

Impact of Larval Behaviors on Dispersal and Connectivity of Sea Scallop Larvae over the Northeast U.S. Shelf

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Highlights:

- Larval swimming within the ocean mixed layer affected the interannual variability of scallop larval dispersal and settlement.
- Ignoring larval swimming behavior in the ocean mixed layer likely overestimates the larval connectivity between Georges Bank (GB) and the Middle Atlantic Bight (MAB).
- Climate-induced warming tends to alter the circulation in ways that intensify larval retention over GB and restrict larval transport from GB to the MAB.

37 **Abstract**

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39 Sea scallops (*Placopecten magellanicus*) are a highly fecund species that supports one of
40 the most commercially valuable fisheries in the northeast U.S. continental shelf region.
41 Scallop landings exhibit significant interannual variability, with abundances widely
42 varied due to a combination of anthropogenic and natural factors. By coupling a pelagic-
43 stage Individual-Based scallop population dynamics Model (hereafter referred to as
44 Scallop-IBM) with the Northeast Coastal Ocean Forecast System (NECOFS) and
45 considering the persistent aggregations over Georges Bank (GB)/Great South Channel
46 (GSC) as source beds, we have examined the dispersion and settlement of scallop larvae
47 over 1978-2016. The results demonstrated that the significant interannual variability of
48 larval dispersal was driven by biophysical interactions associated with scallop larval
49 swimming behaviors in their early stages. The duration, frequency, and stimulus of larval
50 vertical migration in the ocean mixed layer (OML) affected the residence time of larvae
51 in the water column over GB. It thus sustained the persistent aggregations of scallops in
52 the GB/GSC and Southern New England region. In addition to larval behavior in the
53 OML, the larval transport to the Middle Atlantic Bight (MAB) was also closely related to
54 the intensity and duration of northeasterly wind in autumn. There was no conspicuous
55 connectivity of scallop larvae between GB/GSC and MAB in the past 39 years except in
56 the autumn of 2009. In 2009, the significant larval transport to the MAB was produced by
57 unusually strong northeasterly winds. Ignoring larval behavior in the OML could
58 overestimate the scallop population's connectivity between GB and the MAB and thus
59 provide an unrealistic prediction of scallop larval recruitment in the region. Both
60 satellite-derived SST and NECOFS show that the northeast U.S. shelf experienced
61 climate change-induced warming. The extreme warming at the shelfbreak off GB tends to
62 intensify the cross-isobath water temperature gradient and enhance the clockwise subtidal
63 gyre over GB. This change can increase the larval retention rate over GB/GSC,
64 facilitating enhanced productivity on GB.

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1. Introduction

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Sea scallops (*Placopecten magellanicus*), which occur on the northeast continental shelf of North America, support the most valuable wild scallop fishery in the world (Shumway and Parsons, 2016). Georges Bank (GB) is one of two areas with the highest scallop abundances in the Northwest Atlantic (Stokesbury *et al.*, 2004; Hart and Rago, 2006; NFSC, 2018) (Fig.1). Based on drop-camera surveys with a coverage area of 27×10^3 km² over the period 2016-2018, Stokesbury and Bethoney (2020) estimated the scallop population over the northeast shelf, accounting for ~34 billion individual scallops, ~71% of which were on GB. Over GB, the scallop landings exhibited considerable interannual variability, with an annual value of hundreds of million dollars (Naidu and Robert, 2006; NFSC, 2018). Benefiting from the implementation of closed areas as well as fishing effort and gear restrictions, U.S. sea scallop stocks rapidly recovered from a period of severe overfishing during the 1990s (Murawski *et al.*, 2000; Hart and Rago, 2006; Hart *et al.*, 2013; Davies *et al.*, 2015; NFSC, 2018). However, even in light of the recovery, sea scallop abundances have varied significantly, largely due to high recruitment variability affected by a combination of anthropogenic and natural factors (Hart and Rago 2006; NFSC, 2018).

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Recruitment, which is estimated by the survivorship of scallop larvae in their early life stages, is crucial in determining the population size. The early scallop life stages consist of pelagic and benthic phases. Adult scallops spawn eggs near the bottom. After external fertilization, trochophores hatch within 1-2 days, develop small cilia a few hours after hatching, and then start to migrate upward towards the sea surface (McGarvey *et al.*, 1992; Hart and Chute, 2004; Cragg, 2006). Once arriving at the sea surface, they undergo vertical migrations within the surface oceanic mixed layer (OML) (Tremblay and Sinclair, 1990a, 1990b; Gallager *et al.*, 1996). The veliger stage is reached over 4-5 days with the development of shell velum (Silva-Serra, 1995; Pearce *et al.*, 2004). At the ages of 30-35 days, veligers develop into pediveligers with foot and byssus development (Stewart and Arnold, 1994). Pediveligers can actively swim across the thermocline and descend towards the bottom for settlement (Tremblay *et al.*, 1994). During this pelagic

100 phase, changes in the flow-driven larval dispersal and retention are primary factors in
101 controlling interannual variability in spatfall and abundance (*McGarvey et al.* 1993).
102 After settlement, the survivorship of spat (settled larvae) and juveniles crucially
103 influences the adult sea scallop population size and distribution (*Caddy, 1975; Hart and*
104 *Chute, 2004*). During this benthic phase, the substrate motility, water temperature,
105 currents/storms, predation, and starvation can affect the survivorship of newly settled spat
106 and juveniles (*Merrill and Edwards 1976; Larsen and Lee 1978, Hart 2006, Shank et al.*
107 *2012*).

108 The interannual variability of scallop abundance and recruitment on GB/GSC is
109 influenced considerably by changes in both physical and biological processes (*Hart and*
110 *Chute, 2004*). Understanding the driving mechanisms of these variabilities and their
111 connectivity with the Middle Atlantic Bight (MAB) can provide insights into the
112 biophysical reasons for persistently high scallop abundance over GB/GSC and primary
113 factors attributing to abundance reductions. It can also scientifically guide the
114 management of rotationally closed areas, optimal seeding of sea scallops, and protection
115 of seeded sea scallop's settling regions. It is a significant challenge to predict
116 environment-driven variability in the GB/GSC scallop population. The environmental
117 factors reflect the complex nonlinear physical-biological interaction processes, such as
118 global warming, climate-induced shelf-basin scale interactions, local wind/tidal mixing,
119 ocean acidification, ecosystem regime shift, and prey/predator fields, etc. (*Hart and Rago,*
120 *2006; Shank et al. 2012; Stokesbury et al., 2016; Rheuban et al., 2018*).

121 The sea scallop fishery in the U.S. Northeast is currently managed using fishing effort
122 limitations combined with rotational closures (*Hart and Rago 2006*). Areas are closed
123 based on observations of strong recruitment from surveys, and then reopened to fishing
124 after the scallops have grown to more optimal sizes for harvesting. There have been a few
125 modeling studies carried out to assess the marine environmental impact on recruitment
126 processes (reproduction, the timing of spawning, pre and post-settling larval stages) on
127 GB/GSC (*Tian et al., 2009a, 2009b, 2009c; Gilbert et al., 2010; Davies et al., 2014,*
128 *2015*) and in the MAB (*Munroe et al., 2018, Hart et al., 2020*). *Tian et al. (2009a)*
129 developed a scallop population individual-based model (hereafter referred to as Scallop-
130 IBM). The model was coupled with the unstructured grid, Finite-Volume, Community

131 Ocean Model (FVCOM) for the Gulf of Maine (GoM) (hereafter referred to as GoM-
132 FVCOM) (*Tian et al.* 2009a, 2009b, 2009c). Spawning on GB in autumn, they ran this
133 coupled Scallop-IBM/GoM-FVCOM model for 1995-2005. The dispersal of simulated
134 scallop larvae varied interannually, with significant transport to the MAB (*Tian et al.*,
135 2009c). Driving a simplified passive and pycnocline-seeking, temperature-dependent,
136 scallop larval transport model by FVCOM-simulated monthly climatological flow and
137 temperature fields, *Gilbert et al.* (2010) examined the influences of flow-driven retention
138 and larval vertical migration on the larval dispersion in the GB/GSC region for both fall
139 and spring spawning seasons. They found that pycnocline-seeking behavior could alter
140 the larval dispersal by factors of 2-5, and thermal history could significantly affect the
141 planktonic larval duration.

142 The flow and temperature fields used in previous scallop larval transport simulations
143 (e.g., *Tian et al.*, 2009a, 2009b, 2009c; *Gilbert et al.*, 2010) were from the first-
144 generation GoM-FVCOM for the region, which did not consider the physical processes
145 relating to regional-scale climate forcing. Specifically, the GoM-FVCOM hydrodynamics
146 missed two remote boundary conditions: 1) the advective transport from the upstream
147 Labrador Sea and the Arctic Ocean, and 2) the Gulf Stream-shelf interactions along the
148 southeastern part of the domain (Fig. 1). Regarding the population dynamics, although
149 Scallop-IBM included the pre-settling pycnocline-seeking behaviors of scallop larvae,
150 age-at-size-specific pre- and post-settling swimming within the OML or near the bottom
151 were not taken into account (*Stewart and Arnold*, 1994; *Gallager*, 1996; *Gallager et al.*,
152 1986a, 1986b, 1996). Additionally, the spawning distribution for the 1995-2005
153 simulations was based only on a scallop dataset produced by video surveys from the
154 University of Massachusetts/School for Marine Science and Technology (UMASS-D/
155 SMAST) (*Stokesbury et al.*, 2004). This dataset does not contain the data from either the
156 Canadian waters over the eastern flank of GB or NOAA surveys conducted
157 independently every year with records back to 1979. The larval behaviors and spatial
158 distributions of spawning are known to have a significant role in the bulk transport of
159 larvae (*Gilbert et al.* 2010). It is necessary to conduct an in-depth analysis of the
160 responses of dispersal patterns to different behaviors by using a model initialed with
161 complete coverage of spawning locations from all available scallop data.

162 High levels of adult biomass on GB/GCS, including the closed areas over Nantucket
163 Lightship Closed Area (NLCA), Closed Area I (CA-I), Closed Area II (CA-II), and
164 Habitat Area of Particular Concern (HAPC) in the northern part of CA-II are well
165 established (*Hart and Rago* 2006; *Hart et al.* 2013; *Stokesbury et al.*, 2015; *Gallager*,
166 2016). For data mining, we collected the scallop abundance data from NOAA, Canadian,
167 and SMAST surveys, and expanded the database to cover a period from 1979 to 2017.
168 For model development, we, a joint research team at UMASS-D and Woods Hole
169 Oceanographic Institution (WHOI), developed the Northeast Coastal Ocean Forecast
170 System (NECOFS). The 39-year (1978-2016) hindcast simulation of NECOFS was
171 conducted using a global-regional nested FVCOM system, which improved the numerical
172 simulation of the regional circulation by including the Gulf Stream-shelf interaction and
173 flows from the upstream Labrador Sea and the Arctic Ocean. The availability of a
174 complete scallop abundance dataset and 39-year NECOFS hydrodynamic fields allows us
175 to re-examine the influences of physical processes and scallop larval behaviors on the
176 early life stages of scallop larvae in the region. In particular, how do the Gulf Stream-
177 shelf interaction and flows from the upstream Labrador Sea and the Arctic Ocean
178 influence the transport of larval in GB/MAB in the context of realistic larval motility?
179 How do these factors change the population connectivity between GB, Southern New
180 England (SNE) shelf, and the MAB compared to previous estimates? Does the short-term
181 vertical migration affect the dispersal and settlement of scallop larvae in their early life
182 stages? What is the relative importance of these physical and biological factors for
183 understanding and predicting changes due to dispersal and retention? Ultimately, could a
184 coupled physical and individual-based fishery model reproduce and predict biophysical
185 processes in terms of interannual variability and future management implications?

186 In this research, we have upgraded the Scallop-IBM with improvements of larval
187 behavior parameterizations in the pre-settling stage and coupled it with the third version
188 of GoM-FVCOM of NECOFS (hereafter referred to as GoM3-FVCOM). Using this
189 upgraded coupled model, we examined the dispersal and settlement of scallop larvae with
190 eggs spawning on GB over 39 years from 1978 to 2016. The NECOFS-produced hourly
191 physical fields include the Gulf Stream-shelf interaction and the upstream flows from the
192 Labrador Sea and the Arctic Ocean. The simulation aimed to assess the impacts of

193 various migrating larval behaviors within the surface OML on the scallop larvae's
194 dispersal and settlement in their early life stages.

195 The remaining sections are organized as follows. Section 2 describes the data and the
196 model. Section 3 presents the results of model simulations, including the discussion on
197 the sensitivity of larval dispersal and retention to larval behaviors in constant and varying
198 OMLs and the scallop population's connectivity between GB/GSC, SNE, and MAB.
199 Section 4 highlights the biological and physical processes affecting the interannual
200 variability of larval dispersal. Finally, section 5 summarizes the findings with
201 conclusions.

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2. The Coupled NECOFS-Scallop-IBM Model and Data

204 2.1. NECOFS

205 NECOFS is an integrated atmosphere, surface wave, and ocean forecast model
206 system designed for the U.S. northeast coastal region. For the NECOFS version used in
207 this study, the computational domain covers the continental shelf with boundaries over
208 the northern coast of Chesapeake Bay on the south and the Scotian Shelf on the north,
209 including a portion of the MAB (Fig. 2). NECOFS was placed in experimental 24/7
210 forecast operations in late 2007. The present version of NECOFS includes 1) a
211 community mesoscale meteorological model named "Weather Research and Forecasting
212 (WRF-AWR)"; 2) the regional ocean model of FVCOM (GoM3-FVCOM) (*Chen et al.*
213 2003); 3) the unstructured-grid surface wave model (FVCOM-SWAVE) with the same
214 domain as GoM-FVCOM (*Qi et al.*, 2009); 4) the Mass Coastal FVCOM with the
215 inclusion of estuaries, inlets, harbors, and intertidal wetlands; and 5) four subdomain
216 coupled wave-current FVCOM inundation forecast systems in Scituate, MA; Boston
217 Harbor, MA; Hampton-Seabrook Estuary, NH, and Saco Bay, ME. The GoM3-FVCOM
218 grid covers the scallop aggregation areas over GB/GSC, SNE, and the MAB. The grid is
219 constructed using unstructured triangular meshes with a resolution of ~ 0.3-25 km in the
220 horizontal and 45 layers in the vertical.

221 The 39-year (1978-2016) hindcast simulations of NECOFS were conducted using a
222 global-regional nested FVCOM system with the core models of Global-FVCOM and
223 GoM3-FVCOM (Fig. 2). Global-FVCOM is a fully coupled atmosphere-ice-wave-ocean,

224 unstructured-grid primitive equation global ocean model with a horizontal resolution
225 varying from ~2 km within the Canadian Archipelago, shelfbreak, and coastal region to
226 ~50 km in the interior open ocean. This model was driven by *a*) astronomical tidal
227 forcing with eight constituents (M_2 , S_2 , N_2 , K_2 , K_1 , P_1 , O_1 , and Q_1), *b*) surface wind stress,
228 *c*) net heat flux at the surface plus shortwave irradiance in the water column, *d*) surface
229 air pressure gradients, *e*) precipitation (P) minus evaporation (E), and *f*) river discharges
230 (*Chen et al.*, 2016; *Zhang et al.*, 2016a, 2016b). A 39-year NECOFS hourly hindcast
231 product is now available on the NECOFS Web Map Server
232 (<http://porpoise1.smast.umassd.edu:8080/fvcomwms/>). This database includes
233 meteorological and oceanic components. The meteorological database includes hourly
234 fields of physical variables such as wind velocity, air pressure, precipitation minus
235 evaporation, shortwave radiation, longwave radiation, sensible and latent heat fluxes, and
236 air temperature, etc. The oceanic database contains hourly fields of three-dimensional
237 water currents, temperatures, salinity, horizontal/ vertical turbulent diffusion rates, and
238 surface elevation.

