

Spatially variable growth responses to warming in Atlantic Sea Scallops (*Placopecten magellanicus*)

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Abstract

The Atlantic sea scallop (*Placopecten magellanicus*) is an important commercial species in the U.S., and its growth rate is strongly modulated by temperature. In this study, we quantified the interannual variations in scallop growth rate and thermal stress intensity in the Mid-Atlantic Bight (MAB) from 2000 to 2018. The results showed that the scallop growth rate variations were overall synchronized between the shallow (< 60 m) and deep (≥ 60 m) MAB prior to 2015. The response of growth rate to warming in 2015 and 2016 showed marked spatial heterogeneity: scallops in the shallow subregions grew more slowly in the warm years, whereas higher temperature contributed to elevated growth rates in the deep subregions. We developed a dynamic energy budget model to explain the distinct growth rate response to warming at different depths. The model results showed that, in 2016, bottom temperatures in the shallow subregions exceeded the optimal range, limiting the energy available for growth. In contrast, warming created more favorable thermal conditions in the deep habitats. This work reveals the spatiotemporal patterns of scallop growth rates in the MAB and contributes to a quantitative understanding of thermal stress, supporting the development of science-based management strategies under future warming.

Keywords Atlantic sea scallop, growth rate, temperature, bioenergetics

Introduction

The Atlantic sea scallop (*Placopecten magellanicus*), distributed on the continental shelf between Cape Hatteras, NC, and the Gulf of St. Lawrence, is one of the most economically valuable fishery species in the U.S. (Stewart and Arnold 1994). Although the total sea scallop landings in the U.S. fishing grounds (e.g. Mid-Atlantic, Georges Bank, and the Great South Channel) dropped dramatically from the 1970s to the mid-1990s due to overfishing (Edwards and Murawski 1993), the implementation of a series of effective fishery management strategies has rebuilt the depleted populations since 1994 (Hart and Rago 2006). Sea scallop stocks successfully recovered due to this species' rapid individual growth rate and low natural mortality, and the ex-vessel revenues of sea scallop fisheries have increased to around \$500 million per year in recent decades (Hart 2003, Coleman et al. 2022, NOAA 2023).

Sea scallop growth rate is one of the most crucial biophysiological traits indicating the health condition of individuals and suitability of surrounding habitats (MacDonald and Thompson 1985, Hart and Chute 2009, Hodgdon et al. 2020, Coleman et al. 2021).

Given the importance of scallop growth in biomass production, management plan development, and aquaculture site selection, previous studies assessed the spatiotemporal patterns of scallop growth rates in natural habitats and their associated drivers. Laboratory experiments, field measurements, and model results indicated that various factors (e.g. temperature, salinity, water depth, density, food condition, current fields, CO₂ level, fishing activities, sediment resuspension, and parasite infection) and their joint effects modulate sea scallop growth (Grant et al. 1997, Pilditch and Grant 1999a, 1999b; Hart and Chute 2009, Inglis et al. 2016, Siemann et al. 2019, Coleman et al. 2022, Pousse et al. 2023). Among all these factors, temperature has received much attention due to its profound effects on scallop metabolic processes (e.g. clearance, ingestion, respiration, and excretion rates; MacDonald and Thompson 1985, Shumway et al. 1988, Cranford and Grant 1990, Grant and Cranford 1991). The optimum temperature for scallop growth is between 10–15°C, and higher temperatures limit scallop growth and survival due to net energy loss associated with decreased feeding and elevated energy costs of maintenance (Zang et al. 2022a, Stewart and Arnold 1994, Hart

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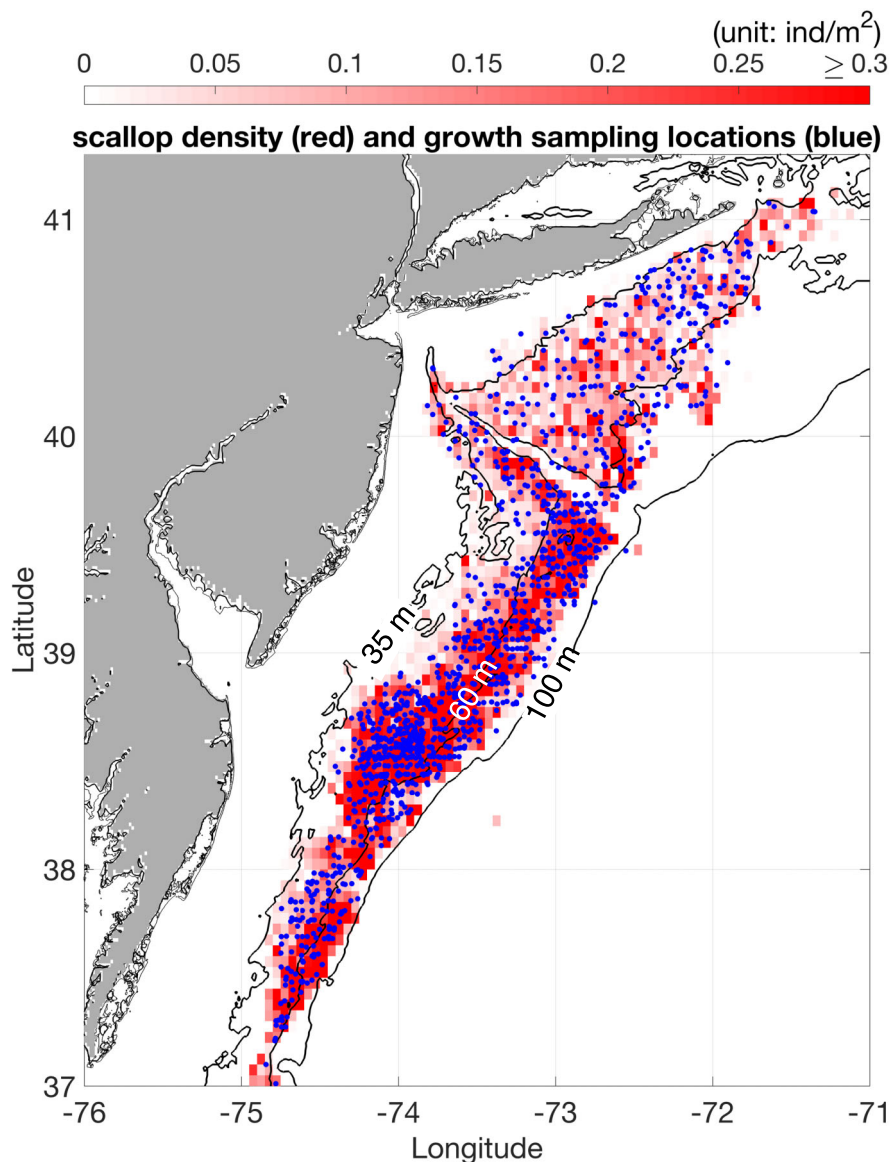


Figure 1 The climatology of Atlantic sea scallop density in the MAB from 2000 to 2018 (data source: NOAA sea scallop dredge survey data). The three black contour lines are 35, 60, and 100 m isobaths, respectively. The blue dots indicate locations where sea scallop shells were collected for growth measurements. The total number of scallop shells collected for growth measurements in the MAB is 8 045.

and Chute 2004, Coleman et al. 2022). Additionally, the intricate nonlinear interactions between temperature and other stressors further confound the thermal response of scallop growth. Coleman et al. (2021) found that at high scallop density, a negative correlation between temperature and growth could occur during some of the most productive periods when temperature was favorable to the rapid growth of individuals, indicating that high density lowered the optimal temperature for scallop growth. Food availability is another critical factor that interacts with temperature effects on scallop growth: sufficient high quality food supply may buffer the negative impacts of thermal stress on growth, while food deficiency can amplify its negative effects, especially for large individuals with higher commercial values (Zang et al. 2022a, MacDonald and Thompson 1985, 1986, Pilditch and Grant 1999a). Furthermore, laboratory experiments under various temperature and pCO₂ conditions showed that the interactive ef-

fects of warming and ocean acidification (OA) contributed to declines in scallop growth and survivorship (Cameron et al. 2022, Pousse et al. 2023).

