et al., 2015; Tarran and Bruun, 2015; Bruun et al., 2017). Here, DFSA was used to construct a predictive dynamic model of the seasonal and decadal properties of how the Oithona spp. population responds to the L4 environment. The L4 zooplankton time series provides a robust record of the dominant cyclic trend of Oithona spp. density. From the time series data (1988–2017), estimates of the primary Oithona spp. cyclic processes, and the baseline Oithona spp. dynamic, were derived using DFSA. The few data gaps were pre-process filled using a 6-week running median. An autoregressive noise model was used to distinguish the baseline Oithona spp. dynamic from noisy and unrelated processes. The physical and biological systems at L4 are influenced by natural cyclic variability on seasonal and decadal scales. Following this natural cyclic variability, the stable hysteresis assumption (Bruun et al., 2017) was applied to the DFSA-derived baseline Oithona spp. dynamic to predict the 2018 dynamic had typical environmental conditions prevailed.

Reproductive and Mortality Rate Estimation

All equations used to calculate rates of reproduction and mortality are provided in Table 2. In situ EPRs (eggs female⁻¹ d⁻¹) of O. similis were calculated from adult female and egg densities, using the egg-ratio method (Edmondson et al., 1962; Checkley, 1980). Egg hatch times were determined from the equation of Nielsen et al. (2002). Total reproductive output (TRO, eggs m⁻³ d⁻¹) was calculated as the product of EPR and adult female density (Maud et al., 2015). The Bêlehrádek equation was used to calculate development time as a function of temperature for each life stage from egg to copepodite CV. For egg and post-hatch stages, we used the value α = 7.6998, as determined for eggs by Nielsen et al. (2002). Values for a were derived from the stage durations at 15°C in Sabatini and Kierboe (1994).

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Equation</th>
<th>References</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hatch time (HT, d⁻¹)</td>
<td>HT = 1504.5 (T + 7.6998)²⁻⁰⁵</td>
<td>Bêlehrádek equation using parameters described for O. similis by Nielsen et al. (2002)</td>
</tr>
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<tr>
<td>Egg production rate (EPR, eggs female⁻¹ d⁻¹)</td>
<td>EPR = E/(F x HT)</td>
<td>Edmondson et al., 1962; Checkley, 1980</td>
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<tr>
<td>Total reproductive output (TRO, eggs m⁻³ d⁻¹)</td>
<td>TRO = EPR x F</td>
<td>Maud et al., 2015</td>
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<tr>
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<tr>
<td>Development time (D, d)</td>
<td>D = a(T + α)⁻ᵇ</td>
<td>Bêlehrádek equation using parameters described for O. similis by Hirst and Ward (2008)</td>
</tr>
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</tr>
<tr>
<td>Mortality rate, sub-adult stages (β, d⁻¹)</td>
<td>β = (ln(N₀/Nₙ)/N₀) + 1</td>
<td>Mullin and Brooks, 1970</td>
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<tr>
<td>Mortality rate, juvenile–adult pair (β, d⁻¹)</td>
<td>β = (ln(N₀/Nₙ)/N₀) + 1</td>
<td>Aksines and Ohman, 1996</td>
</tr>
</tbody>
</table>

 bảng 2: Equations used to calculate Oithona similis reproduction and mortality rates.

$E = \text{egg density (ind m}^{-3}); F = \text{female density (ind m}^{-3}); T = \text{temperature (°C)}; b = 2.05; \alpha = 7.6998 \text{ for eggs and post-hatch stages}; values for } \alpha \text{ from Table 2 in Hirst and Ward (2008); b = mortality rate (d}^{-1}); N = \text{stage density (ind m}^{-3})$. 

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For further detail on these parameters and their application in calculating mortality rates, we refer the reader to Hirst and Ward (2008). Depth-integrated temperature was used in all calculations to incorporate the complete range of temperatures to which the population is potentially exposed.