239 The NECOFS-simulated physical fields were validated through comparisons with
240 available observations. It has demonstrated success in capturing tidal- and shelfbreak
241 density fronts, residual clockwise gyres, wind-driven upwelling, buoyancy-driven river
242 plume, the Gulf Stream-shelf interaction (*e.g.*, warm-core rings), and volume and mass
243 transports entering the Gulf of Maine over the Nova Scotia shelf from the upstream
244 Labrador Sea or even the Arctic Ocean. The model-data comparisons included 1) water
245 elevations at tidal gauges (*Chen et al.*, 2011, *Sun et al.*, 2013), 2) temperature and salinity
246 in the water column (*Li et al.*, 2015), 3) hurricane and extratropical storms (*Chen et al.*,
247 2013, *Beardsley et al.*, 2013), 4) the surface currents measured by CODAR from 2000 to
248 2008 (*Sun et al.*, 2016), and 5) upstream conditions in the Arctic Ocean (*Chen et al.*,
249 2009; *Chen et al.*, 2016; *Zhang et al.*, 2016a,b). The success of scallop-IBM depends on
250 the accuracy and reality of the flow fields predicted by the physical model. We have
251 conducted a model-drifter comparison to validate the reliability of the FVCOM-produced
252 flow field over 1995-2013. Six hundred eighty-four drifters were deployed in the GoM
253 and GB regions, which returned valuable trajectory data (J. Manning, personnel
254 communication). A non-parametric Kolmogorov-Smirnov test was used to judge “good”

255 and “bad” comparisons (*Van Sebille et al., 2009*). The results showed that 75% of drifters
256 were in fair comparison with the model-predicted drifter trajectories (*Sun, 2014*). These
257 validation experiments provide us with confidence in using the NECOFS-produced flow
258 field to study the impact of physical processes on the interannual variability of sea scallop
259 recruitment over GB/GSC, SNE, and MAB.

260 2.2. Scallop-IBM

261 The model used in this study is an upgraded Scallop-IBM coupled with the GoM3-
262 FVCOM model. Scallop-IBM consists of four phases: egg, trochophore, veliger, and
263 pediveliger (Fig. 3). Ages defined individual development in each stage: eggs <2 days,
264 trochophores 2–4 days, veligers 5–40 days, and pediveligers > 40 days (*Stewart and*
265 *Arnold, 1994*). We used fixed development times on pelagic stages under the assumption
266 that the relatively small interannual changes in water temperature would produce
267 insignificant modulation in larval development times. Similarly, the food limitation was
268 not considered for larvae since that food was abundant during the pelagic stages.

269 Modeled larval behavior and their vertical migrations were considered for each life
270 stage based on our empirical understanding. Eggs are spawned on the seabed, neutrally
271 buoyant, and drift passively via vertical currents and turbulence but without vertical
272 migration (*Culliney, 1974; Silva and O’Dor, 1988; Tremblay, 1988; Tremblay et al.,*
273 *1994*). Trochophores have no directionality in their swimming and only randomly spin
274 (*Tian et al., 2009a*), and so were also treated passively. Laboratory experiments have
275 found that once the first shell formed (*prodisoconch*) and the larvae appear in a ‘D’
276 configuration, their gravity centers are below the velum, causing them to swim upwards
277 across the thermocline (*Gallager, 1993; Gallager et al., 1996*). Veligers are subject to
278 horizontal drift in the surface OML above the thermocline, in which they actively
279 switched between upward swimming and sinking to produce a distinct vertical migration
280 pattern. Veligers are sensitive to light transitions, not to any prolonged state of light
281 intensity like day or night (*Gallager et al., 1996*). Larvae between the ages of 5 and 40
282 days vertically migrate within the OML with various patterns such as thermocline-
283 seeking aggregation (*Tremblay and Sinclair, 1990a*), diel (*Tremblay and Sinclair, 1990b*),
284 and semidiurnal cells (*Gallager et al., 1996; Manuel et al., 1996*). *Tremblay and Sinclair*
285 (1990b) used a pump to make profile samplings of scallop larval abundance at eight

286 stations on GB in October 1986 and 1987, respectively. Four of the stations were located
 287 in the stratified region. They observed an aggregation of bivalve scallop larvae in the
 288 thermocline at a depth of the subsurface chlorophyll maximum. In laboratory mesocosm
 289 experiments, over a diel cycle, veligers stayed near the surface at night, moved down,
 290 and remained at the thermocline during the day (Manuel et al., 1996) (Fig. 4). Over
 291 semidiurnal migration cycles, they stayed near the surface when daybreak, moved to the
 292 thermocline around noon, came up towards the surface at sunset, and were back to the
 293 thermocline around mid-night, forming bio-convective cells within the OML after dark
 294 (Manuel et al. 1996) (Fig.4). Larvae also respond to turbulence's ephemeral pulses
 295 greater than 10^{-7} W.Kg⁻¹ by withdrawing their velum and sinking rapidly until the
 296 turbulent energy has subsided (Pearce et al., 1998). The currents in the GB/GSC region
 297 are dominated by the semidiurnal M₂ tidal currents. During the autumn, the thermocline
 298 varied significantly due to winds. The flow differed at the surface and thermoclines so
 299 that migration behaviors influenced larval retention. However, these extensive suites of
 300 swimming behaviors have never been captured in a model to date. In the past, the larvae
 301 were treated as particles with a random walk (e.g., Stewart and Arnold, 1994; Tian et al.,
 302 2009a) or simple thermocline seeking behavior (Gilbert et al., 2010; Davies et al., 2014,
 303 2015; Munroe et al. 2018). Swimming behaviors could contribute significantly to the
 304 overall larval transport potential since they are always responding to the stimuli by
 305 changing their depth (Gallager et al., 1996). Late-stage pediveligers (>40 days) migrate
 306 downwards to settle on the seabed (1.7 mm s⁻¹), but may remain at the thermocline for
 307 more than 100 days and delay metamorphosis if thermal conditions are not suitable
 308 (Pearce et al., 1996). Such a delay in the settlement could lead to higher retention if
 309 larvae are in a gyre circulation. Mortality throughout the pelagic phase is carefully
 310 parameterized based on data and conditions provided in the literature (e.g., Gallager et
 311 al., 1986a,b, 1988; McGarvey et al. 1992).

312 The Scallop-IBM consists of a super-individual tracking equation given as

$$313 \quad P_i(\vec{x}_{n+1}, t_{n+1}) = P_i(\vec{x}_n, t_n) + \int_{t_n}^{t_{n+1}} \vec{v}(\vec{x}, t) dt + W_b(x, y, z, t) \Delta t + R_H + R_K \quad (1)$$

314 where $P_i(\vec{x}, t)$ is the egg or larval number in the i th super-individual at the location $\vec{x} =$
 315 $x\vec{i} + y\vec{j} + z\vec{k}$ at the time t ; x , y , and z are the east, north and vertical axes of the

316 Cartesian coordinates; \vec{i} , \vec{j} , and \vec{k} are unit vectors in x, y and z directions; subscript n
 317 represents the n th time step; \vec{v} is the three-dimensional velocity vector; Δ is the time
 318 step equaling $t_{n+1} - t_n$; W_b is the vertical migration speed due to larval behavior; R_H
 319 and R_K are the horizontal and vertical random walks as functions of model-produced
 320 horizontal and vertical diffusion coefficients. The formulations of R_H and R_K were
 321 described in *Tian et al.* (2009c). Eq. (1) is solved by the 4th-order, 4-stage explicit Runge-
 322 Kutta (ERK) method with the detail given in the FVCOM User Manual (*Chen et al.*,
 323 2013). The time step used in larval tracking was 120 sec, with the random walk time step
 324 of 6 sec.

325 The super-individual approach is commonly used in larval transport studies (*Scheffer*
 326 *et al.*, 1995; *Bartsch and Coombs*, 2004; *Woods*, 2005; *Tian et al.*, 2009a), which has a
 327 similar meaning as the simulated larvae defined in *North et al.* (2008). A super-individual
 328 was defined as an ensemble particle containing a total of 1.0×10^8 individual eggs. In the
 329 Scallop-IMB, the spawning undergoes two phases before and after larval release (*Tian et*
 330 *al.* 2009c), and the larval numbers in each super-individual are given as

$$331 \quad P_i(\vec{x}, t) = \begin{cases} N_s E_s \int_{t_o}^t \frac{1}{\sqrt{2\pi}\sigma} e^{-\frac{1}{2}(\frac{t-t_m}{\sigma})^2} d & \text{Spawning period} \\ P_i(n, t - \Delta) e^{-Mt} & \text{Release period} \end{cases} \quad (2)$$

332 where N_s is the total adult scallops in a spawning cell at \vec{x} ; E_s is the total eggs spawned
 333 by an individual scallop; t_o is the initial time at which the i th super-individual forms; t_m is
 334 the maximum spawning time; σ is the standard deviation; Δ is the numerical integration
 335 time step. M is the instantaneous mortality rate given as a constant of 0.25 d^{-1} . This
 336 constant number was adopted from *McGarvey et al.* (1992) and *Tian et al.* (2009c). A
 337 super-individual formed as total spawned eggs reached 1.0×10^8 . The super-individual
 338 approach helps us reduce the requirement for a computer's memory to handle a large
 339 number of particles.

340 2.3. Data

341 We obtained the sea scallop biomass and distribution data in the study region over
 342 1979-2017. The data were from three sources: 1) SMAST/UMASSD, 2) U.S. NOAA,
 343 and 3) Bedford Institution of Oceanography (BIO). The SMAST/UMASSD drop camera
 344 data covered 2003-2017, NOAA dredge survey data covered 1979-2017, and BIO dredge
 345 survey data covered 2003-2017. The BIO data covered the survey areas on the eastern

346 flank of GB in Canadian waters. We received these data from the Bedford Institute of
347 Oceanography (BIO), Population Ecology Division (PED), Department of Fisheries and
348 Oceans (DFO), Canada.

349 2.4. Design of numerical experiments

350 We have conducted a set of the coupled scallop-IBM/NECOFS model experiments to
351 examine 1) how sensitive the dispersal and settlement of scallop larvae are to the
352 parameterizations of scallop larval behavior in the early stages, 2) how the interannual
353 variability of the subtidal circulation can influence the settlement of scallop larvae, and 3)
354 what are the physical processes affecting the larval connectivity between GB/GSC and
355 MAB. The simulation covered the period 1978-2016. Physical variables and parameters
356 include the flow-induced advection, water temperature, mixing intensity, and OML depth.
357 To distinguish the physical and biological impacts, we drove the Scallop-IBM by
358 spawning based on the multiyear-averaged abundance and distribution of adult sea
359 scallops over 1979-2017 (Fig. 5). The scallop data used to create the multiyear-averaged
360 field included video and dredge surveys from SMAST/UMASSD, NOAA, and
361 BIO/Canada. Different efficiency estimates were made for video and dredge data.

362 Adult sea scallops spawn in the spring and fall seasons, with the dominant spawning
363 in the autumn (*Posgay and Norman, 1958*). Here we only consider the fall spawning
364 season. Following the previous approach used in *Tian et al. (2009a)*, in each year, we
365 specified the scallop spawning to satisfy a normal distribution starting at 00:00 GMT,
366 September 1 and ending at 24:00 GMT, October 10 (Fig. 6). Peak spawning was set on
367 September 20, with a 1-week standard deviation. The major spawning, which accounted
368 for an amount of 95% of the total spawning, was completed over four weeks, a spawning
369 time range observed in the field measurements (*Posgay and Norman, 1958; Posgay,*
370 *1976; Mullen and Morning, 1986; DiBacco et al., 1995*).

371 The simulation was repeated yearly. Each year, Scallop-IBM was integrated over
372 three months from September 1 to November 30, considering a time scale of ~40 days for
373 larval settlement. Two types of experiments were made (hereafter referred to as "Exp-I
374 and Exp-II"). For Exp-I, the model parameters were the same as those used in *Tian et al.*
375 *(2009a)*. Active vertical migration was specified for each life stage. At the age of 2 days,
376 the larvae started migrating upward towards the surface at a speed of 0.3 mm/s. At the

377 age of 5 days or later, the rate of upward larval migration was decreased to 0.1 mm/s. At
378 the age of 40 days, veligers developed into pediveligers, which actively migrated
379 downwards to the seabed at a speed of 1.7 mm/s and settled on a suitable substrate. For
380 Exp-II, in addition to the parameters considered in Exp-I, we included the vertical
381 migration of scallop larvae during early stages within the surface OML following the
382 schematic patterns shown in Fig. 4. Once larvae entered the OML, the upward larval
383 migration speed was replaced by larval vertical migration behaviors specified in the OML
384 in all Exp-II cases. During the spawning period in September, the water was generally
385 well mixed in the shallow regions (< 40 m) over GB and stratified in the deeper water
386 between tidal mixing and shelfbreak fronts (~40-100 m) on the southern flank of GB.
387 During that period, the wind-induced surface OML could deepen to ~20-40 m in the
388 stratified region. We included a vertical larval migration in the model to examine how
389 this type of larval behavior may affect larval settlement after 40 days.

390 The numerical experiments were done for eight cases (Table 1). C#1 is defined as the
391 case for Exp-I in which vertical migrations in the OML were not included. Exp-II was
392 made for seven cases. C#2, C#3, C#4, and C#5 are defined as the cases with diel or
393 semidiurnal vertical migration behavior in a fixed 10 or 30-m depth OML, respectively.
394 C#6 and C#7 refer to the cases with diel and semidiurnal vertical migration behaviors in
395 the physical model's predicted, spatiotemporally-varying OML. We also did an
396 experiment by constraining larvae at the bottom of the model-predicted OML after they
397 migrated upward to the surface at the age of 5 days, and referred it to as a "thermocline-
398 seeking behavior" case (C#8). For C#6, C#7, and C#8, the hourly OML depth was
399 determined by vertical profiles of the model-simulated water density through an
400 empirical method described in Appendix A. The calculated OML depth was validated via
401 modeled temperature, salinity, and density profiles, with examples shown in Figs. A1-A4.

402
403

3. Influences of the Surface OML on Larval Dispersal

3.1. Comparisons between the cases with and without constant thickness OMLs

404 The results indicate that the dispersal and settlement of scallop larvae varied
405 significantly with scallop larval behaviors in their early stages and the thickness of the
406 OML. It is elucidated from the abundance distributions of pediveliger settling at the
407 seabed for the cases with and without diel or semidiurnal migration (C#1, C#2, C#3, C#4,
408

409 and C#5). Examples are displayed here for 2008, 2009, 2012, and 2013 simulated
410 numbers and concentrations of settled super-individual particle/larvae (Figs. 7-10).
411 During the autumn of these four years, the top of GB and in other shallow regions was
412 vertically well-mixed by tides. The OML depth in the mixed areas was equal to the local
413 water depth. In the following discussion, the positive and negative signs of the flow and
414 transport referred to x - and y -directions in rotated figures (e.g., Figs. 7-10: lower panels).