The Mid-Atlantic Bight (MAB; Fig. 1) represents the southern extent of sea scallop habitat on the Northeast U.S. Shelf with relatively strong thermal stress (maximum bottom temperature > 18°C; du Pontavice et al. 2023), making its scallop populations more vulnerable to warming than those at higher latitudes. Bottom temperature in the MAB has increased at an average rate of ~0.35°C per decade with strong spatial heterogeneity (Pershing et al. 2015, Saba et al. 2016, Kleisner et al. 2017, du Pontavice et al. 2023). Marine heatwaves in 2012 and 2016 acutely increased water temperature in the MAB and surrounding areas, significantly affecting thermal conditions and fishery yields (Mills et al. 2013, Pershing et al. 2018). Climate model projection results suggested that warming will continue in the MAB at the rate of ~0.05°C yr⁻¹

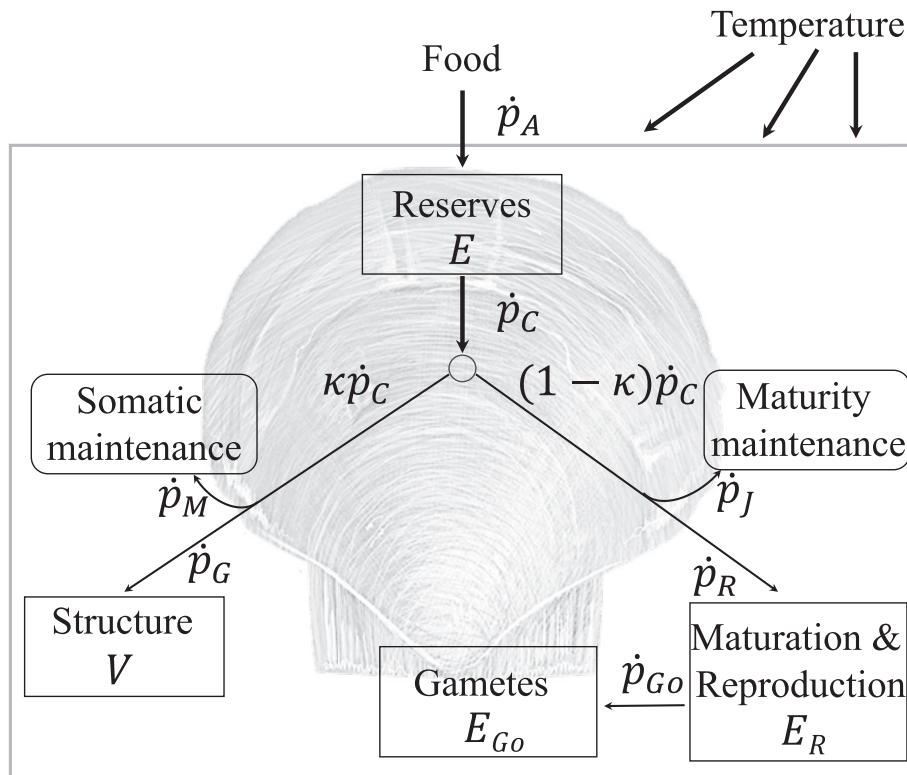


Figure 2 Conceptual scheme of the DEB model applied to sea scallop. Four state variables are Reserves (E), Structure (V) and Maturity & reproduction (E_R), and Gametes (E_{Go}) in rectangular boxes. \dot{p}_A , \dot{p}_C , \dot{p}_M , \dot{p}_G , \dot{p}_J , \dot{p}_R , and \dot{p}_{Go} are energy fluxes estimated in the model (see Table S1 in supplementary material for more detailed information). Arrows indicate energy flux directions.

layers in the vertical direction. Measured temperature, salinity, and current profiles are assimilated into the model results to enhance the quality of its outputs (Chen et al. 2009, 2021b). The modeled water temperature products have been well-calibrated and widely used in previous studies (Zang et al. 2022b, Chen et al. 2011, Li et al. 2015, 2021b). Since the simulations of FVCOM-GOM3 were terminated in 2016, bottom temperature data after 2016 were extracted from FVCOM-GOM4. FVCOM-GOM4 is an updated version of the FVCOM-GOM3 with the same spatial coverage (Fig. S7b), and its grid resolution is higher in nearshore areas (e.g. Penobscot Bay, Boston Harbor, and Narragansett Bay) to better simulate coastal processes. The hourly hydrodynamic model outputs were downloaded from the data server of the University of Massachusetts Dartmouth (<http://fvcom.smast.umassd.edu>) and converted to daily average results.

To assess thermal stress intensity in the MAB, the modeled bottom temperature data were spatially interpolated to the locations of scallop samples using the corresponding shell height increment year determined in Section 2.1. Since the optimum temperature for sea scallop growth is below 15°C (Zang et al. 2022a, Coleman et al. 2022), we estimated sample-weighted Degree Days > 15°C (DD hereafter) to represent thermal stress intensity:

$$DD_{Y,n} = \sum_{i=1}^{365} \max(T_{i,Y,n} - 15, 0)$$

where $T_{i,Y,n}$ is the daily averaged bottom temperature on Julian day i of year Y for sample n . Since sea scallops in the MAB lay their rings when temperatures peak in the fall season (October to December; Chute et al. 2012), DD for year Y was estimated using tem-

perature data from Nov 1st of year $Y-1$ to Oct 31st of year Y to ensure that the temporal coverage of DD matches that of shell height increment (Section 2.1). To evaluate the sensitivity of DD estimates to the selected time window, we tested two alternative periods (from Oct 1st of year $Y-1$ to Sept 30th of year Y and from Dec 1st of year $Y-1$ to Nov 30th of year Y). Since scallop sampling sites varied among years, we also estimated area-weighted DD in the four subregions to provide sampling-independent estimate of thermal stress.