Development times of each life stage were incorporated into the vertical life table (VLT) method (Mullin and Brooks, 1970; Aksnes and Ohman, 1996) to estimate mortality rates (β, d⁻¹) across egg, juvenile and adult stages, in consecutive pairings of these life stage categories. The VLT method determines mortality across stages from the density and duration of the stages (Aksnes et al., 1997). We grouped naupliar and copepodite stages (as juveniles), with the aim of reducing errors in estimating stage duration, indeed grouping the naupliar and copepodite stages reduced the number of negative mortality rate values seen using higher stage resolution. The primary assumption of the VLT method is that there is no trend in recruitment to a stage over the total duration of the stage pair. We therefore applied the method of Hirst et al. (2007) to determine the impact of recruitment rate on our mortality rate estimates for the egg-juvenile stage pair. Data for egg production were grouped into the periods December 2017–February 2018, March–May 2018, June–August 2018 and September–October 2018. For each period, the slope of the natural log of EPR over time was obtained via linear regression. All slopes were an order of magnitude lower than the mortality rates of the egg-juvenile stage pair, thus recruitment trends should not have significantly affected the mortality rate estimates. A major advantage of the VLT method is that it reduces the complication of advection, providing that advection equally affects all individuals in a sample (Aksnes et al., 1997), making it appropriate for our analysis.

RESULTS

L4 Physical Environment: Time Series Comparison

The year 2018 was characterised by exceptionally low water temperatures between weeks 7 and 19, followed by a rapid rate of increase in the depth-integrated temperature at 0.41°C week⁻¹, from a minimum of 7.84°C in week 10, to more typical summer temperatures of 14.8°C by week 27 (Figures 1A,B). In comparison, the baseline rate of increase in mean depth-integrated temperature over the same time frame was 0.24°C week⁻¹ (Figure 1B). Surface NHF also underwent a rapid rate of increase (32.7 W m⁻² week⁻¹) in 2018, from −254 W m⁻² in week 9 to 237 W m⁻² in week 24, substantially faster than the baseline of 13.7 W m⁻² week⁻¹ over the same time period (Figure 1C).

Oithona similis Population Dynamics: Time Series Comparison

Across the time series (1988–2018), mean (± SE) annual Oithona spp. density was generally stable (Figure 2). One exception was the year 2000, for which population densities were exceptionally high for several weeks in spring and autumn, although for most of the year densities were more typical of the time series. Linear regression analysis showed that mean annual Oithona spp. density had no significant correlation with mean annual SST or total prey biomass (P > 0.05). The day at which Oithona spp. density reached the 25th cumulative percentile of the annual density, representing the start of the season, was not correlated with mean SST in the period February to April or May to July (P > 0.05). The duration index of the population (time between the 25th and 75th cumulative percentiles) ranged from 24 to 193 days. The duration index lacked any significant linear correlation with mean SST between February to April or May to July (P > 0.05). Mean annual density, timing of the start of the season and duration index of Oithona spp. also exhibited no correlation with the rate of water column stratification at the start of the season.

Dominant frequency state analysis determined that the statistically significant modes of variability providing the baseline seasonal climatology for both physical and biological systems at L4 were 1, 1/2 and 1/3 year cyclic processes (Figures 3A,B), plus 4 and 10.5 year cycles (Figure 3C). These five frequency states for the Oithona spp. dynamic explain 27% (R²) of the signal variability. There was no significant trend term across the 1988–2017 time series, indicating that mean Oithona spp. density has not increased or decreased across the time series. Over 1988–2017, the baseline Oithona spp. dynamic exhibited a clear bimodal cycle, with a maximum in week 20 and secondary peak in week 38 (Figure 3B). The Oithona spp. population density from the DFSA-derived 2018 prediction was 19% larger than in 2017 (Figure 3D). Timing of the spring peak in the 2017–2018 study (Figure 3E) was consistent with the DFSA-derived Oithona spp. dynamic for both the baseline (Figure 3B) and prediction (Figures 3D,E). However, the secondary peak in the 2017–2018 study for both the Niskin bottle and WP2 net data occurred several weeks earlier than the DFSA-derived prediction (Figure 3E). The median Oithona spp. density in 2018 was ~37% larger for the Niskin bottle data than both the actual and predicted values obtained from the WP2 net data; evidence for the improved sampling efficiency of our 2017–2018 study. Due to this incompatibility between Niskin bottle and WP2 net data, we do not directly compare Oithona spp. densities between these two sampling strategies, but instead focus on the seasonal trends, which appear to be equally well-represented in both sampling strategies.