415 In 2008, for C#1, the scallop larvae were all retained on GB and the SNE shelf, with
416 about 49.1. and 50.9% settling in these two areas, respectively. The larvae were most
417 abundant on the eastern side of GSC and the northeast flank of GB as well as inside the
418 cold pool area (Fig.7f). The cold pool is a relatively uniform cold water body ($< 13^{\circ}$ C)
419 near the bottom that persists from spring through fall over the mid and outer shelf regions
420 (Lentz, 2017). For C#2 and C#3, for a specified 10-m OML, the diel or semidiurnal larval
421 migration in the OML strengthened the larval retention within the clockwise residual gyre,
422 resulting in 75.8 and 80.5% settling on GB/GSC, respectively (Figs. 7g, 7h). Although
423 the difference in larval retention rates on GB/GSC for these two cases was only $\sim 4.7\%$,
424 the spatial distributions of settled larvae differed considerably. For C#2, highly abundant
425 larvae were settled on the western GB and within the GSC and the cold pool areas over
426 the Nantucket Shoal. For C#3, in addition to these three areas, a large portion of larvae
427 was settled down on the northern flank of GB. Without considering vertical migrations in
428 the OML, many larvae were advected southward within the cold pool to the SNE shelf,
429 with a southmost boundary off Long Island. When vertical migrations in the OML are
430 taken into account, the larvae entering the SNE significantly reduced, accounting for
431 $\sim 24.2\%$ for the diel migration case and 19.5% for the semidiurnal migration case. In both
432 cases, a relatively high abundance zone shifted northward and even entered the Long
433 Island Sound.

434 When the OML was deepened to 30 m, the distributions of settled larvae significantly
435 changed (Figs. 7i, 7j). The larvae tended to settle within tidal mixing and shelfbreak front
436 zones. Although the settled larval number remained high around the clockwise gyre over
437 GB, the highest larval abundance concentrated around the western and eastern shelves of
438 GSC. The settled larval number reduced to 56.2% and 71.5% on GB/GSC and increased
439 to 43.8% and 28.5% over the SNE shelf for C#4 and C#5, respectively. The OML

440 deepening enhanced the larval retention around the GSC, and restricted the southward
441 larval transport from GB/GSC toward the MAB. In the diel migration case, the larvae
442 over Nantucket Shoal were advected to the shelf break. That did not happen in the
443 semidiurnal migration case. The differences shown in abundance for C#1-C#5 were
444 observed alternatively from the larval density distributions shown in Figs. 7a-e).

445 The model predicts that the dispersal and settlement of scallop larvae varied
446 significantly from year to year, which was evident in a comparison between 2009 and
447 2008. In 2009, regardless of larval vertical behaviors, many scallop larvae were advected
448 to the SNE shelf and entered the MAB (Fig. 8). The main difference among C#1-C#5
449 was the distributions of larval settling locations, abundance, and pathways from GB/GSC
450 to the MAB. The distributions of larval density in C#1, C#2, and C#3 were similar (Figs.
451 8f, 8g, 8h), except for the higher density spots occurring east of Long Island and over the
452 MAB in C#2 and C#3. As the OML was deepened to 30 m, the larval dispersal
453 dramatically changed. Over GB, a large portion of larvae was settled and concentrated
454 within the mixed area in the diel migration case (C#4) (Fig. 8i), while they expanded to
455 cover the most area of the bank in the semidiurnal migration case (C#5) (Fig. 8j).
456 Furthermore, the OML deepening caused larvae to shift toward the shelfbreak on their
457 journey to the MAB. The highest larval density was found in the MAB in C#5, but not in
458 C#4. Although significant larvae were advected southward to the MAB, the cases with
459 larval vertical migration behaviors in the OML still provided a higher larval retention rate
460 on GB. In C#1, 33.0% of larvae were settled over GB/GSC. The retention rate varied
461 with the OML depth and larval behaviors. For C#2-C#4, it was increased from 39.6% to
462 56.2% when the OML deepened from 10 m to 30 m, while for C#5, it remained similar
463 for the 10- and 30-m OML cases. The features described here can be viewed alternatively
464 from the larval density distributions for C#1-C#5 shown in Figs. 8a-e).

465 2012 was a warm year during which the nearshore sea temperature increased by ~1.0-
466 2.0°C. Warming intensified the cross-isobath gradients of the bottom temperature over
467 the middle shelf and shelfbreak. The settlement of larvae is influenced considerably by
468 larval behaviors in the OML and the OML depth. For C#1, many larvae were transported
469 to the SNE shelf and even entered the MAB, with the highest abundance over GB and
470 within the cold pool south of Long Island (Figs. 9a, 9f). When diel and semidiurnal larval

471 vertical migration behaviors were considered in a fixed 10-m depth OML (C#2 and C#3),
472 the larvae over GB were aggregated around GSC, with a portion entering the SNE shelf
473 (Figs. 9b, 9g, 9c, 9h). Although the larval distribution patterns for C#2 and C#3 were
474 similar, the larval dispersal was more extensive in the semidiurnal migration case than in
475 the diel migration case. As the OML depth deepened to 30 m, most larvae were retained
476 on GB and around GSC. No larvae were advected southward to enter the MAB. For a
477 given OML depth, the larval distributions varied with larval behaviors in the OML. For
478 C#4, the settled larvae showed a dispersive distribution on GB, with the highest
479 abundance in the cold pool area over Nantucket Shoal west of GSC (Figs. 9d, 9i). For
480 C#5, the larvae were settled around the tidal-mixing front on GB, with a dense
481 aggregation around GSC (Figs. 9e, 9j). The results for C#4 and C#5 were correlated well
482 with the extremely high recruitment found in NLCA from 2012 (*Bethoney et al.*, 2016).

483 Changes in the larval dispersal and settlement with the OML depth and larval
484 behaviors in 2013 were similar to that found in 2012 (Fig. 10). Either ignoring larval
485 behaviors in the OML (C#1) or having larval behaviors in a thin OML (C#2 and C#3)
486 overestimated the southward larval transport. The deeper OML favored larval retention
487 over GB/GSC and Nantucket Shoal (C#4 and C#5). For a given 30-m OML, the larval
488 dispersals significantly differed for the diel (C#4) and semidiurnal (C#5) migration cases.
489 For C#4, the highest larval aggregation area was on the SNE (Figs. 10d, 10i), while for
490 C#5, it was around the GSC (Figs. 10e, 10j). Over GB, similar to 2012, the settled larvae
491 were distributed on the top and western areas in the C#4 case, while they occupied the
492 entire bank in the C#5 case.

493 The significant difference among C#1-C#5 for 2008, 2009, 2012, and 2013 illustrates
494 that the larval dispersal and settlement varied not only by the changes in physical
495 environments but also with larval behaviors in the OML. Larval behaviors in the OML
496 made larvae stay longer in the vertical column before settling, increasing the larval
497 residence time on GB. Thus, ignoring it will overestimate the larval transport to the SNE
498 shelf and MAB.

499 **3.2. Influences of larval behaviors in the varying-thickness OML**

500 The OML depth varied significantly in time and space, especially during spring and
501 autumn (Flagg, 1987). In these two seasons, it was in a range of 10-40 m over the shelf

502 (Li *et al.*, 2020). The vertically well-mixed and stratified areas were distinct in the model-
503 predicted mean water density profilers throughout September-November. In 2013, for
504 example, the water was vertically well-mixed in areas where bottom depths were
505 shallower than 50 m over GB and Nantucket Shoal, while it was strongly-stratified on
506 the southern flank of GB, in GSC, and over middle/outer shelves of SNE and MAB (Fig.
507 11). Three sections labeled A, B, and C were selected to show the variability of the OML
508 on the eastern and southern flanks of GB and the SNE shelf over September-November
509 (Fig. 12). Over GB, in the areas between tidal and shelfbreak fronts, the OML depth was
510 ~10 m in September and then gradually increased to ~30-40 m or deeper in November
511 (Fig.12: see A and B). Within the shelfbreak front, the OML depth remained steady after
512 October. On Section-B, the OML thinned rapidly in November, suggesting a local scale
513 onshore intrusion of the stratified Gulf Stream water during that period. The temporal
514 variability of the OML at Section-C over the SNE shelf was similar to that at Section-A
515 on the eastern flank of GB.

516 To examine the influence of larval behaviors in a varying OML on the dispersal and
517 settlement of scallop larvae, we repeated the 2013-2016 experiments with the real-time
518 OML provided hourly from NECOFS (C#6 and C#7). We also ran the model with a
519 thermocline-seeking larval behavior in the same model-predicted OML (C#8). These
520 additional cases were conducted over the same period, starting on September 1 and
521 ending on November 30. The comparison was made among results obtained for eight
522 cases (C#1-C#8) with and without the inclusion of larval behaviors.

523 The results showed that the variability of the OML had a marked influence on the
524 scallop larval dispersal. An example was exhibited here for 2013 simulation results.
525 Although the settled larval distributions were similar between C#6 (Figs. 13a, 13d) and
526 C#4 (Figs. 10d, 10i) and also between C#7 (Figs. 13b, 13e) and C#5 (Figs. 10e, 10j), the
527 spatiotemporal variation of the OML pushed larvae in the highly abundant area
528 northward to the Nantucket Sound in C#6 (Figs. 13a, 13d) and aggregated larvae on the
529 western shelf of GSC in C#7 (Figs. 13b, 13e). C#8 considered a case for constraining
530 larvae at the bottom of the OML. In this case, most of the larvae aggregated on southern
531 and western flanks of GB, within the region between 50- and 100-m isobaths (Figs. 13c,
532 13f). The highest larval density area was in the GSC area, but the abundance was much

533 smaller than those found for C#7. For C#7 and C#8, either semidiurnal migration or
534 thermocline-seeking behavior consistently predicted a larval aggregation in the closed
535 area around GSC. This feature was not captured in the case without larval behaviors in
536 the OML.

537 Changes in the residence time of larvae in the water column on GB were one of the
538 reasons for distinct differences in the larval dispersal and settlement for C#1-C#8. For
539 example, tracking a super-individual originating from the same initial location on GB for
540 these eight cases, we examined horizontal and vertical movements of this super-
541 individual under different biophysical environments (Fig. 14). In each case, the tracking
542 period was 41 days, with its trajectory sampled daily. For C#1, the super-individual
543 migrated upward to the sea surface at the 5-day age and then stayed there until they grew
544 to the 40-day age. The near-surface flow rapidly advected this super-individual
545 southward along the shelf, with a residence time of ~15 days on GB (Fig.14a). When
546 larval behaviors in the OML were considered, the daily larval trajectory varied with the
547 sampling method. Here sampling was taken at noon each day. At this time, the larvae
548 were mainly at the bottom of the OML regardless of diel, semidiurnal, and thermocline-
549 seeking larval behaviors.

550 For C#2 and C#3, the super-individual migrated upward to the subsurface at a depth
551 of 10 m at the 5-day age and moved southward following a daily mean trajectory at the
552 bottom of the OML (Figs. 14b, 14c). After 40 days, it settled to the seabed around GSC.
553 Compared with the diel migration behavior, the semi-diurnal migration behavior favored
554 retaining the larvae on GB, even though their trajectories almost coincided during the
555 first 7 days. As a result, the super-individual settled on the western shelf of GSC in C#2,
556 but within the GSC in C#3 (Fig. 14b).

557 Similar features were also found for C#4 and C#5 when the OML depth was
558 deepened to 30 m. In the diel vertical migration case (C#4), after the super-individual
559 migrated upward to enter the OML, it followed a daily trajectory at the bottom of the
560 OML to move southward along the bank (Fig. 14c). This super-individual then settled
561 down near the shelf break of the SNE shelf. Differing from C#4, the super-individual in
562 C#5 was trapped locally after 8 days and eventually settled around 60-m isobath area on
563 the southern flank of GB after 40 days (Fig. 14c). For a given fixed-depth OML, the

564 longer distance in vertical migration tended to make the larvae move slowly in the
565 horizontal. This feature was also observed in the spatiotemporally-varying OML cases,
566 even though horizontal and vertical trajectories of the super-individual significantly
567 differed.

568 The diel vertical migration behavior (C#6) was less favorable to retain the larvae on
569 GB compared with semidiurnal (C#7) and thermocline-seeking (C#8) vertical migration
570 behaviors (Fig. 14d). For C#6, the super-individual followed the clockwise gyre
571 circulation to drift along the bank during the first 35 days, then turned northward on the
572 western GB, and eventually settled at the seabed east of the GSC. The trajectory of this
573 super-individual varied significantly in the vertical before settling. For C#7 and C#8, the
574 semidiurnal or thermocline-seeking vertical migration pushed the super-particle offshore
575 toward the shelfbreak front, retained it in the deeper depth, and eventually made it settle
576 on the southeastern flank of GB, an area close to its origin. In these two cases, the
577 thermocline-seeking behavior was more favorable to restrain the horizontal movement
578 than the semidiurnal behavior. It explains why similar aggregation patterns were found
579 for C#7 and C#8 around the GSC. The comparison of horizontal and vertical trajectories
580 of the same super-individual in these eight cases again highlights the importance of
581 including larval behaviors in the OML in the Scallop-IBM, especially for the early life
582 stage simulation.

583 **3.3. Statistics and connectivity of scallop larvae over GB/GSC, SNE, and the** 584 **MAB**

585 Dividing the model domain into 2×2 km boxes, we statistically calculated the mean,
586 percentage, and standard deviation of larval density over 39 years from 1978 to 2016 for
587 C#1-C#5, respectively. Probability is represented by the settling percentage of larvae in
588 each box over 39 years, ranging from 0 (0%) to 1 (100%). Standard deviation was
589 estimated relative to the 39-year mean, which illuminated the range of the interannual
590 variability. For C#1, the mean larval density remained high over GB/GSC and SNE, with
591 a significant interannual variability occurring in the SNE and MAB region (Figs. 15a-c).
592 In this case, the probability rate of larvae entering the MAB was up to 50%. For C#2 and
593 C#3, the diel vertical larval migration tended to retain larvae over GB/GSC and SNE,
594 with maximum interannual variability occurring over the SNE shelf and northern area of

595 the MAB (Figs. 15d-i). In these two cases, the model showed that including the larval
596 behavior in the OML considerably reduced the probability rate of larvae entering the
597 MAB. The major difference between these two cases was in the spatial distribution of
598 settled larvae over GB/GSC and SNE. In the semidiurnal case, more larvae accumulated
599 in the eastern portion of NLCA and the center of GB. For C#4 and C#5, deepening of the
600 OML favored the larval retention over GB/GSC and SNE and restricted larval transport
601 from entering the MAB, even though it happened occasionally (Figs. 15j-o). Similar to
602 the 10-m OML case, the primary difference between diel and semidiurnal migration cases
603 was in the spatial distribution of settled larvae. The semidiurnal migration behavior in the
604 OML led to denser larval accumulation in the three closed areas, especially in the
605 northern portion of CA-II over the northeastern flank of GB. Regardless of whether
606 larval swimming behaviors in the OML were considered, the SNE was a region featuring
607 the maximum larval interannual variability.