Sea scallop DEB model

We developed a sea scallop bioenergetic model based on the Dynamic Energy Budget theory (Kooijman 2010). The DEB model simulates energy fluxes through four state variables: 1) the structural volume V ; 2) the reserve E ; 3) the energy allocated to maturity and reproduction E_R , and 4) the energy used to create gametes E_{Go} (Fig. 2). Temperature and food availability are the forcing variables. Energy from food is assimilated into the reserve compartment, and a fixed fraction (κ) of energy flux from the reserve is allocated to support somatic maintenance and structure growth. The rest of the energy ($1 - \kappa$) is used for maturity maintenance, maturation, and reproduction. The primary DEB model parameters for *P. magellanicus* were retrieved from the AmP collection (Lavaud and Kooijman 2020). We adopted the DEB model structure in Lavaud et al. (2014, 2017, 2019, 2021) to simulate sea scallop growth and its response to temperature variation, and this DEB model structure has been successfully applied to simulate the growth and reproduction of other bivalve species (e.g. king scallop and eastern

oyster). Readers are referred to Lavaud et al. (2014, 2017) for more detailed information regarding the DEB model structure. Due to the distinct optimum temperatures for sea scallop feeding and maintenance (Shumway et al. 1988), we applied different parameters in the Arrhenius relationships (i.e. T_{AL} and T_{AH}) for energy assimilation and consumption to better resolve the effects of warming on scallop growth. Two spawning events per year (spring and fall) were incorporated into the DEB model to represent the documented bimodal reproductive dynamics of sea scallops in the MAB (Kirkley and Dupaul 1991). All model equations and parameters are listed in Tables S1 and S2. Daily bottom temperature climatology in the entire MAB scallop habitats was estimated and used to drive the DEB model. The simulation was initiated with a scallop shell height of 40 mm (year 0) and lasted 14 years. The modeled shell height was compared with observed shell height in the MAB based on von Bertalanffy growth parameters estimated by Hart and Chute (2009) ($K = 0.508$ and $L_{\infty} = 133.3$ mm). The simulated tissue weight and gonad weight were compared with the data using the \log_{10} - \log_{10} relationships between shell height and these variables in Langton et al. (1987). It should be noted that scallop growth rates vary spatiotemporally in response to local environmental conditions (Hart and Chute 2009). As the DEB model is designed to simulate an average individual within the population, the modeled growth curve represents the average growth trajectory in the entire MAB rather than individual-level variability. The fixed model structure and parameters (Tables S1 and S2) provides a consistent benchmark that enables us to assess the influence of environmental factors on scallop growth across our study area.

To quantitatively estimate warming effects on sea scallop growth and its spatial heterogeneity, we designed sensitivity tests driven by daily bottom temperature climatology and bottom temperature in the warm year 2016 in the four subregions. All other inputs and parameters were the same to ensure that the differences between model results only resulted from temperature inputs. The simulation period of sensitivity tests was one year, and we used four initial sea scallop shell heights (i.e. 40 mm, 60 mm, 80 mm, and 100 mm) to assess the impacts of warming on scallop growth in different size groups. Since food supply plays a vital role in modulating sea scallop growth while its magnitude is still difficult to quantify in natural habitats (Zang et al. 2022a, Hart and Chute 2009), a baseline value of food limiting factor $f = 0.70$ was used to represent moderate food conditions in the entire MAB. We conducted four additional sensitivity tests with $f = 0.60, 0.65, 0.75,$ and 0.80 to assess the impacts of food availability on scallop growth and its spatial pattern in the MAB.

Results

Spatial pattern of scallop growth rate in the MAB

The median scallop growth rate decreased gradually from ~ 37 mm yr⁻¹ in the smallest size bin (40–45 mm) to 6 mm yr⁻¹ in the largest size bin (125–130 mm) (Fig. 3). In both the SMAB and the NMAB, scallop growth rates across almost all size bins were significantly higher in the shallow subregions (Fig. 3). Although multiple independent comparisons may increase the probability of Type I error, the overall pattern of growth rate differences between shallow

and deep habitats is consistent across size bins. Growth rate differences between the shallow and deep subregions were most pronounced in the 60–80 mm size bins, reaching ~ 5 –9 mm yr⁻¹; whereas differences for smaller and larger scallops were generally below 4 mm yr⁻¹. Unlike the significant growth rate difference between shallow and deep areas, the growth rate differences across latitudes within the same depth range were much smaller (Fig. S8). In the shallow subregions, individuals between 70 and 100 mm grew 1–3 mm yr⁻¹ faster in the NMAB, while only marginal differences were detected in other size groups (Fig. S8a). Growth rates in the deep SMAB and the deep NMAB were quite comparable across all size bins (Fig. S8b). Overall, sea scallop growth rates showed a much stronger and consistent gradient across depths than across latitudes in the MAB, with higher growth rates observed in shallow areas in all size groups.

Interannual variations in normalized scallop growth rate and thermal stress

The significant positive correlations between normalized growth rates in the two shallow subregions (shallow SMAB—shallow NMAB: $r = 0.76, p < 0.05$) and the two deep subregions (deep SMAB—deep NMAB: $r = 0.64, p < 0.05$) indicated that the interannual variations in scallop growth rates were synchronized within the same depth range (Fig. 4). In contrast, no significant correlation was detected between the shallow and deep subregions within the NMAB (shallow NMAB—deep NMAB: $r = 0.41, p > 0.05$) or the SMAB (shallow SMAB—deep SMAB: $r = 0.24, p > 0.05$) from 2000 to 2018. Interestingly, the decoupling of normalized growth rates across depths did not last throughout the entire study period. Correlation analyses using 14-year moving time windows indicated that the interannual variations in normalized growth rates were positively correlated between the shallow and deep subregions prior to 2015 (Table 1), while the correlation coefficients gradually declined and became statistically insignificant after 2015 (Table 1). The time series showed that normalized growth rates in the shallow subregions gradually declined below 0 after 2015 (Fig. 4a and c), whereas growth rates in the deep subregions remained high (deep NMAB: 0.3–0.6; deep SMAB: 0.7–1.0; Fig. 4b and d). The contrasting post-2015 growth patterns between shallow and deep subregions contributed to the insignificant correlations when the time window included years after 2015. We also estimated the interannual variations in normalized growth rates based on only small (≤ 80 mm) and only large (> 80 mm) individuals (Fig. S9). The results showed similar trends to those across all size classes, indicating overall consistent spatiotemporal patterns in scallop growth rates in different size groups. In addition, we estimated scallop growth rate anomalies in the four subregions using observational data and simulated growth rate based on von Bertalanffy growth curve in Hart and Chute (2009) to exclude depth and latitude effects (Supporting Text in Supplementary Material). The results were similar to our normalized growth rate interannual variations with elevated/decreased growth rates in the deep/shallow subregions after 2015, respectively (Fig. S10). A linear mixed-effects model was also used to reconstruct interannual growth anomalies independent of size and depth effects while explicitly accounting for non-independence among shell height increment measurements (Supporting Text in Supplementary Material). The interannual growth anomalies patterns in the