L4 Trophic Environment in 2017–2018

The pico- micro-plankton community exhibited distinct seasonality in biomass and vertical distribution (Figure 4). Ciliate carbon biomass maxima occurred in week 24, with a relatively wide depth-distribution. Subsequently, carbon biomass of round-shaped dinoflagellates exhibited a single maximum in week 32, concentrated around 10 m. Two peaks in ellipsoid-shaped dinoflagellate carbon biomass occurred between weeks 22–32 at around 25 m, followed by a third peak between weeks 36 and 38 in the 10 m layer. Diatom carbon biomass was highest between weeks 18 and 27, predominantly in the lower half of the water column. The fluorescence signal recorded by the CTD showed a corresponding depth profile over this time period,
FIGURE 1 | (A) Water column temperature for the 2017–2018 study (Ocean Data View, Schlitzer, 2018); (B) depth-integrated temperature for the 2017–2018 study and weekly mean (± SD) depth-integrated temperature for the time series baseline (2002–2017); (C) surface net heat flux (NHF) for the 2017–2018 study and weekly mean (± SD) surface NHF for the time series baseline (1988–2017). In (B, C), baseline standard deviation is represented by the grey band. Vertical dashed lines mark the onset and breakdown of stratification, indicated by the switch in surface NHF from negative to positive (week 13), and positive to negative (week 39) during the 2017–2018 study.

FIGURE 2 | Mean (± SE) annual Oithona spp. density for each year of the L4 time series (1988–2018), compared against the mean (± SD) Oithona spp. density over the full time series. In the latter, the mean is represented by the solid line, and the standard deviation is represented by the grey band.

evidence that the diatom cells were alive, and not a sinking mass of a decaying bloom. Large flagellates exhibited several biomass peaks between weeks 15 and 32, reaching maximum biomass in weeks 35–38 at 10 m. The pico- nano-plankton occurred predominantly at the top 10 m of the water column, although heterotrophic taxa had a generally wider vertical distribution compared to autotrophic taxa. An initial peak in autotrophic flagellate biomass in week 27 was followed by a
biomass maximum in week 38. Heterotrophic nanoeukaryotes were relatively abundant from week 26 onwards, with maximum biomass in week 35. Heterotrophic bacteria biomass has highest in weeks 24 and 35–37. *Synechococcus* was rare for most of the year, with a single biomass maximum in week 36. Of the above taxa, autotrophic flagellates had the greatest contribution to carbon biomass.

**2017–2018 Study: Oithona similis Population Dynamics**

In spring (weeks 11–22), depth-integrated mean (± SE) *O. similis* densities were 1750 ± 325 ind m⁻³ for nauplii, 2830 ± 621 ind m⁻³ for copepodes, 660 ± 167 ind m⁻³ for adult females, and 82.7 ± 20.1 ind m⁻³ for adult males. Over summer (weeks 23–35) these densities increased substantially in the naupliar (2080 ± 553 ind m⁻³) and copepodite stages (4710 ± 966 ind m⁻³), whereas the increase in adult density was negligible for both females (675 ± 144 ind m⁻³) and males (84.1 ± 17.0 ind m⁻³). The initial increase in *O. similis* population density between weeks 15 and 19 preceded that of prey biomass, whereas the population decline in weeks 35–38 occurred when prey biomass was maximal (Figure 5A). Over these time periods, mean (± SE) depth-integrated total prey carbon biomass, excluding bacteria, increased from 112 ± 17.3 to 233 ± 75.02 µg C L⁻¹. The *O. similis* population exhibited clear ontogenetic differences in vertical
FIGURE 4 | Prey carbon biomass over the 2017–2018 study (Ocean Data View, Schlitzer, 2018). Vertical dashed lines mark the onset and breakdown of stratification, indicated by the switch in surface NHF from negative to positive (week 13), and positive to negative (week 39) during the 2017–2018 study.
distribution, with all life stages being relatively scarce in the top few metres (Figure 5B). Nauplii were most abundant at 10 m, whereas copepodites had a wider depth range of 10–25 m. Adult stages had the widest depth range, occurring predominantly in the lower half of the water column. It therefore appears that the depth distribution of *O. similis* at L4 increases with development to later life stages.