608 We estimated the percentage of larvae settling in three geographic zones of GB/GSC,
609 SNE, and the MAB (see the boundary of each zone in Fig. 1) for C#1-C#5, respectively.
610 The model consistently predicted that GB/GSC was a high retention area (Fig.16 and
611 Table 2). C#2 and C#3, also C#4 and C#5, exhibited a similar interannual variability
612 pattern. On GB/GSC, the mean differences over 1978-2016 were 7.0% between C#2 and
613 C#1, and up to 10.2 between C#3 and C#1, indicating that the semidiurnal migration
614 behavior increased the retention by ~3.2% (Fig. 16a). When the OML depth was
615 deepened to 30 m, the retention rate on GB/GSC was decreased by 3.7% for the diel
616 migration case and 7.0% for the semidiurnal migration case. The SNE shelf was also a
617 high aggregation area of scallop larvae (Fig. 16b). In this region, considering larval
618 behaviors in the OML increased the larval settlement rate. The rate became higher as the
619 OML deepened. The 39-year mean difference was 6.9% between C#2 and C#1, and 5.6%
620 between C#3 and C#1. The difference was up to 23.3% between C#4 and C#1, and
621 18.8% between C#5 and C#1.

622 The most considerable difference among C#1, C#2, C#3, C#4, and C#5 was the larval
623 settlement rate in the MAB. For C#1, the model predicted a sizeable larval transport to
624 the MAB, with a 39-year mean of 22.1% and a maximum of up to 40% (Fig. 16c). The
625 larval transport to the MAB was considerably reduced by taking larval behaviors in the

626 OML into account. Except for 2009, it was about 10% or less than for C#2 and C#3, 5%
627 or less for C#4, and close to zero for C#5. The 39-year means for C#2-C#5 were 8.2, 6.3,
628 1.8, and 0.7%, respectively. These results suggest that the GB/GSC and MAB scallop
629 populations were poorly connected by larval transport. The high scallop abundance
630 observed in the MAB might have been produced by a high recruitment rate of larvae
631 spawned in the local region.

632 We started implementing a method to determine the real-time OML depth in the
633 simulation in 2013. The experiments for varying OML were done for 2013-2016. The
634 statistics of these four-year results for C#6-C#8 showed that regardless of vertical
635 migration patterns, the GB/GSC and SNE had high scallop larval settlement, with the
636 maximum interannual variability occurring over the SNE shelf (Fig. 17, Table 3). In
637 particular, the spatiotemporal variability of the OML led to denser larval accumulation in
638 the NLCA. No larvae were advected into the MAB in all three cases of C#6, C#7, and
639 C#8. We also estimated the percentage of larvae settling in three geographic zones of
640 GB/GSC, SNE, and the MAB for these three cases and compared the results with C#5.
641 For the semidiurnal migration case, the interannual variability for C#5 and C#7 exhibited
642 a similar pattern in the GB/GSC and SNE regions (Fig. 18). The spatiotemporally-
643 varying OML produced a high retention rate on GB/GSC, with a 5.4% difference
644 between GB/CSC and SNE regions for these two cases. Also, C#7 predicted less larval
645 transport to the MAB than C#5, even the transports for both cases were close to zero. For
646 the diel migration case, although the settled larvae percentages in the GB/GSC and SNE
647 regions showed a similar variation for C#6 and C#4, the spatiotemporally-varying OML
648 produced a more favorable condition to retain the larvae on GB/GSC than the fixed-depth
649 OML. The difference was up to 9.5% between GB/GSC and SNE regions for these two
650 cases. The larval settlement showed relatively large variability in C#8. The mean
651 percentages over 2013-2016 were 62.9% over GB/GSC, 37.2% over the SNE shelf, and
652 0.0% entering the MAB.

653

654

4. Discussion

655 Our results indicate that the larval vertical migration in the OML can significantly
656 influence the dispersal and settlement of scallop larvae over GB/GSC and SNE, as well

657 as larval transport to the MAB. In the GB/GSC and SNE regions, although the 39-year
658 mean difference was in the range of ~10% or less between C#1 and C#2-C#5, their
659 dispersal patterns differed considerably. Vertical migration made scallop larvae stay
660 longer in the water column on GB/GSC as compared to passive larvae, because it
661 exposed them to different currents in the deeper water, which were slower and more
662 cyclonic (Werner *et al.*, 1993; Page *et al.*, 1999). As a result, the larvae originating from
663 eggs spawned on GB, mainly drifted around the bank following the clockwise residual
664 flow and eventually settled on GB and surrounding SNE areas. Only a few moved
665 southwards to enter the MAB.

666 The conclusions in Tian *et al.* (2009a, 2009c) were similar to our findings for C#1
667 (without swimming behaviors) but very different from the results for C#2-C#8
668 (swimming that oscillated between subsurface depths). We believe that the difference
669 was due to the physics and larval behaviors. Tian *et al.*'s (2009a-c) simulations did not
670 include the Gulf Stream-shelf interaction and inflow from the upstream Labrador Sea and
671 the Arctic Ocean. The currents used to drive the Scallop-IBM significantly differed from
672 the NECOFS fields used in this study, especially at the shelf break where the Gulf Stream
673 influences were significant. Tian *et al.* (2009c) implemented a thermocline-seeking
674 larval behavior in the Scallop-IBM. They assumed that the OML depth remained constant,
675 with thermoclines always at a depth of 23 m. Once larvae migrated to 23 m, they drifted
676 as passive particles along with the horizontal flow at that depth. The simulation covered
677 1995-2005, and the results showed significant larval transport to the MAB in 1998, 2001,
678 2004, and 2005. Especially in 2005, the larval settlement in the MAB was even more than
679 larvae settled over GB/GSC. Comparing our simulation results with Tian *et al.* (2009a,
680 2009c) for the same period 1995-2005, we found that no matter how the OML depth was
681 specified, the models predicted a high aggregation over GB/GSC and SNE, and a weak
682 connection between GB/GSC and the MAB. Even in 2005, the larval transport to the
683 MAB was only around 10% for C#2 and C#3 and close or equal to zero for C#4 and C#5.
684 Over 2013-2016, we repeated the thermocline-seeking larval behavior experiment (C#8)
685 with a similar approach used in Tian *et al.* (2009c), but we considered the spatiotemporal
686 variation of the OML depth (Fig.17). In this case, larval transport to the MAB was non-
687 existent.

688 *Tian et al. (2009c)* argued that vertical migration played a less critical role in the
689 dispersal and settlement of scallop larvae originating from GB/GSC. Their argument was
690 based on two pieces of evidence observed by *Gallager et al. (1996)* and *Tremblay and*
691 *Sinclair (1990a)*. *Gallager et al. (1996)* detected the larvae migration in the OML,
692 aggregating twice at the sea surface during the night and at the bottom of the OML during
693 the day (e.g., Fig. 4). The measurements were made in a thin OML of ~4 m (mesocosm).
694 *Tian et al. (2009c)* assumed that such a short-distance vertical migration would not affect
695 the larval dispersal since the horizontal drifting velocity zone or the residence time
696 remained unchanged. The fact was that the OML depth varied significantly in autumn,
697 especially during a storm event (*Li et al., 2020*). *Tremblay and Sinclair's* profiler
698 sampling showed a high larval abundance within thermoclines at depths varying in the
699 range of 12-23 m on GB. Based on this observation, *Tian et al. (2009c)* questioned
700 whether active larval vertical migration was a general feature on GB. The fact was that
701 profiler sampling was done at different times, and each was completed in 74 min. A few
702 in-situ observations were not sufficient to cover the daily migration period. Small
703 amplitude diel vertical migration was also found in a shallow area of < 25 m off Grand
704 Mann Island in the Gulf of Maine by *Tremblay and Sinclair (1990b)*. Therefore, it may
705 have been premature to conclude that no vertical migration of larvae existed in the region.

706 The scallop larval dispersal and settlement results for cases with semidiurnal and
707 thermocline-seeking migrations (C#7 and C#8) suggest that there was almost no larval
708 connectivity between GB/GSC and the MAB. Although the larval distributions for these
709 two cases differed and the settlements showed more considerable variability in C#8 than
710 in C#7, the 4-year mean settled larval percentages in either GB/GSC or SNE regions
711 were 5.4% or less for these two cases.

712 Our simulation results with larval migrations within the OML show that 2009 was a
713 year with a significant larval transport from GB/GSC to the MAB. Since that year, the
714 retention rate of migrating larvae in the GB/GSC and SNE regions remained a high value,
715 with almost no larvae transporting southward into the MAB. The bottom temperature
716 over the northeast shelf was characterized by a cold pool, forming in spring, and
717 gradually decaying through autumn (*Lentz et al., 2003, Lentz, 2017*). Although this cold
718 pool's intensity was considerably weak in autumn, it was still visible as a relatively

719 uniform cold temperature region bounded by 12-13°C contours in Fig.19. Compared with
720 the climatological mean bottom temperature over 1978-2008 (Fig. 19a), in 2009, the cold
721 pool area expanded onshore over the SNE shelf and shrank towards the shelfbreak south
722 of Long Island (Fig. 19b). 2012 was a warm year with a ~2°C rise of the bottom
723 temperature in the tidally well-mixed area of GB and nearshore regions (Fig. 19c).
724 Warming significantly shrank the area of the cold pool and pushed it offshore. The well-
725 defined cold pool disappeared on the southern flank of GB due to the warming-induced
726 intensification of the cross-isobath gradient of bottom temperature. This feature was
727 sustained over 2013-2016 (Fig. 19d). The cold pool functioned as an index for the
728 GB/GSC, SNE, and MAB connectivity. The weakening of the cold pool's intensity and
729 intensified cross-isobath gradient of bottom temperature tends to enhance the clockwise
730 gyre circulation over GB, which indirectly supported our finding: warming has restricted
731 the larval transport from GB/GSC to the MAB.

732 The warming tendency was evident in the satellite-derived sea surface temperature
733 (SST) change over the U.S. northeastern shelf in the past decades (Fig. 20). Significant
734 warming occurred in 2012. After that, the water remained warmer. The yearly warming
735 rate of the SST averaged over the shelf bounded at the 300-m isobath was ~0.04 over
736 1982-2020 (Fig. 20a). Assuming 2012 as a year for warming regime shift, the mean SST
737 after that was about 1.0°C higher than the climatological SST mean averaged over 1982-
738 2011. This warming feature was captured in the NECOFS simulation. The warming rate
739 in the region varied significantly in space, with the maximum around the shelfbreak off
740 GB (Fig. 20b). We examined the NECOFS-predicted subtidal flow field in the region and
741 found a branch of the Gulf Stream that flowed northeastward towards GB. This branch
742 flow has been intensified significantly in recent years, causing extreme warming at the
743 shelfbreak off GB. As we detected in the NECOFS-simulated temperature and flow fields,
744 the warming has intensified the cross-isobath gradient of water temperature on the
745 southern flank of GB and thus strengthened the clockwise gyre over the bank.

746 The model predicted extensive southward water transports in the autumn of 2009.
747 Selecting a cross-shelf section over the SNE shelf (see the location in Fig. 1), we
748 calculated the water transport through that section over 1978-2016. Across that transect,
749 the 39-year mean transport was -0.46×10^{-3} Sv (Sv = 10^6 m³/s). The anomaly exhibited

750 relatively large positive (northward) and negative (southward) phases in 2008 and 2009,
751 respectively, and remained positive since 2011 (Fig. 21). The anomaly's interannual
752 variability explains why the larval transport to the MAB was most extensive in 2009, and
753 no connectivity between GB/GSC and the MAB had occurred since 2010. The wind was
754 a primary driver for the sizeable southward transport in autumn of 2009. The wind
755 records at Buoy#44008 show that differing from other years, the northeasterly wind
756 prevailed over the northeast shelf during autumn of 2009, with a maximum speed of >16
757 m/s (Fig. 22). The extreme northeasterly or northerly winds tended to push the water
758 onshore. It enhanced the southward along-shelf flow under a balance between the
759 pressure gradient and earth rotation-induced Coriolis forces. The flow intensification was
760 the reason why a large number of larvae drifted to the MAB in that year. This result
761 suggests that in addition to larval vertical migration behaviors in the OML, the GB/GSC
762 and MAB connectivity also depends on the intensity and duration of northeasterly winds
763 during the fall spawning season.

764 It should be pointed out that scallop spawning over GB/GSC varies interannually.
765 This variability has not been taken into account in this study. We have not considered any
766 size-dependency of spawning either (*Davies et al.*, 2014). No experiments were done for
767 the case of spawning in the MAB, as it is unlikely that the larvae could be transported
768 northward to SNE, against the prevailing southward along-shelf flow. Recent
769 observations revealed persistent warming in the region. NECOFS shows that warming
770 has produced a positive anomaly of water transport over the SNE shelf since 2011. An
771 enhanced northward flow in autumn could advect larvae in the MAB to the upstream
772 SNE region. It is worth examining these questions in the future using the 39-year hourly
773 hindcast NECOFS product, which can provide insights into the biophysical processes
774 attributing to the mixing and exchanges of larvae between the GB and MAB scallop
775 populations in the SNE region.

776 We did not consider the spring spawn in our experiments. The spawning time of sea
777 scallops varies latitudinally across its range, extending from the Strait of Belle Isle,
778 Newfoundland, to Cape Hatteras, North Carolina (*Posgay*, 1957; *Barber and Blake*, 2006;
779 *Stokesbury and Bethoney*, 2020). Annual autumn spawning is typical in Newfoundland
780 (*MacDonald and Thompson*, 1986), whereas semi-annual spawning is characteristic of

781 the MAB (*DuPaul et al.*, 1989). On GB, the autumn spawn is dominant, while spring
782 spawning varies in magnitude and temporally (*Chute et al.*, 2012; *Hennen and Hart*, 2012;
783 *Davis et al.*, 2014; *Thompson et al.*, 2014; *Davis et al.*, 2015). Depending on mortality
784 estimates, spring-spawning contributes minimally up to about one-third of the annual
785 total larval settlement (*Davis et al.*, 2014). For example, *Chute et al.* (2012) examined 14
786 scallops with stable isotopes, 13 of which were fall spawned, including 6 from GB and
787 Nantucket Shoals. The one that was spring spawned was likely spawned in the MAB. The
788 spawning cycle, fertilization success, larval survival, and dispersion are all influenced
789 heavily by the environment. As oceanographic conditions change on GB, spring-
790 spawning may become increasingly important as it is in the MAB. It could also affect the
791 larval connectivity between the GB/GSC and the MAB like that detected by *Davies et al.*
792 (2014).

793 Our studies considered various larval swimming behaviors, which require additional
794 field confirmation. Recently, *Norton et al.* (2020) examined the impact of ocean
795 conditions on the recruitment of Dungeness crab (*Metacarcinus magister*) in the U.S.
796 Pacific Northwest. Their studies examined six swimming behaviors. Considering these
797 behaviors in a generalized linear model (GLM) with superior fits to the observations, they
798 found that the ensemble solution with various swimming behaviors in the larval IBM
799 model could improve predicting larval crab dispersion. This ensemble approach could be
800 adopted in the larval scallop simulation, especially in a condition with various
801 unconfirmed swimming behaviors.