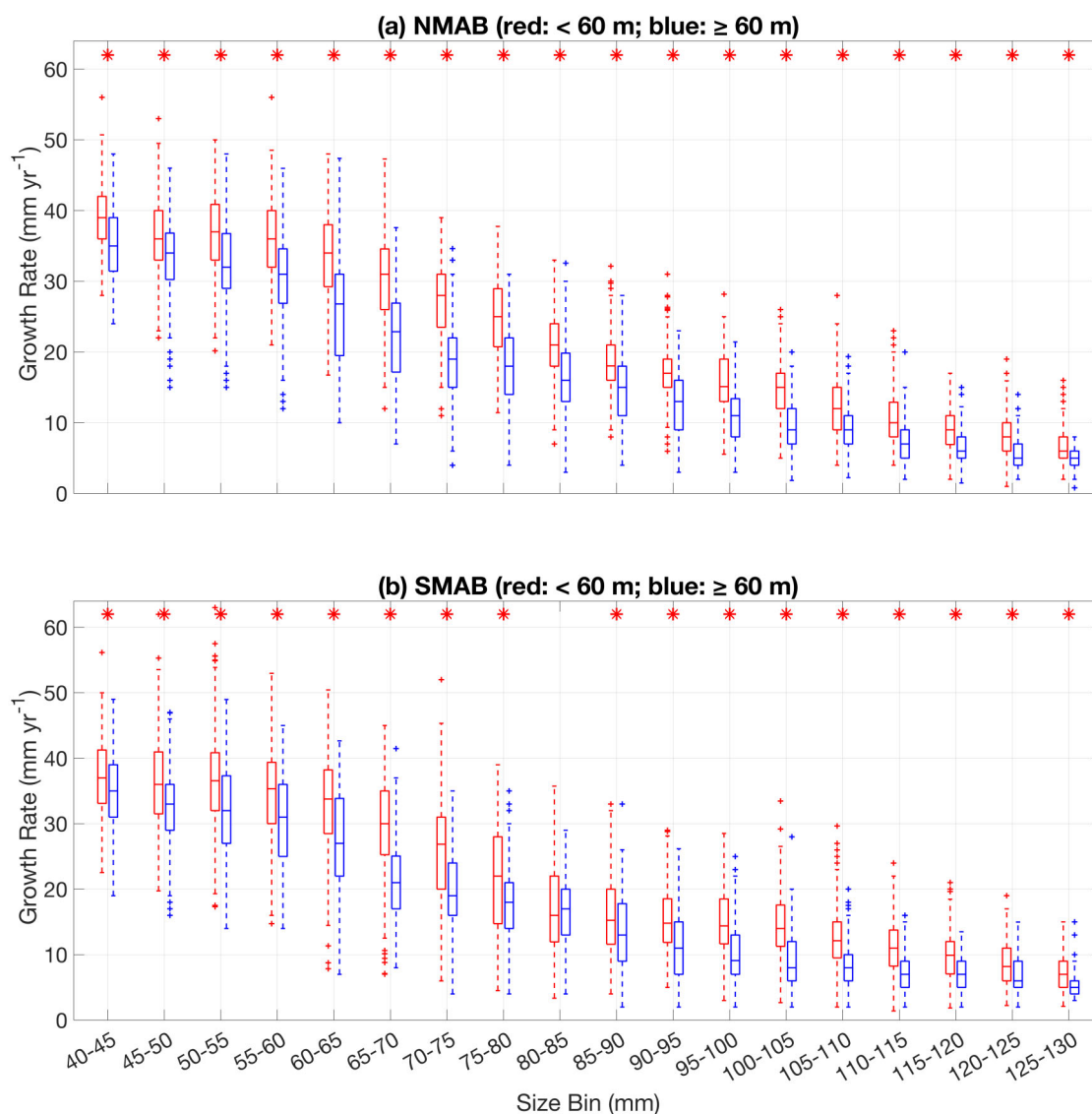


Figure 3 Scallop growth rates across 18 size bins (x-axis) in the northern MAB (NMAB; latitude $\geq 39^\circ\text{N}$; panel a) and the southern MAB (SMAB; latitude $< 39^\circ\text{N}$; panel b). Red and blue colors indicate data in the shallow and the deep subregions, respectively (cutoff depth: 60 m). Mann–Whitney U tests were performed for each size bin to assess statistical significance. Red asterisks indicate significantly higher values in the shallow subregions.

four subregions were consistent with those obtained from the size-normalized growth results (Fig. S11), providing additional support for the spatially heterogeneous growth responses after 2015.

Interannual variations in sample-weighted DD (i.e. thermal stress intensity) in the four subregions stayed low and fluctuated between 0 and $100^\circ\text{C}\cdot\text{d}$ before 2015 (Fig. 5). In 2015 and 2016, DD in shallow subregions increased to $150\text{--}210^\circ\text{C}\cdot\text{d}$, which was followed by a gradual decrease after 2017 (Fig. 5a and c). In the deep SMAB, DD peaked in 2015 and 2016, but its magnitude was lower than that in the shallow subregions (2015: $140^\circ\text{C}\cdot\text{d}$; 2016: $121^\circ\text{C}\cdot\text{d}$; Fig. 5d). Conversely, DD in the deep NMAB was not elevated in 2015 and 2016 (Fig. 5b), suggesting that the scallops in the deep NMAB did not experience strong thermal stress likely due to its higher latitude and greater depth. Although area-weighted DD differed from sample-weighted DD slightly (Fig. S12), both metrics showed stronger thermal stress in 2015 and 2016 in the shallow subre-

gions and relatively weak thermal stress in the deep subregions during the entire study period. Analyses using different time windows (Section 2.2) indicated that DD estimates were insensitive to the choice of period (Figs. S13 and S14).

DEB model results

To evaluate the performance of our DEB model in simulating sea scallop growth in the MAB, we compared modeled sea scallop shell height, dry tissue weight, wet tissue weight, dry gonad weight, and wet gonad weight with previous statistical results based on laboratory measurements (Langton et al. 1987, Hart and Chute 2009) (Fig. 6). Both the DEB model results and von Bertalanffy growth curve in Hart and Chute (2009) showed that scallop shell height in the MAB reached an asymptote at 133 mm within eight years (Fig. 6a). Modeled dry and wet tissue weights were

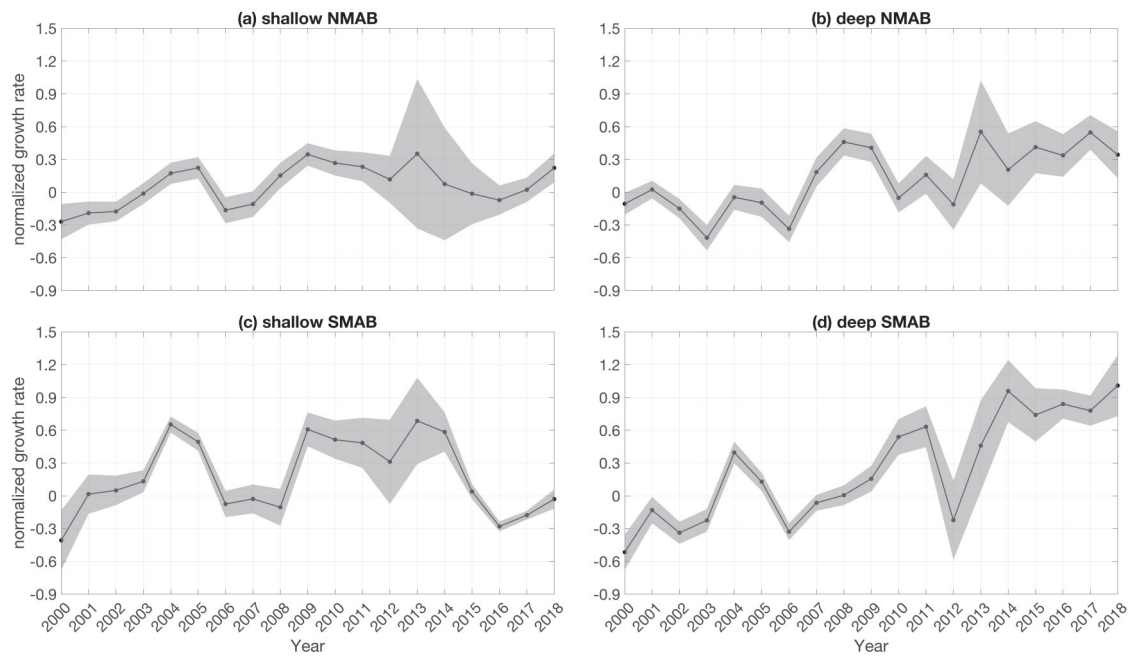


Figure 4 Normalized scallop growth rate interannual variations in four subregions from 2000 to 2018 (panel a: shallow NMAB; panel b: deep NMAB; panel c: shallow SMAB; panel d: deep SMAB). The grey envelopes represent 95% confidence intervals.