**Reproductive and Mortality Rates**

*Oithona similis* EPR seasonality over 2017–2018 resembled that of the baseline EPR dynamic, with minimum rates in winter, followed by an increasing trend towards maximum rates in summer (Figure 6A). In 2018, the summer maximum occurred several weeks earlier than that of the baseline EPR dynamic. However, during the summer (weeks 23–35) there was little difference between mean (± SE) EPR in 2018 (2.59 ± 0.45 eggs female$^{-1}$ d$^{-1}$) and the baseline (2.60 ± 0.35 eggs female$^{-1}$ d$^{-1}$). At the start of 2018, EPR was substantially below the baseline until approximately week 18 (Figure 6A). As TRO is a function of EPR and adult female density, direct comparison between the 2017 and 2018 study and the baseline is not possible on account of the different sampling efficiencies of the Niskin
bottle and the fine mesh nets. Nonetheless, in agreement with
the baseline dynamic, TRO in 2018 was highest in spring
and summer (Figure 6B), coinciding with peaks in adult
density and EPR, respectively. Mortality rates (β) were
considerably higher and more variable in the egg-juvenile
stage pair compared to the juvenile-adult stage pair, although both
exhibited an overall increase in mortality rate towards summer
(Figure 6C). Maximum mortality rate for the egg-juvenile stage
pair was 0.54 d$^{-1}$ in week 38. In contrast, maximum mortality
rate for the juvenile-adult stage pair was just 0.19 d$^{-1}$, and
occurred in week 26.

DISCUSSION

Our study shows that the numerically dominant cyclopoid
species $O. similis$ has maintained generally stable population
dynamics over the L4 time series, despite high variability in both
the physical and biological environment, as evidenced in our
1-year intensive study. The biomass dominant calanoid species
$C. helgolandicus$ has also maintained stable population dynamics
at L4 (Maud et al., 2015). In contrast, important species at
this site, such as $Acartia clausi$, $Pseudocalanus elongatus$ and
$Temora longicornis$, have undergone major declines in population
density (Edwards et al., 2020). Here, we compare the population
dynamics of $O. similis$ and $C. helgolandicus$ at L4 to investigate
which traits may contribute to population stability and resilience
in this highly variable environment.

$Oithona similis$ Population Dynamics:
Time Series Comparison

Despite high climatic variability over the 2017–2018 study,
$O. similis$ maintained similar population densities and bimodal
cycle as that over the whole 30-year time series. Evidence of
$Oithona$ spp. population stability at L4 has been reported
previously (Atkinson et al., 2015; Castellani et al., 2016; Cornwell
et al., 2018). Moreover, bimodal cycles in density are reported
in $O. similis$ populations from the White and Barents Seas
(Dvoretskii, 2007; Dvoretsky and Dvoretsky, 2009), Loch Striven,
Scotland (Marshall, 1949), and the Southern Ocean (Fransz
and Gonzalez, 1995). In contrast, a single density peak in
spring (Castellani et al., 2016), summer (Zamora-Terol et al., 2014), and late autumn (Lischka and Hagen, 2005), has been documented in Mediterranean, subarctic and Arctic *O. similis* populations, respectively.

We found that mean SST was not correlated with either the mean annual density or phaenological timings of the L4 *Oithona* spp. population, supporting the results of a previous L4 time series study of *Oithona* spp. (Castellani et al., 2016). However, temperature is a major driver for other *O. similis* populations, both in terms of density (Ward and Hirst, 2007; Dvoretzky and Dvoretzky, 2009; Castellani et al., 2016), and biomass (Castellani et al., 2007). The importance of temperature in driving population dynamics may depend on the proximity of the population to their thermal range edge, with temperature effects becoming stronger closer to this edge. We conclude that at station L4, it appears temperature is, presently and historically, not the most important driver of *O. similis* population dynamics in terms of density or phaenology, either at the intra- or inter-annual scale.