802

803

5. Conclusions

804 With spawning based on multiyear-averaged abundance and distribution of adult sea
805 scallops over GB/GSC, we examined the impacts of physical processes and larval
806 swimming behaviors within the OML on the interannual variability of the scallop larval
807 dispersal and settlement in the GB/GSC, SNE, and MAB regions over 1978-2016. The
808 study was conducted using the coupled Scallop-IBM and NECOFS model. The results
809 indicate that in addition to the flow-induced advection, larval behaviors in the OML
810 significantly affected larval dispersal and settlement by altering the flow-induced
811 advection experienced at different depths. The thermocline-seeking, diel or semidiurnal

812 migration behaviors of larvae in the OML increased the larval residence time in the water
813 column over GB/GSC. These behaviors led to persistent larval aggregations in the
814 GB/GSC and SNE regions. In addition to larval behaviors, larval transports to the MAB
815 were also closely related to the intensity and duration of northeasterly wind in autumn.
816 No functional connectivity of larvae between GB/GSC and the MAB occurred in the past
817 39 years, except in the autumn of 2009, during which an extreme northeasterly wind
818 prevailed. Neglecting larval behaviors in the OML can exaggerate the connectivity scale
819 of the GB and MAB sea scallop populations. Our studies suggest this connectivity will
820 only matter in intense wind scenarios as expected with future climate change.

821 SNE is the region featuring a maximum interannual variability of larval settlement.
822 The NECOFS has captured the climate change-induced warming over the U.S.
823 northeastern shelf. The extreme warming at the shelfbreak off GB has significantly
824 intensified the cross-isobath gradient of water temperature and enhanced the clockwise
825 subtidal gyre over the bank. This change tends to increase the larval retention rate over
826 GB/GSC, suggesting higher scallop recruitment in the future.

827

828 **Appendix A: A method to calculate the thickness of the ocean mixed layer**

829 The thickness of the surface ocean mixed layer (OML) is defined as a depth above
830 which the water density remains essentially unchanged in the vertical. In practice, it is
831 usually determined using a threshold approach with a criterion relative to a reference
832 value (*e.g.*, *de Boyer Montégut et al.*, 2004). Here we introduced a method based on the
833 density profile.

834 Defining H as the bathymetric depth at a particular geographic location, ρ as the
 835 water density that varies vertically from $\mathbf{z} = \mathbf{0}$ at the surface to $\mathbf{z} = -\mathbf{H}$ at the bottom
 836 and ρ_o as the surface water density, we can estimate the mixed layer depth (\mathbf{h}_m by

$$837 \quad \mathbf{h}_m = \mathbf{H} - \sqrt{2\mathbf{h}_{diff}/\gamma} \quad (\text{A.1})$$

838 where $\mathbf{h}_{diff} = \mathbf{h} - \rho_o\mathbf{H}$; $\mathbf{h} = \int_{-\mathbf{H}}^0 \rho d\mathbf{z}$; and γ is defined as the maximum increase rate of
 839 the density with depth. Once γ is determined from a density profile, we can precisely
 840 estimate h_m . To demonstrate how this method work, examples are given below for three
 841 idealized cases.

842 Case 1: A vertically well-mixed case with a density profiler shown in Fig. A1. In this
 843 case, ρ is constant throughout the water column, so that

$$844 \quad \rho = \rho_o; \mathbf{h} = \rho_o\mathbf{H}; \text{ and } \mathbf{h}_{diff} = \mathbf{0}.$$

845 Substituting h and h_{diff} into (A.1), we have $h_m = H$. Note here that $\gamma = 0$. For a real
 846 application, one can directly assume h_m equals the local depth.

847 Case 2: A stratified case with a linear density profiler shown in Fig. A2. In this case,

$$\rho = \rho_o - (\rho_H - \rho_o) z/H.$$

Substituting it into (A.1), we have

$$848 \quad \mathbf{h} = \int_{-\mathbf{H}}^0 [\rho_o - (\rho_H - \rho_o) z/H] d\mathbf{z} = (\rho_H + \rho_o) z/H; \mathbf{h}_{diff} = \mathbf{0.5} (\rho_H - \rho_o) \mathbf{H}.$$

849 Also, $\gamma = (\rho_H - \rho_o) /H$, so that $\mathbf{h}_m = \mathbf{H} - \sqrt{2\mathbf{h}_{diff}/\gamma} = \mathbf{0}$.

850 Case 3: A two-layer with a density profiler shown in Fig. A3. In this case, the density
 851 profiler is given as

$$852 \quad \rho = \begin{cases} \rho_o, & -\mathbf{h}_m \leq \mathbf{z} \leq \mathbf{0} \\ \rho_o - (\rho_H - \rho_o) (\mathbf{z} + \mathbf{h}_m) /(\mathbf{H} - \mathbf{h}_m), & \mathbf{z} \leq -\mathbf{h}_m \end{cases}$$

853 and $\gamma = (\rho_H - \rho_o) /(\mathbf{H} - \mathbf{h}_m)$, then, we have

854

855
$$h = \rho_o h_m + 0.5 (\rho_H + \rho_o) (H - h_m)$$

856 and

857
$$h_{diff} = h - \rho_o H = \frac{\rho_H - \rho_o}{2} (H - h_m),$$

858 so that

859
$$h_m = H - \sqrt{2h_{diff}/\gamma} = H - (H - h_m) = h_m.$$

860 With demonstrations from these three idealized cases, we applied this method to
861 calculate the thickness of the OML based on the NECOFS-produced hourly density
862 profile. The result was validated by comparing it with the simulated temperature, salinity,
863 and density profiles at nodes of the triangular mesh. Examples are shown in Fig. A4 for
864 selected three sites across GB. Using (A.1), we calculated h_m at these sites. They equaled
865 14.8, 5.0, and 9.1 m, respectively. Marking the calculated h_m using red dashed lines in the
866 profiles, we found that they matched well with the depth of model-simulated OML.

867

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886

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1166 **Figure Captions**

1167
1168 Figure 1: Schematic of the near-surface (red arrows) and deep (white arrows) flows over
1169 the US northeast shelf. GB: Georges Bank, GSC: Great South Channel, SNE:
1170 Southern New England, MAB: Middle Atlantic Bight. The red color patch
1171 represents the Gulf Stream northward meander water. Red color rings represent
1172 the warm-core ring separated from the Gulf Stream. Gray thick lines are the
1173 boundaries between GB/GSC, SNE, and MAB. The solid black thin line is the
1174 transect where the transport was calculated. The 3-D icon represents the NOAA
1175 buoy, and the number on the right is the buoy number.

1176 Figure 2: The unstructured meshes for Global-FVCOM and GoM-FVCOM. The cells
1177 marked with red colors represent the common cells nesting between Global-
1178 FVCOM and GoM-FVCOM.

1179 Figure 3: Structures of the scallop-IBM early life stage model. Four pelagic stages are
1180 considered: 1) egg, 2) trochophore, 3) veliger, and 4) pediveliger. U , V , and W
1181 are the x, y, and z components of the water velocity. T is the water temperature,
1182 and K_m is the vertical eddy viscosity. The dashed line box presents the pelagic
1183 stages, and the gray shadow area indicates benthic stages.

1184 Figure 4: The diel and semidiurnal larval vertical migration sub-models in the surface
1185 mixed layer during the period of 5 through 40 days from eggs to veliger stages.
1186 Diel and semi-diurnal vertical migration patterns were based on the observations
1187 made by Tremblay and Sinclair (1990b), Manuel et al. (1996), and Gallager et al.

1188 (1996). The number in the figure indicates the time of a day defined by a 24-hour
1189 clock.

1190 Figure 5: Scallop abundance (scallop#/m²) (a) and gridded density (individual/m²) (b) for
1191 spawning. The individuals in each cell were determined using the combined
1192 scallop data from BIO, NOAA, and SMAST. In the upper panel, shapes bounded
1193 by red lines are the closed areas; CA-I: closed area I, CA-II: closed area II, and
1194 NLCA: Nantucket Lightship closed area. In the lower panel, the dashed thick line
1195 is the boundary between the US and Canadian waters.

1196 Figure 6: Illustration of the egg spawning period starting at 00:00 September 1 and
1197 ending at 24:00 October 10. The spawning process satisfies a normal probability
1198 distribution with the maximum on September 20 and a one-week standard
1199 deviation.

1200 Figure 7: Distributions of the settled larval density (a-e) and locations/ abundances of
1201 settled super-individuals (f-j) for the cases C#1 (No OML), C#2 (10 m-OML:
1202 diel), C#3 (10 m-OML: semidiurnal), C#4 (30 m-OML: diel), and C#5 (30 m-
1203 OML: semidiurnal). The results were from the 2008 simulation. Two thick gray
1204 lines are the boundaries between GB/GSC, SNE, and MAB. Gray lines with
1205 labels are 50, 100, and 200-m isobath contours.

1206 Figure 8: Distributions of the settled larval density (a-e) and locations/ abundances of
1207 settled super-individuals (f-j) for the cases C#1 (No OML), C#2 (10 m-OML:
1208 diel), C#3 (10 m-OML: semidiurnal), C#4 (30 m-OML: diel), and C#5 (30 m-
1209 OML: semidiurnal). The results were from the 2009 simulation. Two thick gray
1210 lines are the boundaries between GB/GSC, SNE, and MAB. Gray lines with
1211 labels are 50, 100, and 200-m isobath contours.

1212 Figure 9: Distributions of the settled larval density (a-e) and locations/ abundances of
1213 settled super-individuals (f-j) for the cases C#1 (No OML), C#2 (10 m-OML:
1214 diel), C#3 (10 m-OML: semidiurnal), C#4 (30 m-OML: diel), and C#5 (30 m-
1215 OML: semidiurnal). The results were from the 2012 simulation. Two thick gray
1216 lines are the boundaries between GB/GSC, SNE, and MAB. Gray lines with
1217 labels are 50, 100, and 200-m isobath contours.

1218 Figure 10: Distributions of the settled larval density (a-e) and locations/ abundances of

1219 settled super-individuals (f-j) for the cases C#1 (No OML), C#2 (10 m-OML:
1220 diel), C#3 (10 m-OML: semidiurnal), C#4 (30 m-OML: diel), and C#5 (30 m-
1221 OML: semidiurnal). The results were from the 2013 simulation. Two thick gray
1222 lines are the boundaries between GB/GSC, SNE, and MAB. Gray lines with
1223 labels are 50, 100, and 200-m isobath contours.

1224 Figure 11: Ratio of the model-simulated mixed layer to the local depth averaging over
1225 September-November, 2013. The right lower panel shows the cross-isobath
1226 distributions of temperature and salinity on GB. The solid black thick line is the
1227 location of the section. Black lines are 50, 100, and 200-m isobath contours.

1228 Figure 12: Cross-isobath sections (thick white lines) labeled "A, B, and C" and the depths
1229 of the monthly averaged OML for September, October, and November 2013 on
1230 Sections A, B, and C, respectively. Red line: September, blueline: October, and
1231 blackline: November. Black lines are the isobath contours matching with depth
1232 images.

1233 Figure 13: Distributions of the settled larval density (a-c) and locations/ abundances of
1234 settled super-individuals (d-f) for the cases C#6 (varying OML: diel), C#7
1235 (varying OML: semidiurnal), and C#8 (thermocline-migration). The results were
1236 from the 2013 simulation. Two thick gray lines are the boundaries between
1237 GB/GSC, SNE, and MAB. Gray lines with labels are the 50, 100, and 200-m
1238 isobath contours.

1239 Figure 14: Horizontal and vertical trajectories of a super-individual originating from the
1240 same site on the southeastern flank of GB. a: C#1 (No OML); b: C#2 and C#3 (10
1241 m-OML); c: C#4 and C#5 (30 m-OML); d: C#6, C#7, and C#8 (Varying OML). t_d :
1242 diel; t_{sd} : semidiurnal; m_b : thermocline-seeking. The results were from the 2013
1243 simulation. Black lines are the isobath contours matching with depth images.

1244 Figure 15: The 39-year mean, percentage, and standard deviation of settled scallop larvae
1245 over 1978-2016 for C#1-C#5. a-c: C#1 (No OML); d-f: C#2 (10 m-OML: diel); g-
1246 i: C#3 (10 m-OML: semidiurnal); j-l: C#4 (30 m-OML: diel); m-o: C#5 (30 m-
1247 OML: semidiurnal). Two thick gray lines are the boundaries between GB/GSC,
1248 SNE, and MAB. Gray lines are the 50, 100, and 200-m isobath contours (see Fig.
1249 11 for isobath labels).

1250 Figure 16: Model-predicted percentages of the scallop larvae settling in the GB/GSC (a),
1251 SNE (b), and MAB (c) regions, respectively, over 1978-2016 for C#1 (solid black
1252 line), C#2 (solid blue line), C#3 (dashed blue line), C#4 (solid red line), and C#5
1253 (dashed red line).

1254 Figure 17: The 4-year mean, percentage, and standard deviation of settled scallop larvae
1255 over 2013-2016 for C#6, C#7, and C#8. a-c: C#6 (varying OML: diel); d-f: C#7
1256 (varying OML: semidiurnal); g-i: C#8 (thermocline-migration). Two thick gray
1257 lines are the boundaries between GB/GSC, SNE, and MAB. Gray lines are the 50,
1258 100, and 200-m isobath contours (see Fig. 11 for isobath labels).

1259 Figure 18: Model-predicted percentages of the scallop larvae settling in the GB/GSC (a)
1260 and SNE (b) regions, respectively, over 2013-2016 for the cases C#4 (30 m-OML:
1261 diel), C#5 (30 m-OML: semidiurnal), C#6 (varying OML: diel), C#7 (varying
1262 OML: semidiurnal), and C#8 (thermocline-migration).

1263 Figure 19: Distributions of the three-monthly averaged bottom temperature in the region
1264 covering GB, SNE, and the MAB over September-November. a: 1978-2008-
1265 averaged; b: 2009; c: 2012; d: 2013-2016 averaged.

1266 Figure 20: b: distribution of the yearly surface temperature increase rate calculated based
1267 on the satellite-derived SST data over 1982-2020. The temperature increase rate
1268 was estimated based on the annual increase rate calculating over two consecutive
1269 years. a: the change of the satellite-derived SST over the shelf bounded by the
1270 300-m isobath over 1982-2019. Solid black dots: the yearly averaged SST for
1271 each year; thick red line: the linear regression fitting line; thick blue dashed lines:
1272 averaged SSTs over 1982-2011 and 2012-2020, respectively.

1273 Figure 21: Anomalies of the water transport through an across-shelf section over the SNE
1274 shelf (see the location in Figure 1) over 1978-2016. The value listed in the upper-
1275 right area is the 39-year mean water transport.

1276 Figure 22: The wind rose plot at NOAA buoy 44008 for September-November, 2009.

1277 Figure A1: Illustration of the density profile under a vertically well-mixed condition for
1278 Case 1.

1279 Figure A2: Illustration of a linear density profile under a stratified condition for Case 2.

1280 Figure A3: Illustration of a two-layer system in which the water density is constant in the

1281 upper layer and linearly increases with depth in the lower layer for Case 3.
 1282 Figure A4: Vertical profiles of sea temperature (red), salinity (blue), and density (black)
 1283 at three sites across GB at 00:00 GMT, September 1, 2013. The thick dashed line
 1284 represents the OML depth calculated using Eq. A.1 in Appendix A.
 1285
 1286

Table 1: Types of numerical experiments made in this study

Parameters Case	OML	Larva behavior
Case 1 (C#1)	No	No
Case 2 (C#2)	10 m	diel migration
Case 3 (C#3)	10 m	semidiurnal migration
Case 4 (C#4)	30 m	diel migration
Case 5 (C#5)	30 m	semidiurnal migration
Case 6 (C#6)	varying	diel migration
Case 7 (C#7)	varying	semidiurnal migration
Case 8 (C#8)	varying	thermocline-seeking

1287

Table 2: Mean percentages and standard deviations of larvae settling in GB/GSC, SNE, and MAB over 1978-2016 for C#1-C#5.