Table 1 Normalized scallop growth rate interannual correlations within 14-yr time windows between the shallow and deep MAB subregions (*: $p < 0.05$; **: $p < 0.01$)

	14-yr Time Window					
	2000–2013	2001–2014	2002–2015	2003–2016	2004–2017	2005–2018
NMAB (shallow vs. deep)	0.55*	0.54*	0.49	0.34	0.20	0.25
SMAB (shallow vs. deep)	0.82**	0.76**	0.57*	0.24	0.08	-0.01

comparable to the statistical results of Langton et al. (1987), increasing from approximately 1 to 20 g and 5 to 160 g, respectively (Fig. 6b and c). The seasonal fluctuations of dry and weight tissue weights were driven by reproduction (Fig. 6b and c). Dry and wet gonad weights decreased dramatically during two spawning events per year and recovered gradually after spawning (Fig. 6d and e). The fluctuation magnitude of modeled dry gonad weight increased gradually with age from almost 0 g in year 0 to 3 g after year 8 (minimum dry gonad weight = 0.8 g; maximum dry gonad weight = 3.8 g) (Fig. 6d). Wet gonad weight fluctuated between 7 and 25 g after reaching 8 years of age (Fig. 6e).

Fig. 7 shows the modeled scallop shell height (initial shell height = 40 mm; blue lines) over one year based on bottom temperature climatology (red solid lines) and 2016 bottom temperature (red dashed lines) in the four subregions. Bottom temperature in the MAB exhibited strong seasonality with higher temperature in early fall (October–November) and lower temperature from March to May (Fig. 7). Bottom temperatures in 2016 exceeded the historical climatology by 1–2°C during most of the year (red lines in Fig. 7). In the shallow SMAB, the bottom temperature difference between the climatology and year 2016 reached more than 5°C in early September and October (Fig. 7c). In the shallow subre-

gions, scallop shell height was predicted to grow faster in 2016 than under climatological temperature in the first eight months because temperature in 2016 was closer to the optimum temperature (< 15°C) for scallop growth (Fig. 7a and c). From September to November, bottom temperature exceeding 15°C in 2016 almost halted scallop growth, whereas shell height continued to increase under climatological temperature, resulting in approximately 5 mm greater shell height by the end of the simulation (Fig. 7a and c). In the deep subregions, although bottom temperature in 2016 was overall higher than the climatology, it barely exceeded 15°C and facilitated faster scallop growth (Fig. 7b and d). The modeled scallop shell height in 2016 was approximately 4 mm higher than under the climatology conditions by the end of the one-year simulation (Fig. 7b and d). The sensitivity tests based on 60-, 80-, and 100-mm initial shell heights showed similar spatial patterns, although larger individuals in the deep subregions benefited less from warming (Figs S15–S17). Modeled scallop shell height and dry tissue weight growth curves based on various food limiting factors (section 2.3) suggested that scallop growth was sensitive to food supply (Fig. S18). However, high bottom temperature in 2016 consistently reduced scallop growth in the shallow subregions and enhanced growth in the deep subre-

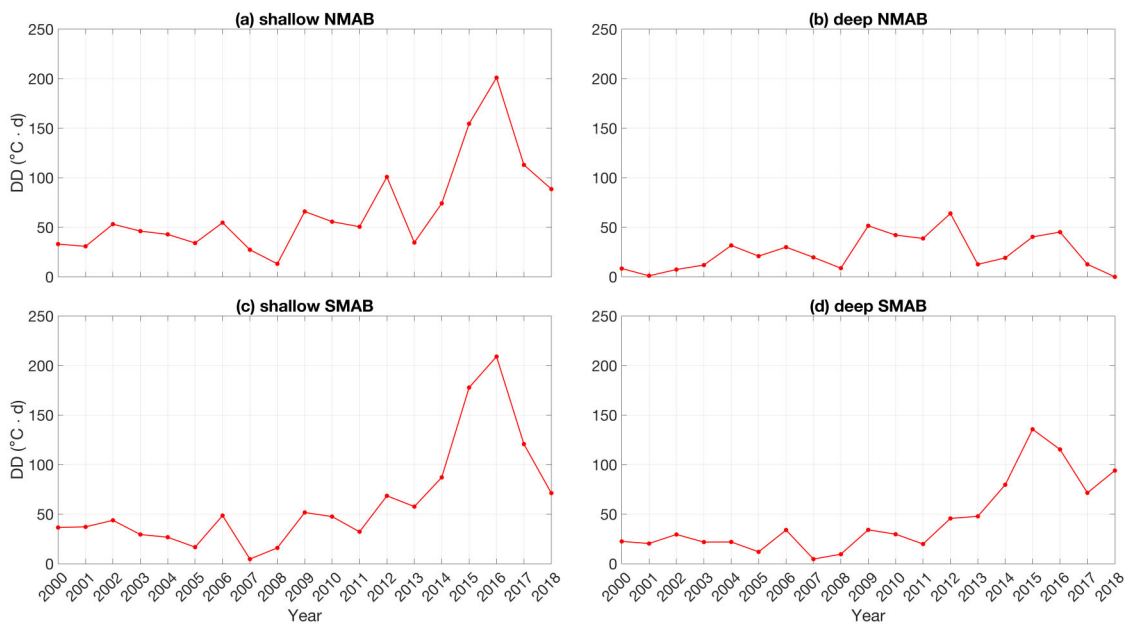


Figure 5 Degree days (DD, unit: °C · d) interannual variations in the four subregions from 2000 to 2018 (panel a: shallow NMAB; panel b: deep NMAB; panel c: shallow SMAB; panel d: deep SMAB).

gions across all tested food conditions, which was consistent with our benchmark run (Figs S19–S22). In general, the DEB model results reflected the same spatial heterogeneity in warming effects on scallop growth as observed in the data, with decreased growth rates in the shallow subregions and increased rates in the deep areas.

To examine the drivers of scallop growth spatial heterogeneity under warming from a bioenergetics perspective, we analyzed three energy fluxes in the model directly related to scallop growth: energy allocated for somatic maintenance and structure growth (κP_C ; Fig. 8), somatic maintenance flux (P_M ; Fig. 9), and allocation flux to growth (P_G ; Fig. 10). κP_C in the shallow subregions increased gradually from 50 J d⁻¹ in January to ~270 J d⁻¹ in September, and its value in 2016 was 10–50 J d⁻¹ higher than the climatological results (Fig. 8a and c). After September, κP_C became stable without a strong difference between the two temperature scenarios (Fig. 8a and c). In the deep subregions, κP_C increased during the entire year, and its value in 2016 was always higher than climatological results, reaching ~400 J d⁻¹ by the end of the simulation (Fig. 8b and d). The fluctuations of κP_C in the MAB were overall positively correlated with bottom temperature, indicating that warming enhanced energy flux for somatic maintenance and structure growth (Fig. 8). The spatiotemporal patterns of P_M in the four subregions were very similar to those of κP_C , and their comparable ranges (κP_C : 50–420 J d⁻¹; P_M : 50–400 J d⁻¹) suggested that a substantial portion of the allocated energy was used for somatic maintenance (Fig. 9). Compared with κP_C and P_M , P_G was considerably lower ranging from 0 to 16 J d⁻¹ (black lines in Fig. 10). P_G increased moderately from 7 J d⁻¹ in January to 15 J d⁻¹ in August when temperature was below 14°C, whereas further temperature increase exceeding 15°C led to the negative correlation between P_G and temperature after August (Fig. 10). In the shallow subregions, P_G in 2016 decreased to zero when bottom temperature was higher than 16°C between September and November, whereas the

climatological P_G during the same period fluctuated around 14 J d⁻¹ (Fig. 10a and c). In the deep subregions, although P_G in 2016 dropped sharply below 6 J d⁻¹ when temperature exceeded 16°C in mid-October, this decline only lasted a few days before recovering to 15 J d⁻¹ (Fig. 10b and d).