### L4 Trophic Environment

The various numerical and conceptual models developed to explain the plankton seasonality at station L4 are primarily based on trophic parameters (Atkinson et al., 2018). Considering the preference of *Oithona* spp. for motile prey (Atkinson, 1995, 1996; Castellani et al., 2005), the L4 *Oithona* spp. population has been suggested to benefit from the increased summer abundance of motile prey, such as ciliates and dinoflagellates (Kenitz et al., 2017). However, our results suggest that other factors may have a greater impact on the population dynamic. For example, the autumn decline in *O. similis* population density in 2017–2018 would not have been predicted based on the high biomass of motile prey at this time. Indeed, the importance of food in driving *O. similis* population dynamics at L4 is uncertain (Castellani et al., 2016; Cornwell et al., 2018; Djeghri et al., 2018), and we propose that other factors likely override the effect of food, as discussed below.

### Vertical Profile: Population Dynamics and Trade-Offs

Depth-resolved sampling enabled us to study the vertical profile and seasonality of all *O. similis* life stages. Nauplii were most abundant in the 10 m layer, where pico-nano-plankton biomass was generally greatest. One explanation for this distribution is that growth is essential for survival among early life stages. Furthermore, occupying the warm upper layer would accelerate ontogenetic development, resulting in an earlier temporal onset of improved motility and escape performance, which could potentially reduce predation mortality (McLaren, 1963; Fiksen and Giske, 1995; Kiørboe and Sabatini, 1995). However, occupying the upper layer increases exposure to visual predators. There is thus a trade-off between growth and survival among earlier life stages, which potentially forces them to adopt more risky feeding behaviours (Hirst and Bunker, 2003). In contrast, our finding that adult stages occupy greater depths may reflect their ability to live for longer periods with less food (Hirst and Bunker, 2003), potentially enabling adults to reduce feeding in favour of predation avoidance. Thus, the trade-off between growth and survival may be different for adults compared to earlier life stages.

All life stages were relatively scarce in the top few metres of the water column despite periods of high food biomass in this layer. Other factors, such as ultraviolet stress (Fileman et al., 2017), visual predators and turbulence, may be forcing the population out of this layer. Although turbulence can enhance prey encounter rates for ambush predators (Kiørboe and Saiz, 1995; Saiz and Kiørboe, 1995), *Oithona* spp. populations have been observed to occupy greater depths during periods of increased surface turbulence (Incze et al., 2001; Visser et al., 2001; Maar et al., 2006), although nauplii may be less able to avoid turbulence due to their low swimming capacity (Lagadeuc et al., 1997; Andersen et al., 2001; Maar et al., 2003). However, considering that nauplii were also relatively rare in the uppermost layer, the impact of turbulence in determining the *O. similis* vertical profile at L4 is debateable. Another factor to consider is salinity, although the generally weak salinity gradient at L4 makes it unlikely that salinity has a strong impact on the *O. similis* vertical profile at this site. In contrast, at freshwater influenced sites, salinity can be a major driver of vertical distribution, with distinct avoidance of low salinity upper layers, as observed for all life stages of *O. similis* in the Bornholm Basin, Baltic Sea (Hansen et al., 2004), and Hylsfjord, Norway (Andersen and Nielsen, 2002). At other sites, the factors driving the *O. similis* vertical profile are less clear. For example, vertical distribution was not significantly correlated with surface salinity, temperature or chlorophyll across the North Sea (Maar et al., 2006), or with phytoplankton biomass or water column stratification in Chaleur Bay, Gulf of Saint Lawrence (Lagadeuc et al., 1997). Evidently, the factors driving *O. similis* vertical profile vary between populations, and depend on the environment in which each population occurs.

It must be acknowledged that in our study, all sampling events were conducted at mid-morning. Thus, our results do not account for potential diel vertical migration, although studies from other sites generally report a lack of diel vertical migration in *O. similis* (Turner and Dagg, 1983; Lagadeuc et al., 1997; Andersen and Nielsen, 2002; Eiane and Ohman, 2004; Hansen et al., 2004; Maar et al., 2006). However, seasonal vertical migration has been observed in the *O. similis* population of Kongsfjorden, Svalbard, which occurs predominantly in the upper 50 m layer in summer and below 100 m in winter (Lischka and Hagen, 2005). Conversely, we found that the vertical profile of each life stage appears relatively constant throughout the year at L4, as also reported for *O. similis* across the Arctic Ocean (Ashjian et al., 2003).