Zone Case	GB/GSC	SNE	MAB
C#1: No OML	43.7±12.4	34.2±12.5	22.1±13.9
C#2: 10-m OML: diel	50.7±6.5	41.1±6.3	8.2±6.3
C#3: 10-m OML: semidiurnal	53.9±7.5	39.8±5.8	6.3±4.9
C#4: 30-m OML: diel	40.7±7.0	57.5±6.6	1.8±2.7
C#5: 30-m OML: semidiurnal	46.3±7.2	53.0±7.5	0.7±2.8

1290

Table 3: Mean percentages and standard deviations of larvae settling in GB/GSC, SNE, and MAB over 2013-2016 for C#6, C#7, and C#8.

Zone Case	GB/GSC	SNE	MAB
C#6: Varying OML: diel	53.5±7.0	46.5±7.1	0.0±0.1
C#7: Varying OML: semidiurnal	57.7±6.1	42.5±6.1	0.0±0.0
C#8: Varying OML: thermocline-seeking	62.9±8.8	37.1±8.8	0.0±0.0

1293

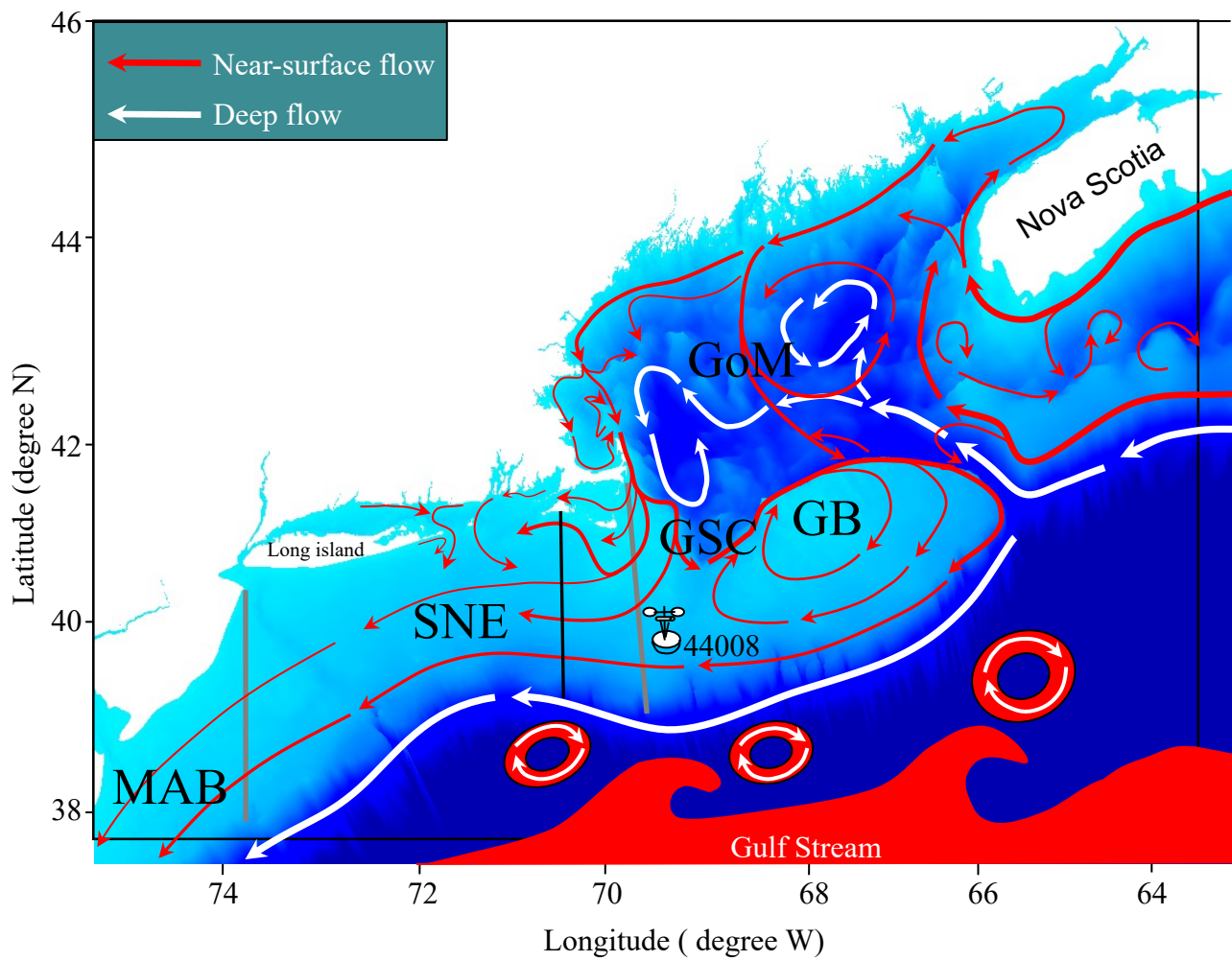


Figure 1

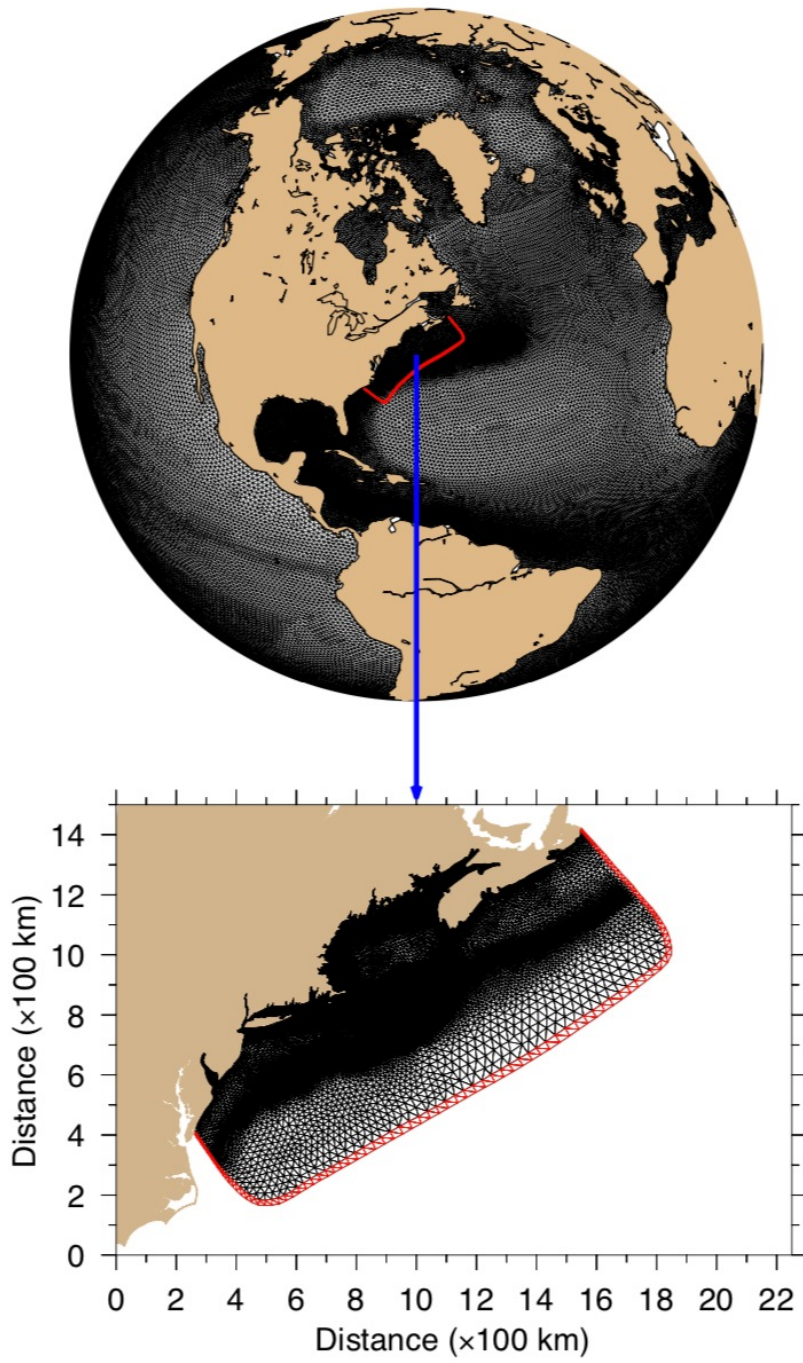


Figure 2

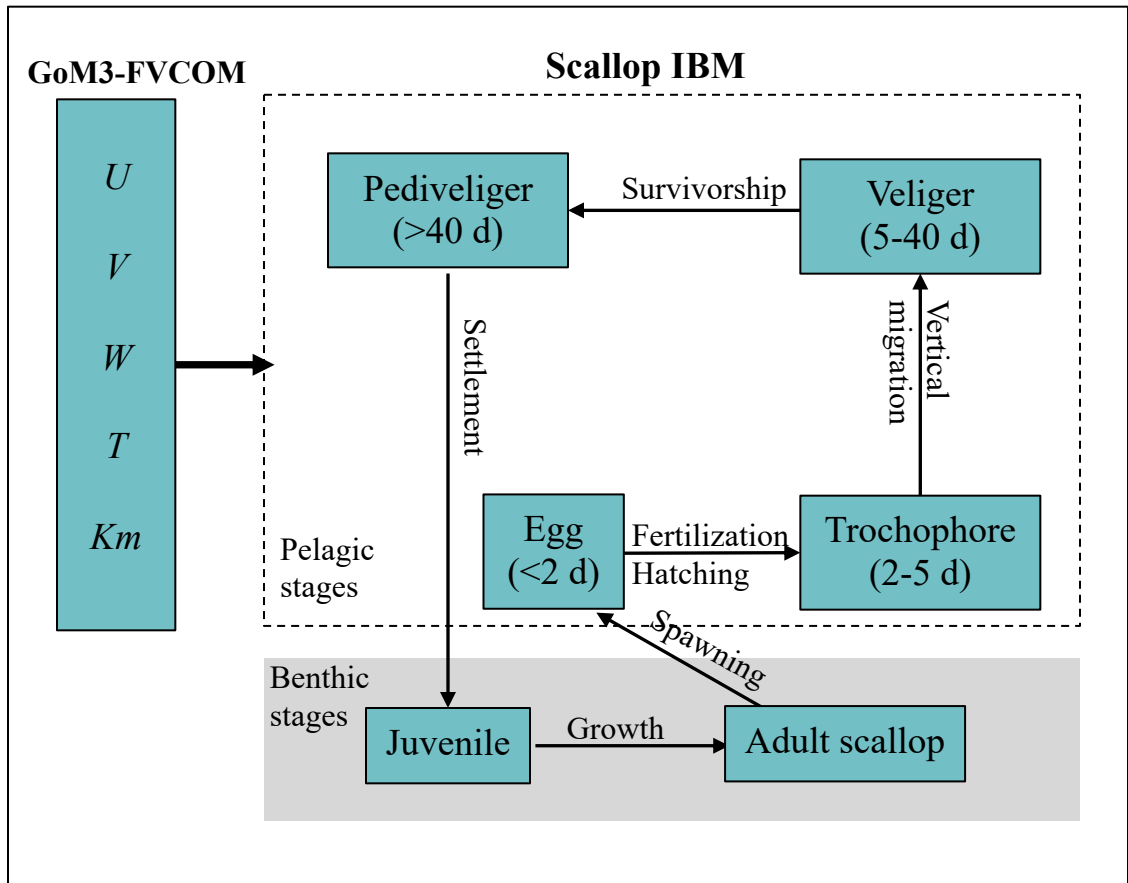


Figure 3

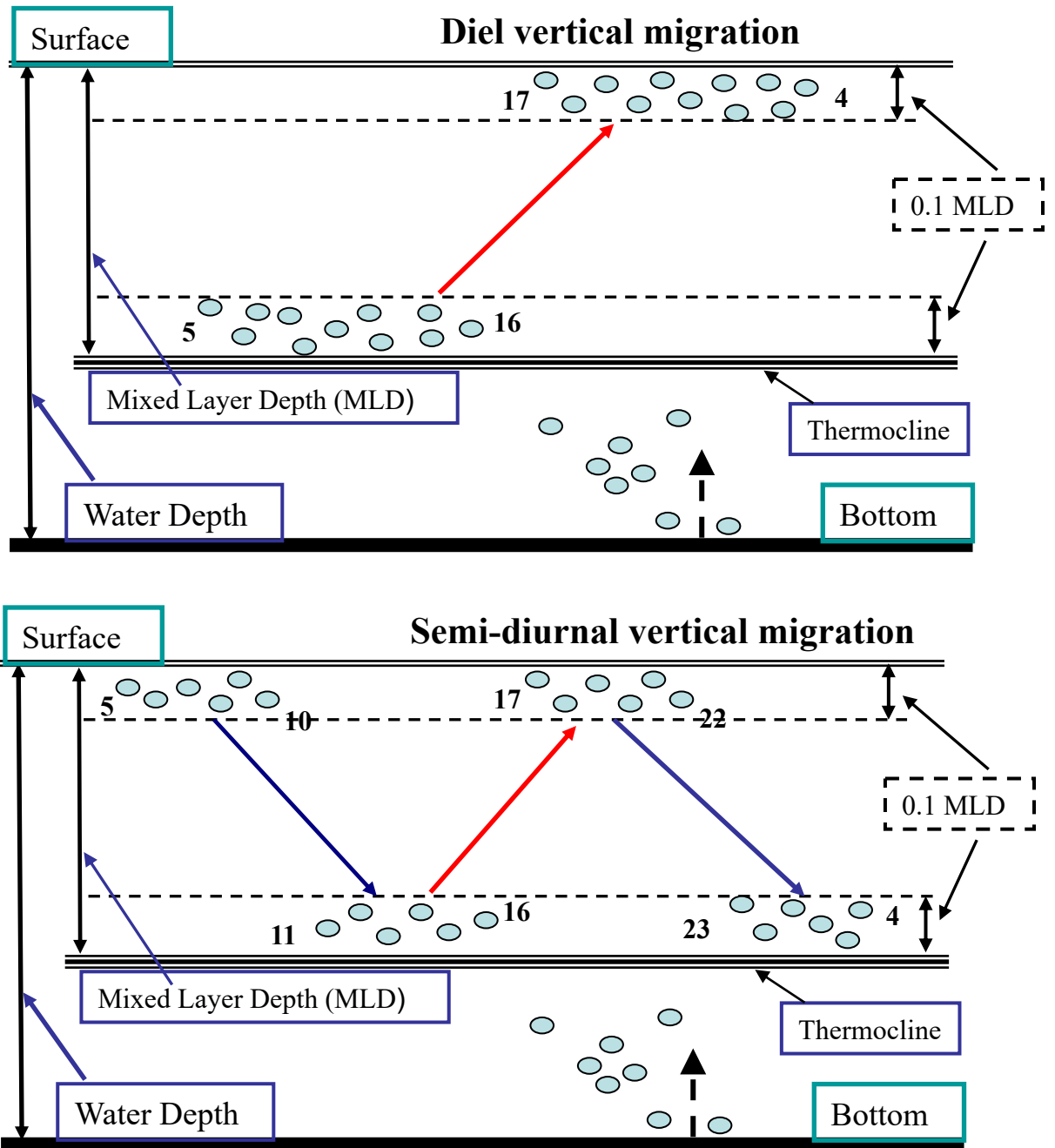


Figure 4

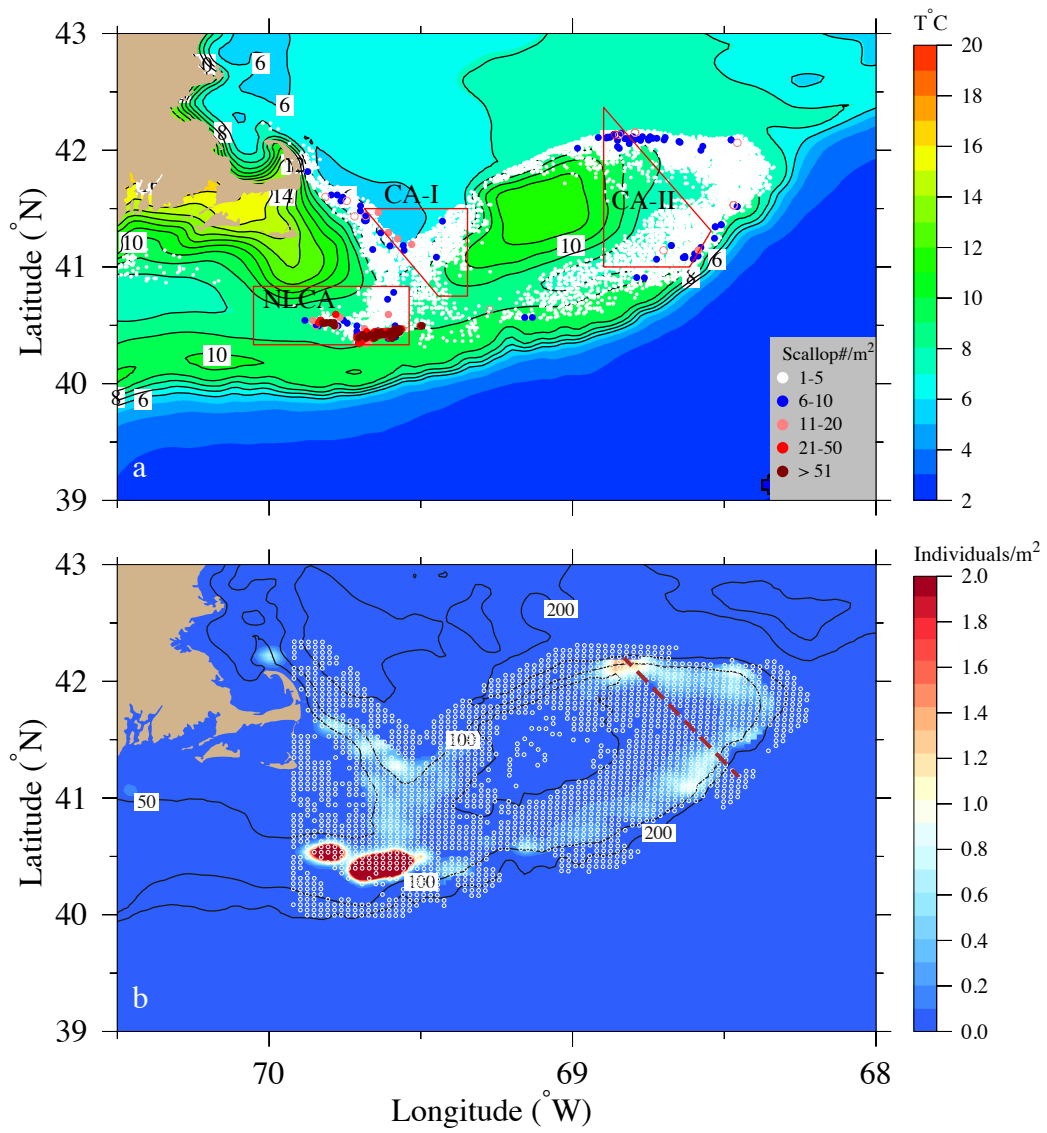


Figure 5