Discussion

Defining and quantifying thermal stress on sea scallop growth

The sea scallop is a cold-water bivalve species, and its physiological rates (e.g. feeding, growth, respiration, and mortality), biogeographic distribution, population size structure, and commercial value are very sensitive to temperature variations (MacDonald et al. 1987, Tanaka et al. 2020, Cameron et al. 2022, Zang et al. 2023). Therefore, appropriately defining and quantifying thermal stress intensity is critically important for understanding how thermal conditions are associated with scallop population dynamics. Mean temperature across various temporal scales has been widely applied to examine the impacts of thermal conditions on scallop growth and spatial distribution in natural habitats (Torre et al. 2019, Hodgdon et al. 2020, Tanaka et al. 2020). Although mean temperature can overall represent thermal environments in scallop habitats during a certain period, it smooths out high frequency variability that also plays a critical role in regulating sea scallop physiology. Laboratory experiment results indicated that temperature variation between 6 and 15°C contributed to faster scallop soft tissue (excluding adductor) growth (Pilditch and Grant 1999a). It should be noted that the temperature fluctuation range in Pilditch and Grant (1999a) was determined based on field measurements in Nova Scotia and did not exceed the optimum tem-

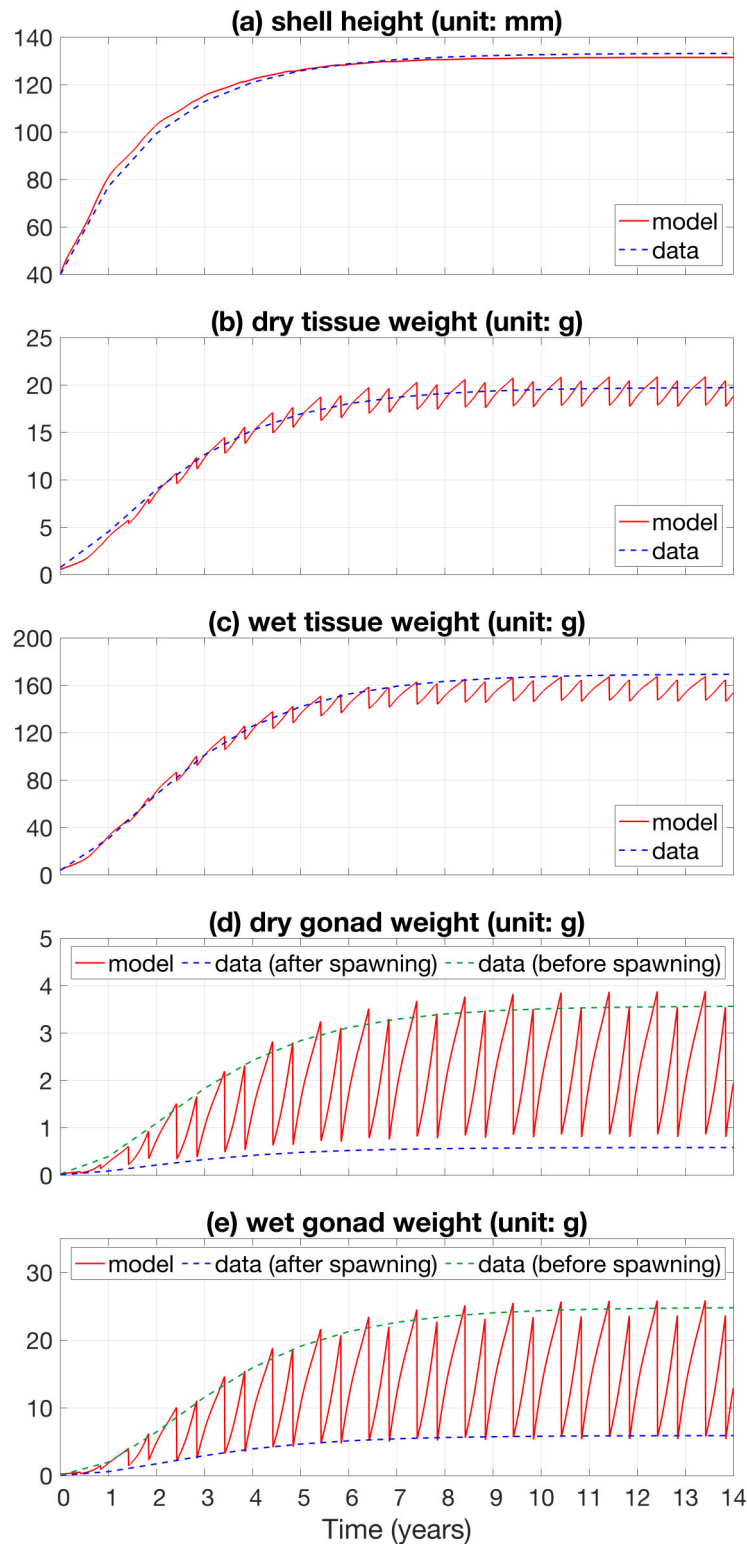


Figure 6 Simulated shell height (SH; a), dry tissue weight (DTW; b), wet tissue weight (WTW; c), dry gonad weight (DGW; d), and wet gonad weight (WGW; e) (red solid lines) in comparison with growth curves based on laboratory-measured data in Hart and Chute (2009) and Langton et al. (1987) (dashed lines). The dashed line in panel a is based on von Bertalanffy growth parameters estimated by Hart and Chute (2009) ($K = 0.508$ and $L_{\infty} = 133.3$ mm). From panels b and e, the measured dry/wet tissue weight and gonad weight are based on \log_{10} - \log_{10} relations in Langton et al. (1987) ($\log_{10}(\text{DTW}) = -4.42 + 2.69 \cdot \log_{10}(\text{SH})$; $\log_{10}(\text{WTW}) = -4.40 + 3.12 \cdot \log_{10}(\text{SH})$; $\log_{10}(\text{DGW}_{\text{before_spawning}}) = -8.01 + 4.03 \cdot \log_{10}(\text{SH})$; $\log_{10}(\text{DGW}_{\text{after_spawning}}) = -7.52 + 3.43 \cdot \log_{10}(\text{SH})$; $\log_{10}(\text{WGW}_{\text{before_spawning}}) = -8.40 + 4.61 \cdot \log_{10}(\text{SH})$; $\log_{10}(\text{WGW}_{\text{after_spawning}}) = -8.13 + 4.19 \cdot \log_{10}(\text{SH})$).