### The Balance Between Reproduction and Mortality

Egg production rate seasonality over the 2017–2018 study was similar to the baseline dynamic, although the summer EPR maximum was several weeks earlier than that of the baseline. In sac-spawning species such as *O. similis*, the production of new egg
clutches is limited by the rate at which the previous eggs hatch (Ward and Hirst, 2007). As higher temperatures increase rates of embryonic development and hatching, and consequently reduces the time between the production of egg clutches, EPR increases as a result (Nielsen et al., 2002). The early EPR peak in 2018 could have resulted from accelerated development of the spring cohort due to the rapid rate of seasonal warming. Coincident with the low water column temperatures at the start of 2018, EPR was substantially below the baseline seasonal average until around week 18, at which point water column temperature had begun to reach more typical levels.

Female body size of copepod species can vary substantially and also impact fecundity (Sabatini and Kiorboe, 1994). At station L4, *O. similis* adult female body size conforms to the temperature–size rule (Atkinson, 1994), decreasing from a maximum in spring following the low winter temperatures, to a minimum in autumn after the annual temperature maximum (Cornwell et al., 2018). The seasonal trend in *O. similis* female body size is therefore counter to that of EPR, indicating that for the L4 *O. similis* population, fecundity appears more influenced by positive temperature effects than reductions associated with smaller female body size. In contrast, EPR of the calanoid copepod *C. helgolandicus* at L4 is positively correlated with adult female body size, with both EPR and female body size peaking in spring (Cornwell et al., 2018). Therefore, future increases in temperature fluctuations may impact fecundity via its effects on physiology of the species, including development rates, and also body size, the strength of these effects being species specific.

The spring and summer peaks in TRO and naupliar density indicate the production of two main generations over the 2017–2018 study, conforming to the bimodal cycle of the baseline dynamic. The production of two generations per year has been previously reported in other *O. similis* populations (Digby, 1954; Lischka and Hagen, 2005; Dvoretsky, 2007; Dvoretsky and Dvoretsky, 2009). The small peak in naupliar density in spring 2018 coincided with maximum adult density, indicative of high recruitment early in the year. Subsequently, there was a general increase in mortality rates over the remainder of the study. Mortality rates were far greater and more variable across the egg–juvenile stage pair compared to the juvenile–adult stage pair, which agrees with previous studies on other *O. similis* populations reporting higher mortality rates in early life stages (Marshall, 1949; Fransz and Gonzalez, 1995; Eiane and Ohman, 2004; Hirst and Ward, 2008; Thor et al., 2008), and low recruitment success (McLaren, 1969). The high mortality rates in autumn, particularly for the egg–juvenile stage pair, may have contributed to the *O. similis* population decline.

**What Traits Provide Population Stability in Copepods?**

Despite the overall decline in copepod densities at station L4 over the 30-year time series (Edwards et al., 2020), *O. similis* has maintained stable population densities and seasonality. The calanoid copepod *C. helgolandicus* has also maintained stable population densities over the L4 time series (Maud et al., 2015; Edwards et al., 2020). The inter-annual variation in mean population densities is similar in both species, with coefficients of variation (CV) of 0.49 and 0.44 for *O. similis* and *C. helgolandicus*, respectively. For context, copepods that have undergone long-term population declines at L4, namely *Pseudocalanus elongatus*, *Temora longicornis* and *Acartia clausi*, show greater inter-annual variability in their mean population densities, with CVs of 0.54, 0.61 and 0.71, respectively. Both *O. similis* and *C. helgolandicus* have been intensively studied at L4, providing a broad data set which we use here to suggest traits contributing to population stability (Table 3).