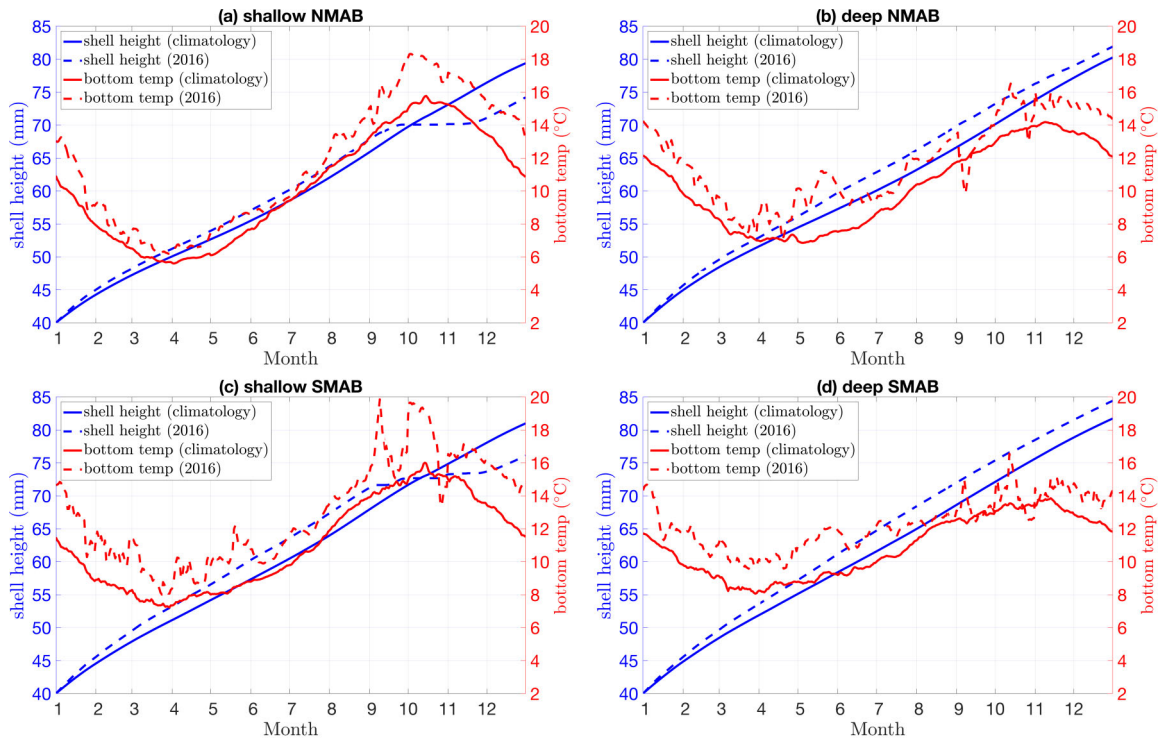


Figure 7 Annual shell height growth (blue) and the corresponding bottom temperature (red) in the four subregions (panel a: shallow NMAB; panel b: deep NMAB; panel c: shallow SMAB; panel d: deep SMAB). The initial scallop shell height is assigned as 40 mm. Solid and dashed lines are the results based on bottom temperature climatology and bottom temperature in 2016, respectively.

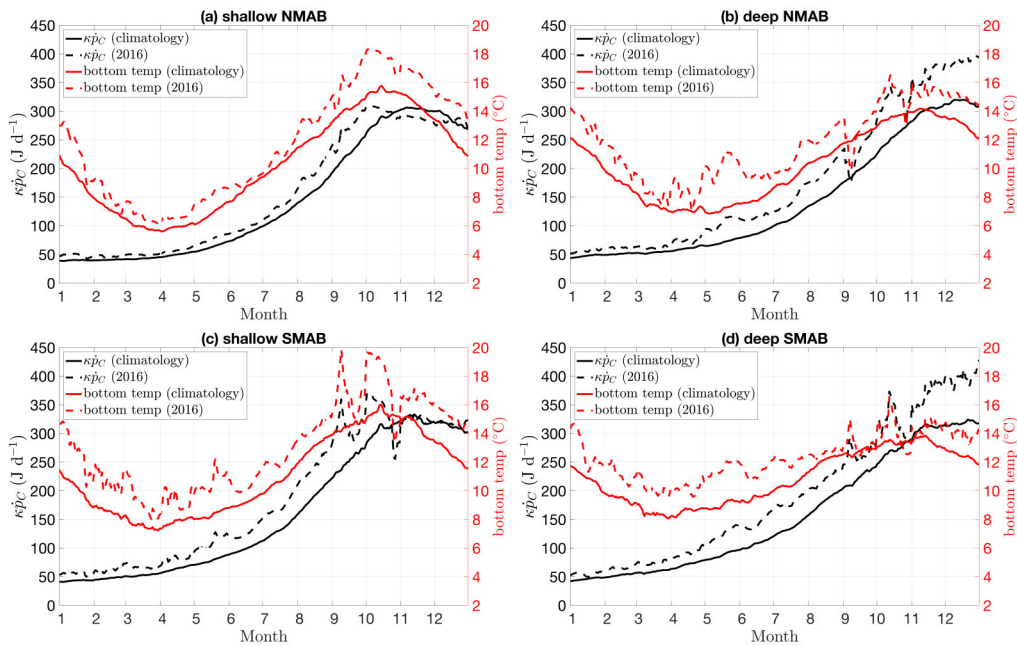


Figure 8 Annual energy flux for somatic maintenance and structure growth ($\kappa \hat{p}_c$; black) and the corresponding bottom temperature (red) in the four subregions. Solid and dashed lines are the results based on bottom temperature climatology and bottom temperature in 2016, respectively.

perature for scallop growth. In the MAB where higher bottom temperatures occur, temperature fluctuations might amplify thermal stress and limit scallop growth due to prolonged exposure to unfavorable thermal conditions (Silina 2023). Local thermal adapta-

tion may influence scallop growth responses to temperature variability. Previous studies have reported distinct genetic clusters between northern populations (Newfoundland and the Gulf of St. Lawrence) and those on Georges Bank and the MAB, which might

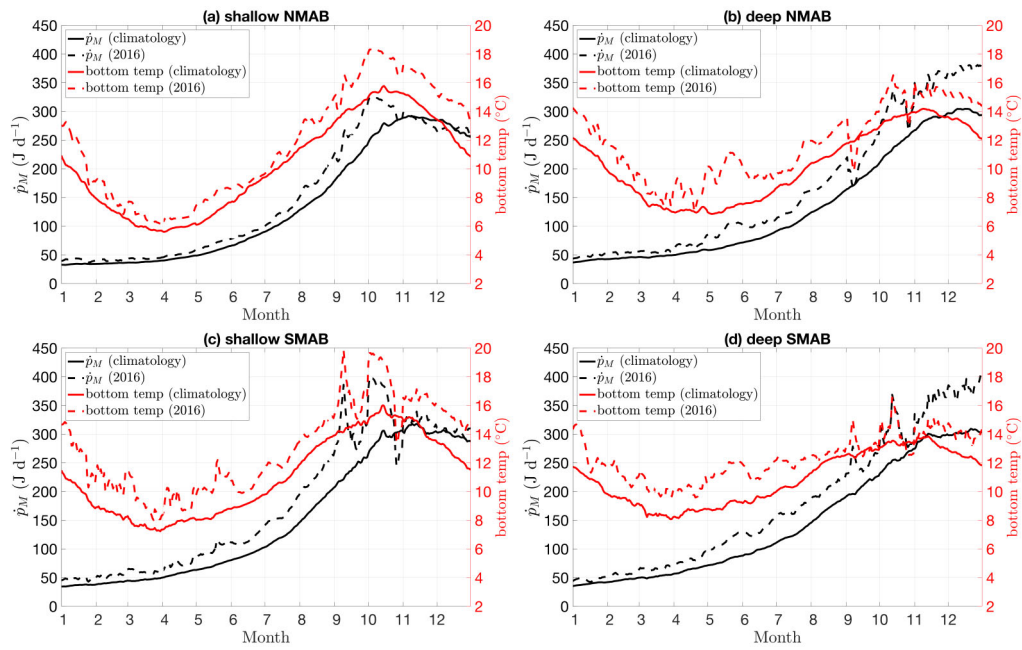


Figure 9 Annual somatic maintenance flux (\dot{p}_M ; black) and the corresponding bottom temperature (red) in the four subregions. Solid and dashed lines are the results based on bottom temperature climatology and bottom temperature in 2016, respectively.

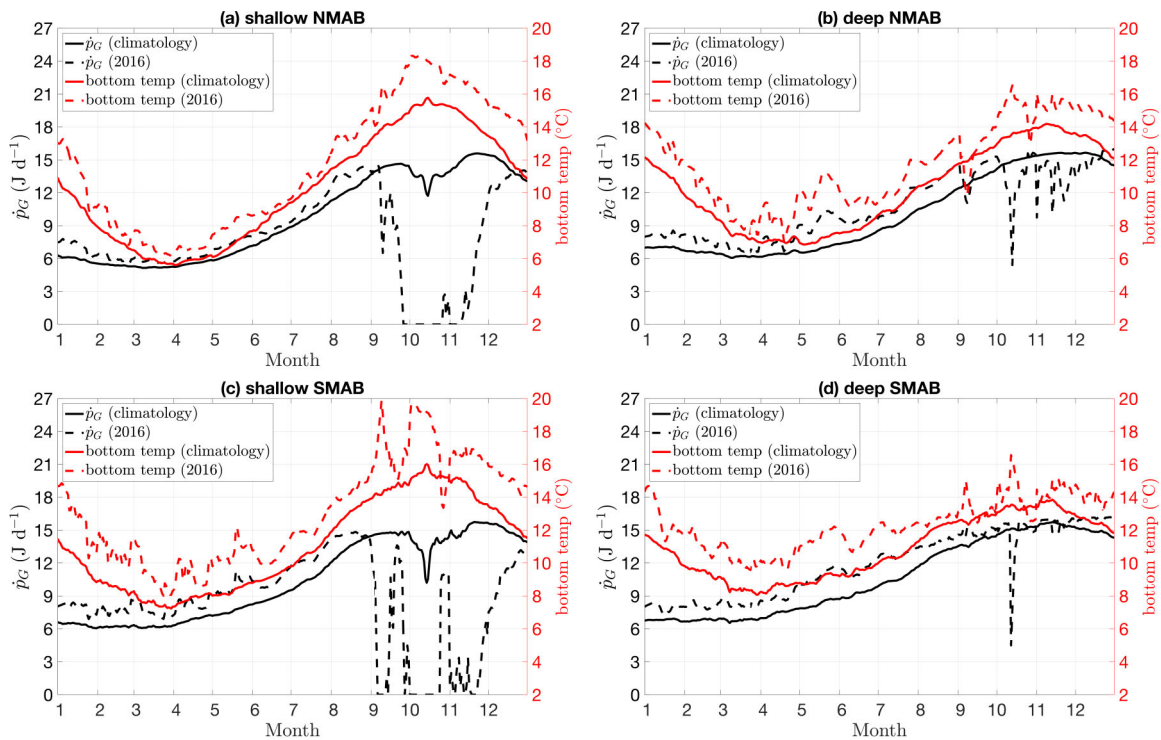


Figure 10 Annual allocation flux to growth (\dot{p}_G ; black) and the corresponding bottom temperature (red) in the four subregions. Solid and dashed lines are the results based on bottom temperature climatology and bottom temperature in 2016, respectively.

be associated with differences in thermal adaptation (Van Wyn-garden et al. 2017; Lehnert et al. 2019; Lowen et al. 2019). Given that adaptive responses have been documented in many bivalve species (Stickle et al. 1989, Pernet et al. 2007), thermal adaptation of scallops in northern habitats may mitigate the negative impacts

of thermal stress to some extent under future climate warming. Additionally, mean temperature fails to capture the phenological characteristics of temperature variability, which is also important in regulating scallop growth. Our DEB model results suggested that warming in the cold season accelerated scallop growth in the

Mills 2000, Robinson et al. 2016, Shumway and Parsons 2016). Increased investments in sea scallop aquaculture research and technology over the last decade have overcome some of the aforementioned challenges and provide a valuable knowledge base for the success of sea scallop aquaculture (Grant et al. 2003, Coleman et al. 2021, 2022, Ishaq et al. 2023). Future efforts to develop adaptive, science-based strategies for aquaculture site selection, cultivation, and operations will enhance scallop productivity and complement wild-capture fisheries to mitigate the impacts of warming on sea scallop industry. Most sea scallop aquaculture activities in the U.S. are concentrated in coastal Massachusetts and Maine, where temperatures are generally lower than in the MAB. Habitat suitability projections based on linear temperature trends suggest that moderate warming may improve thermal conditions for scallops in some nearshore areas of the Gulf of Maine (Tanaka et al. 2020). However, field observations near the mouth of Penobscot Bay show summer temperatures exceeding optimal thresholds for scallop growth at several sites (Coleman et al. 2021). Episodic events such as warm-core ring impingement and their interactions with warm slope water intrusion can further elevate bottom temperatures in the Gulf of Maine, resulting in enhanced thermal stress for cultured scallops (Du et al. 2022). These findings indicate that continued climate warming could pose increasing risks to scallop aquaculture, highlighting the importance of adaptive site selection for scallop aquaculture and long-term temperature monitoring.

Conclusions

We estimated the spatiotemporal patterns of sea scallop growth rate and thermal stress intensity in the MAB from 2000 to 2018. The results suggested that scallop growth rates across all size groups were lower in the deep subregions, with minimal effect of latitude. Scallop growth rates in the entire MAB were generally synchronized until 2014, after which strong thermal stress in 2015 and 2016 coincided with the emerging spatial heterogeneity featured by higher growth rate in the deep subregions and lower growth rate in the shallow areas, respectively. A DEB model was adopted to assess sea scallop growth under 2016 and climatological thermal conditions. The comparisons between model results and observational data indicated the good performance of DEB model in capturing scallop growth dynamics in the MAB. The DEB model results suggested that bottom temperature exceeding 16°C in 2016 profoundly lowered energy allocation to scallop growth in the shallow subregions, while warming in the deep subregions lead to more favorable thermal conditions for scallop growth.

Our findings highlight the importance of appropriately defining and quantifying thermal stress intensity in understanding scallop growth variability across the MAB. Given the projected future warming in many sea scallop habitats on the Northeast U.S. Shelf, improved metrics of thermal stress and consideration of multiple environmental stressors will be important for understanding scallop growth dynamics and informing adaptive management strategies for sustaining sea scallop fishery resources.

Supplementary material

Supplementary material is available at [ICES Journal of Marine Science](https://doi.org/10.1016/j.jms.2026.05.001) online.

Data availability

The sea scallop growth rate data and the DEB model outputs used in this article will be shared upon a reasonable request to the corresponding author.

Author contributions

Z.Z. and R.J. designed the study and performed the statistical analyses. Z.Z. and R.L. developed and calibrated the DEB model. C.C. and S.L. provided physical model setup information and outputs. D.R.H., A.S.C., and R.L.M provided sea scallop growth rate data. R.J. and D.R.H. assisted with the conceptualization of this work. Z.Z., R.J., D.R.H., K.M.K., and R.L. developed the first manuscript draft. All authors reviewed the paper and approved the manuscript submission.

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Conflicts of interest

There is no conflict of interest to state in this research.

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