*Oithona similis* and *C. helgolandicus* contrast markedly in body size, feeding- and reproductive modes, with *C. helgolandicus* generally exhibiting greater reproductive output and high mortality in early life stages. Conversely, small cyclopoid copepods such as *O. similis* possess lower rates of feeding, growth, reproduction and mortality compared to calanoid species of equivalent size, and these lower rates have been suggested to contribute to the stability of cyclopoid populations (Paffenhofer, 1993). In contrast to the bimodal cycle and summer EPR maxima of *O. similis*, as described in this study, the population density

<table>
<thead>
<tr>
<th>Trait</th>
<th>Oithona similis</th>
<th>Calanus helgolandicus</th>
</tr>
</thead>
<tbody>
<tr>
<td>Reproductive mode</td>
<td>Sac-spawning</td>
<td>Broadcast-spawning</td>
</tr>
<tr>
<td>Feeding mode</td>
<td>Ambush-feeding</td>
<td>Suspension-feeding</td>
</tr>
<tr>
<td>Female total length (mm)</td>
<td>0.68–0.96</td>
<td>2.60–3.28</td>
</tr>
<tr>
<td>Mean (±95% CI)% change in CI per 1°C increase in SST</td>
<td>−2.33 ± 0.5</td>
<td>−7.46 ± 1.6</td>
</tr>
<tr>
<td>Phaenological change (days per 1°C increase in SST)</td>
<td>−4</td>
<td>−4</td>
</tr>
<tr>
<td>Mean population density inter-annual variation (CV)</td>
<td>0.49</td>
<td>0.44</td>
</tr>
<tr>
<td>Annual mean mortality rate (de−1) of early life stages</td>
<td>Egg-juvenile: † 0.18</td>
<td>Egg-NI: 1.50 (Hirst et al., 2007)</td>
</tr>
<tr>
<td>Annual mean mortality rate (de−1) of later life stages</td>
<td>Juvenile-adult: † 0.09</td>
<td>CV-adult female: 0.10 (Hirst et al., 2007) 0.09 (Maud et al., 2018)</td>
</tr>
<tr>
<td>Annual mean EPR (eggs female−1 de−1)</td>
<td>*1.79</td>
<td>*14.3</td>
</tr>
<tr>
<td>Mean EPR inter-annual variation (CV)</td>
<td>0.13</td>
<td>0.19</td>
</tr>
</tbody>
</table>

CI = female carbon mass (µg C female−1); SST = sea surface temperature; CV = coefficient of variation; †L4 time series data; † present study. Phaenological change follows the sign convention of minus means earlier when warmer.
of *C. helgolandicus* at L4 follows a considerably weaker bimodal cycle with a summer maximum, and peak EPR during the spring diatom bloom (Irigoin and Harris, 2003; Maud et al., 2015). These inter-specific differences in seasonality are to some extent explained by the factors driving EPR in each species. Temperature is significantly positively correlated with EPR in *O. similis* at L4 (Cornwell et al., 2018) and at other sites (Ward and Hirst, 2007; Dvoretsky and Dvoretsky, 2009; Zamora-Terol et al., 2014). Conversely, *C. helgolandicus* EPR is more dependent on food availability than temperature (Bonnet et al., 2005), as observed in the L4 population (Maud et al., 2015; Cornwell et al., 2018). From this analysis, it is clear that species with contrasting functional traits can maintain stable population densities and seasonality.

Temperature is considered to be a fundamental driver of phaeology and species density at the macroecological scale of ocean basins (Beaugrand and Kirby, 2018; Beaugrand et al., 2019). However, at individual study sites such as L4, these temperature effects on population dynamics are less evident. For instance, the peaks in *O. similis* population density at L4 occur at both the coldest and warmest times of the year. At the scales of seasonal and inter-annual variability at a single site, other factors may be more important in shaping population trends. For example, density-dependent mortality processes have been suggested to provide population stability in *C. helgolandicus* at L4 (Hirst et al., 2007; Maud et al., 2015, 2017, 2018). Likewise, our results lead us to propose mortality as a major driver of the stable population dynamic of *O. similis* at L4, which is upheld even during periods of high climatic variability.

**DATA AVAILABILITY STATEMENT**

The datasets generated for this study are available on request to the corresponding author.

**REFERENCES**


**AUTHOR CONTRIBUTIONS**

LC designed the methodology, performed field and laboratory work regarding data collection, analysed the data, and wrote the manuscript. EF, JB, AH, GT, HF, CL, and AA contributed to the drafting of the manuscript. EF provided support with FlowCam analysis and classification. JB significantly improved the time series analysis by introducing modelling techniques. AH provided guidance in calculating mortality rates. GT provided the flow cytometry data. TS provided the net heat flux data. AM provided the L4 zooplankton time series data.

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**Conflict of Interest:** The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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