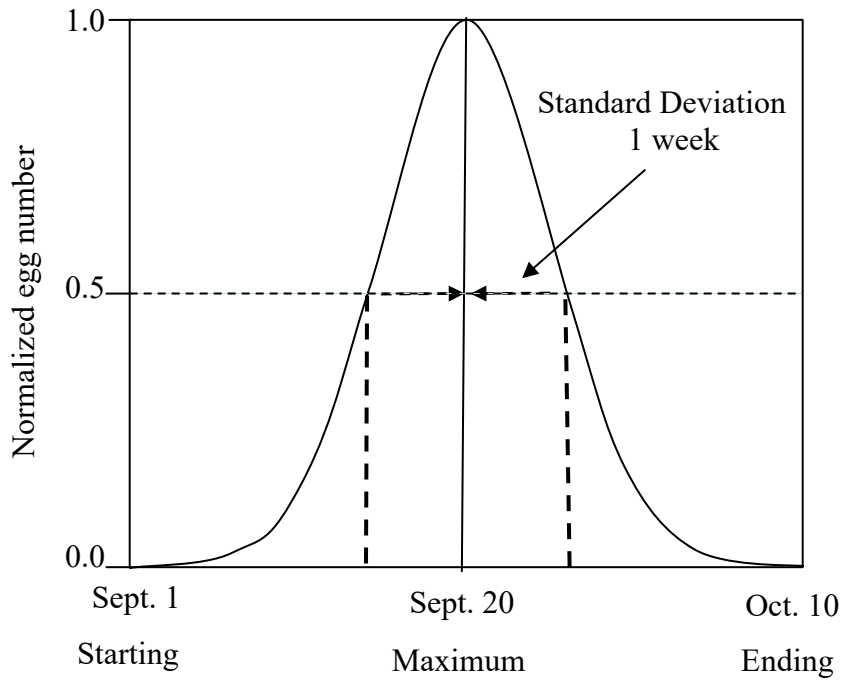


Figure 6

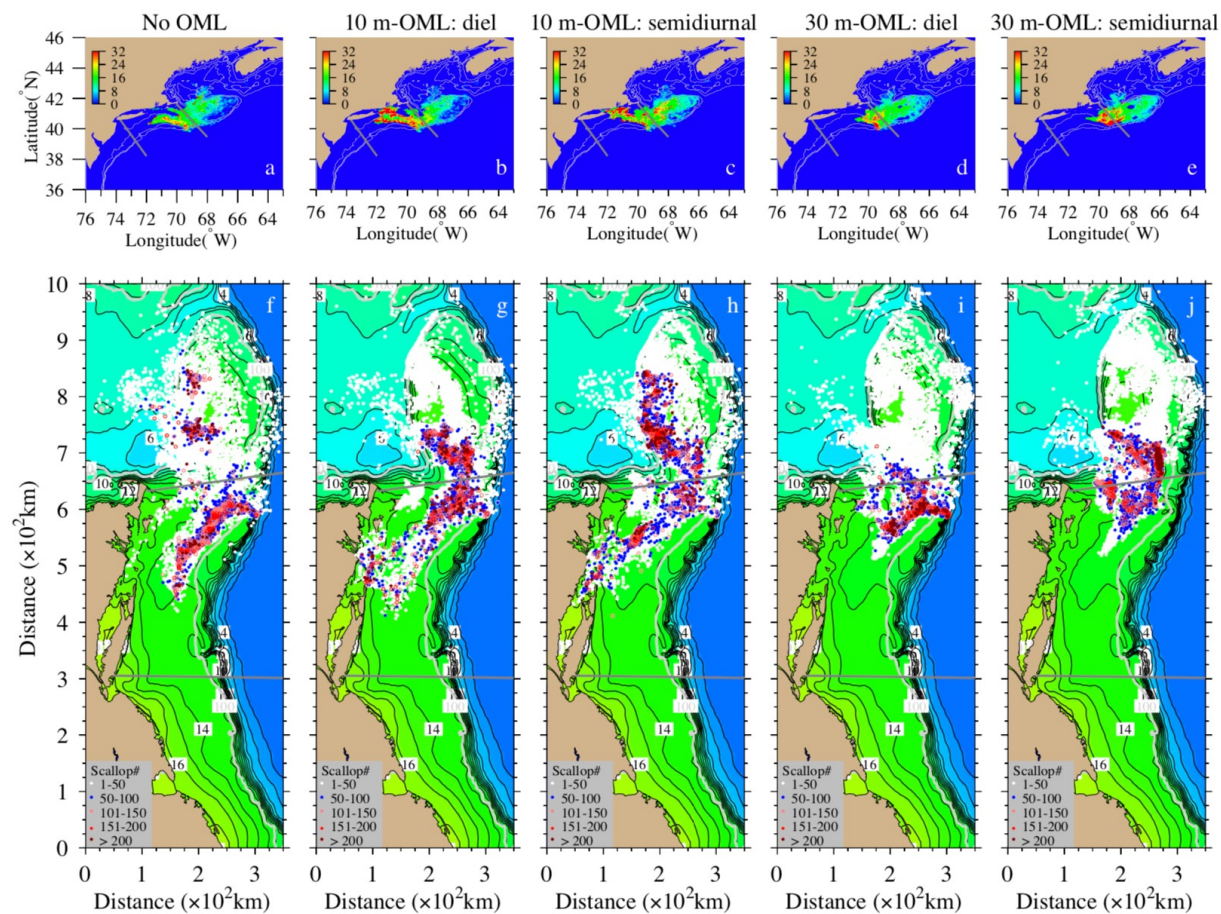


Figure 7

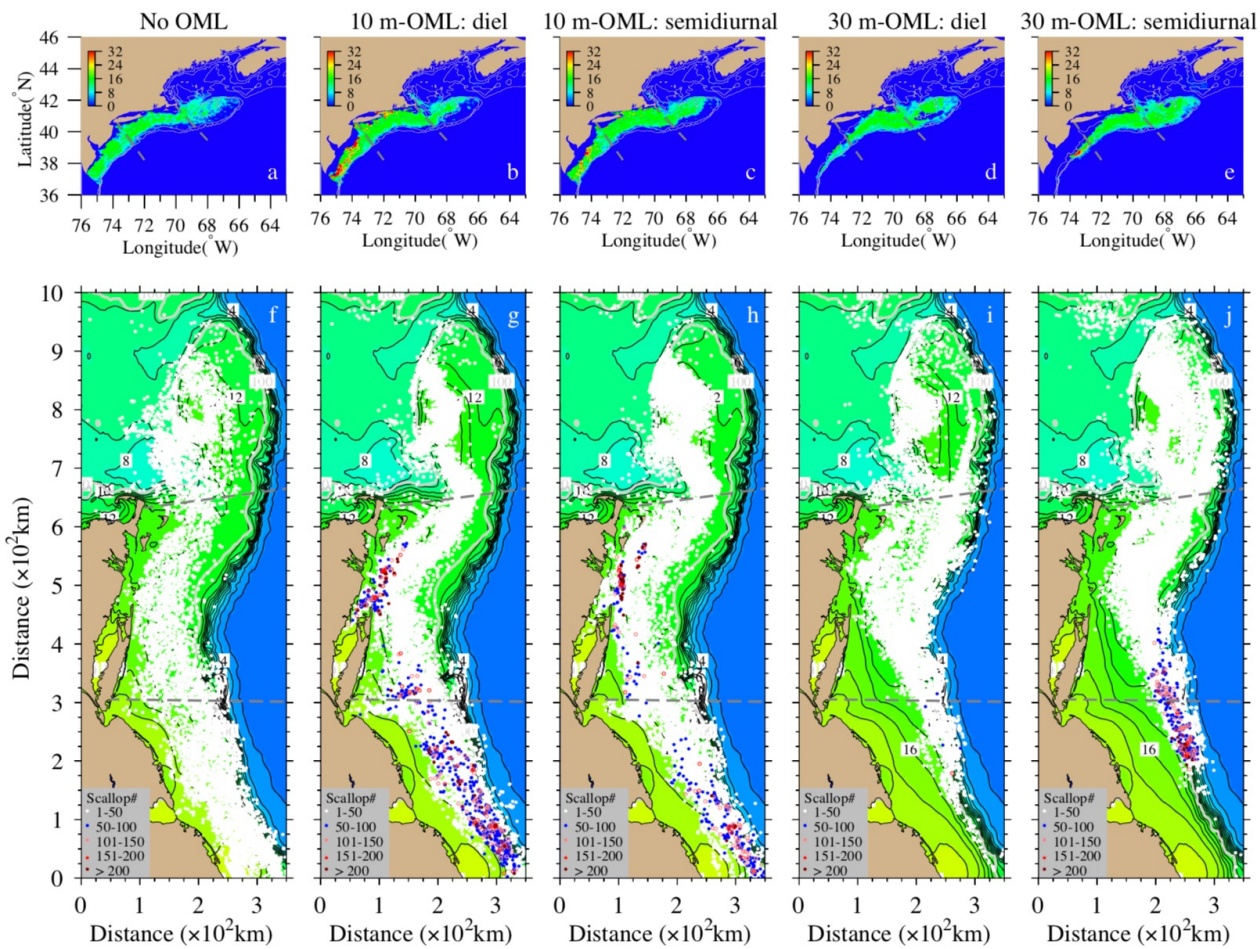


Figure 8

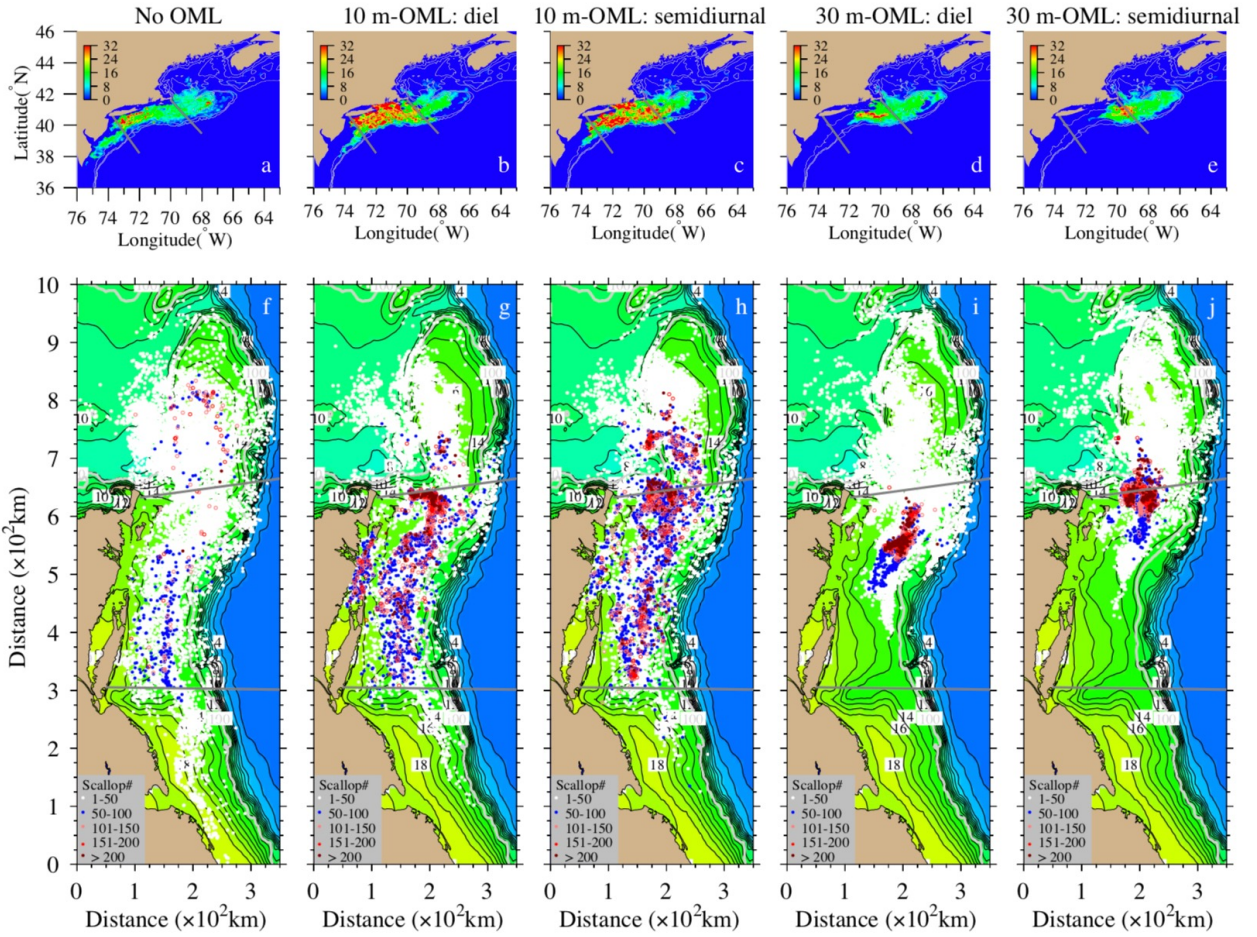


Figure 9

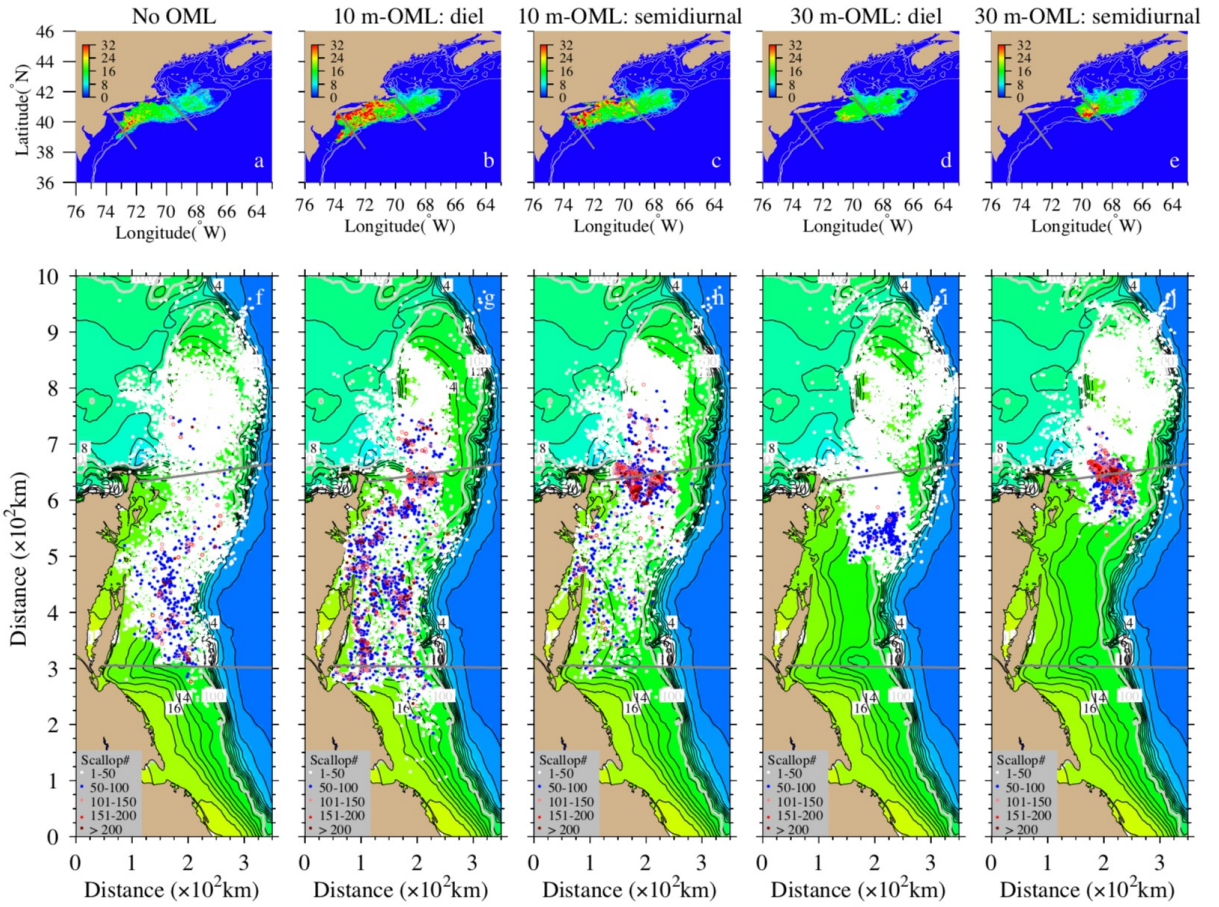


Figure 10

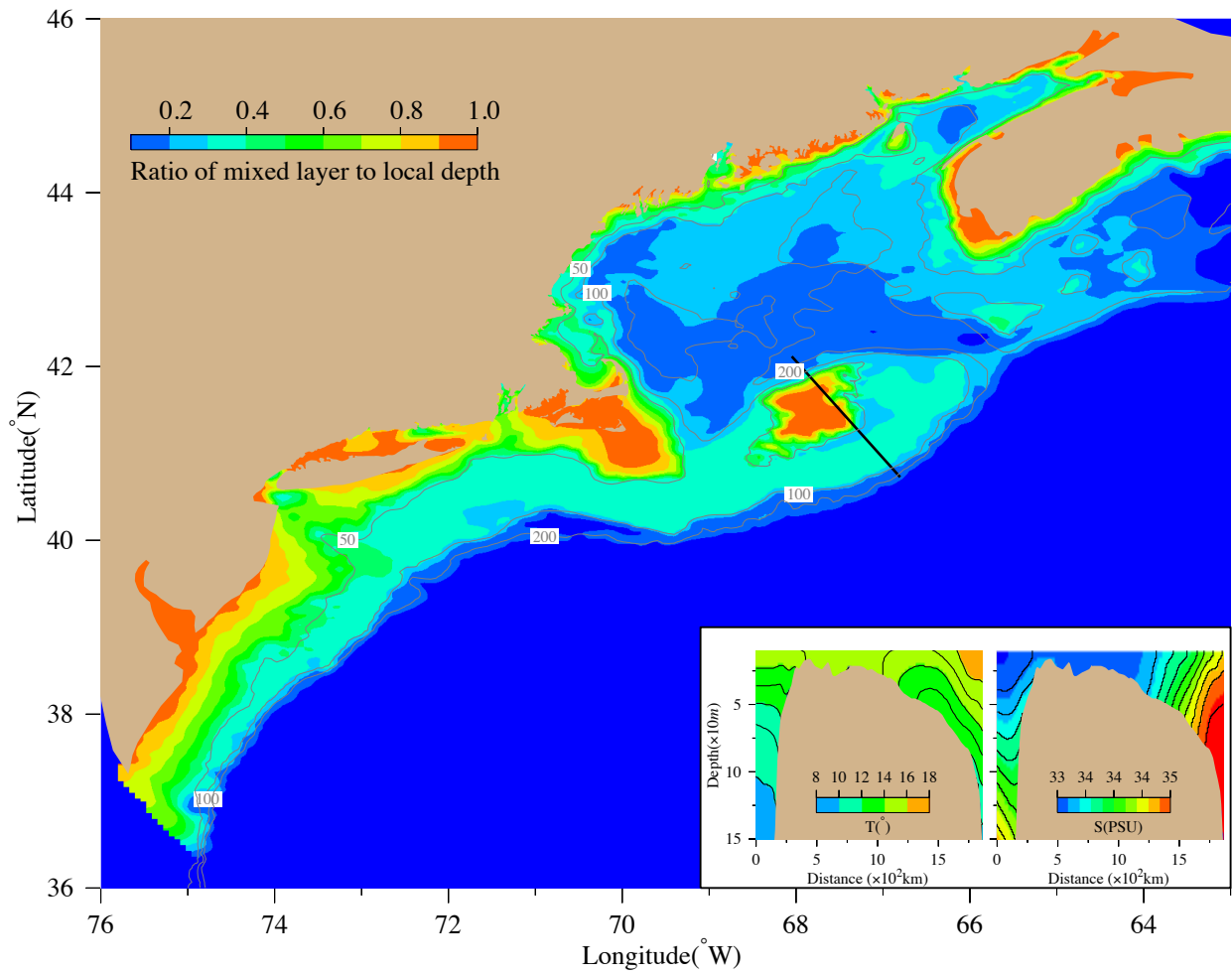


Figure 11

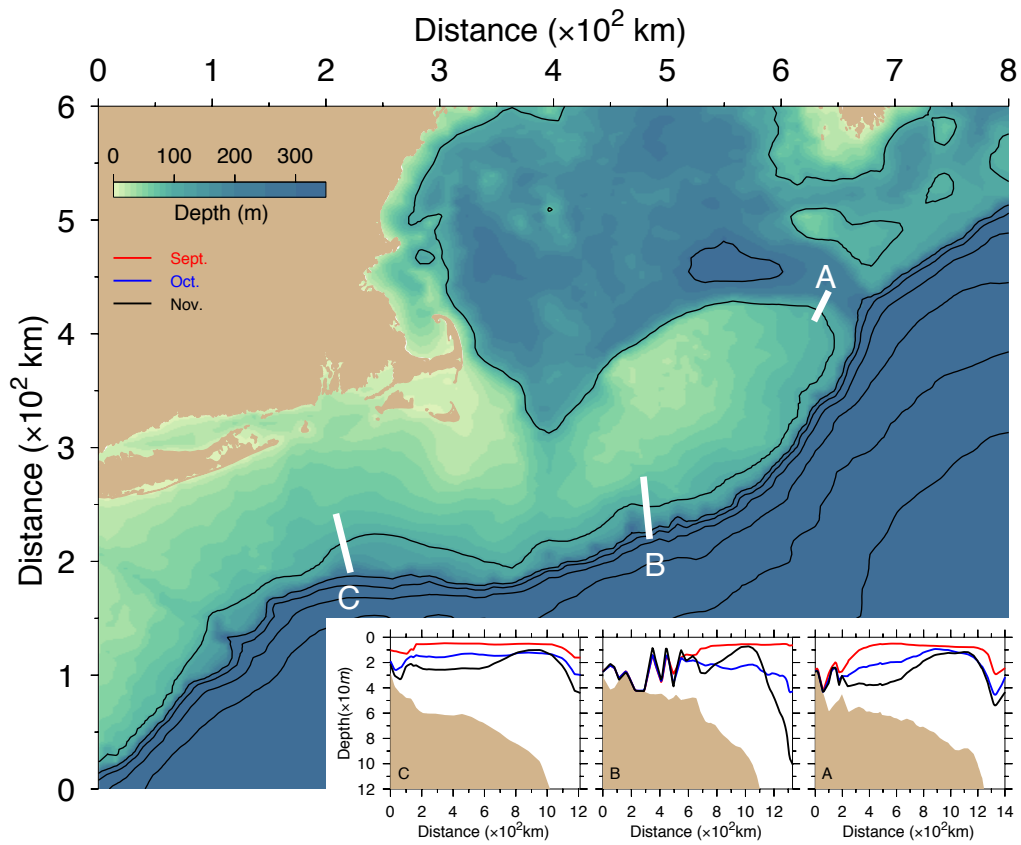


Figure 12

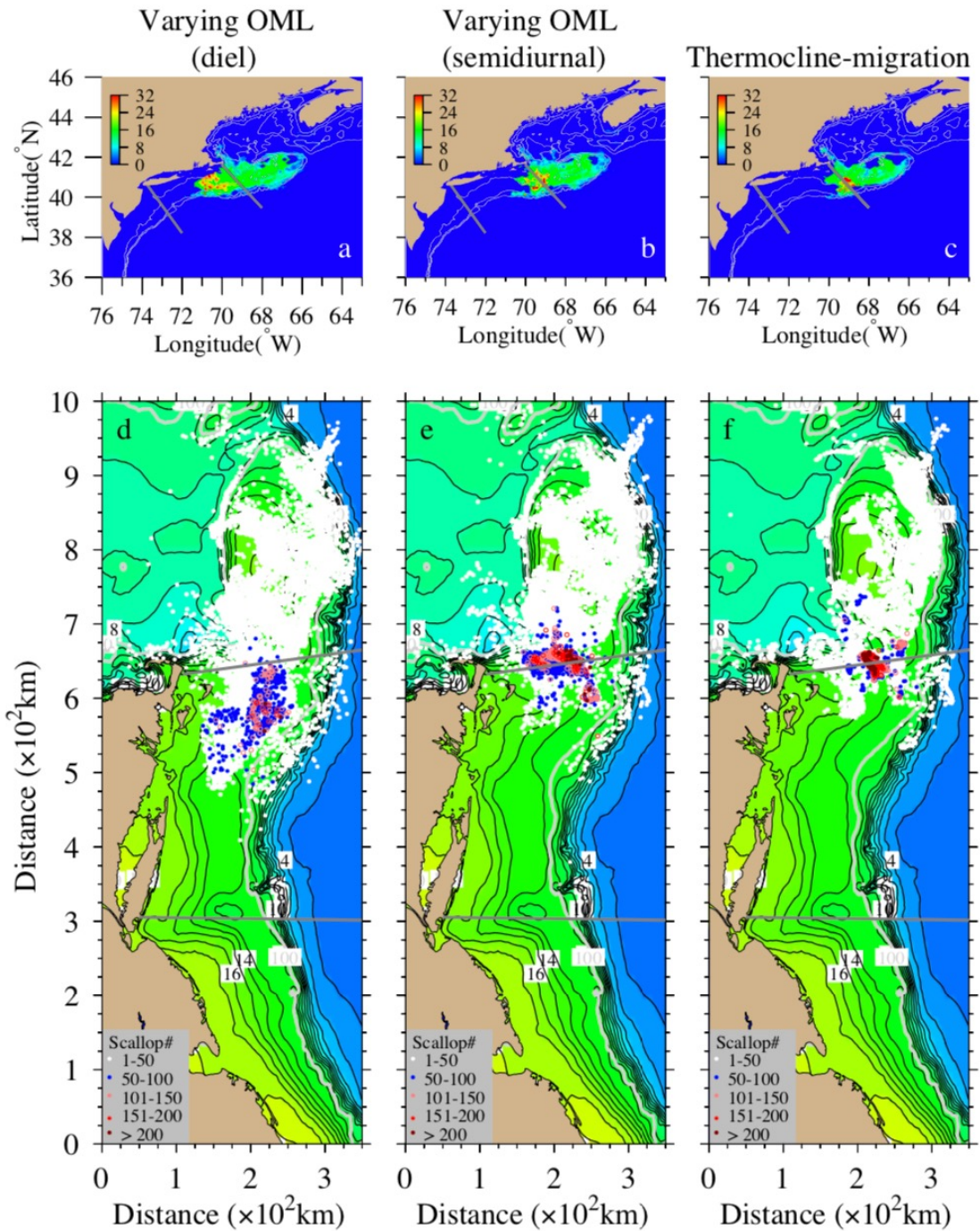


Figure 13

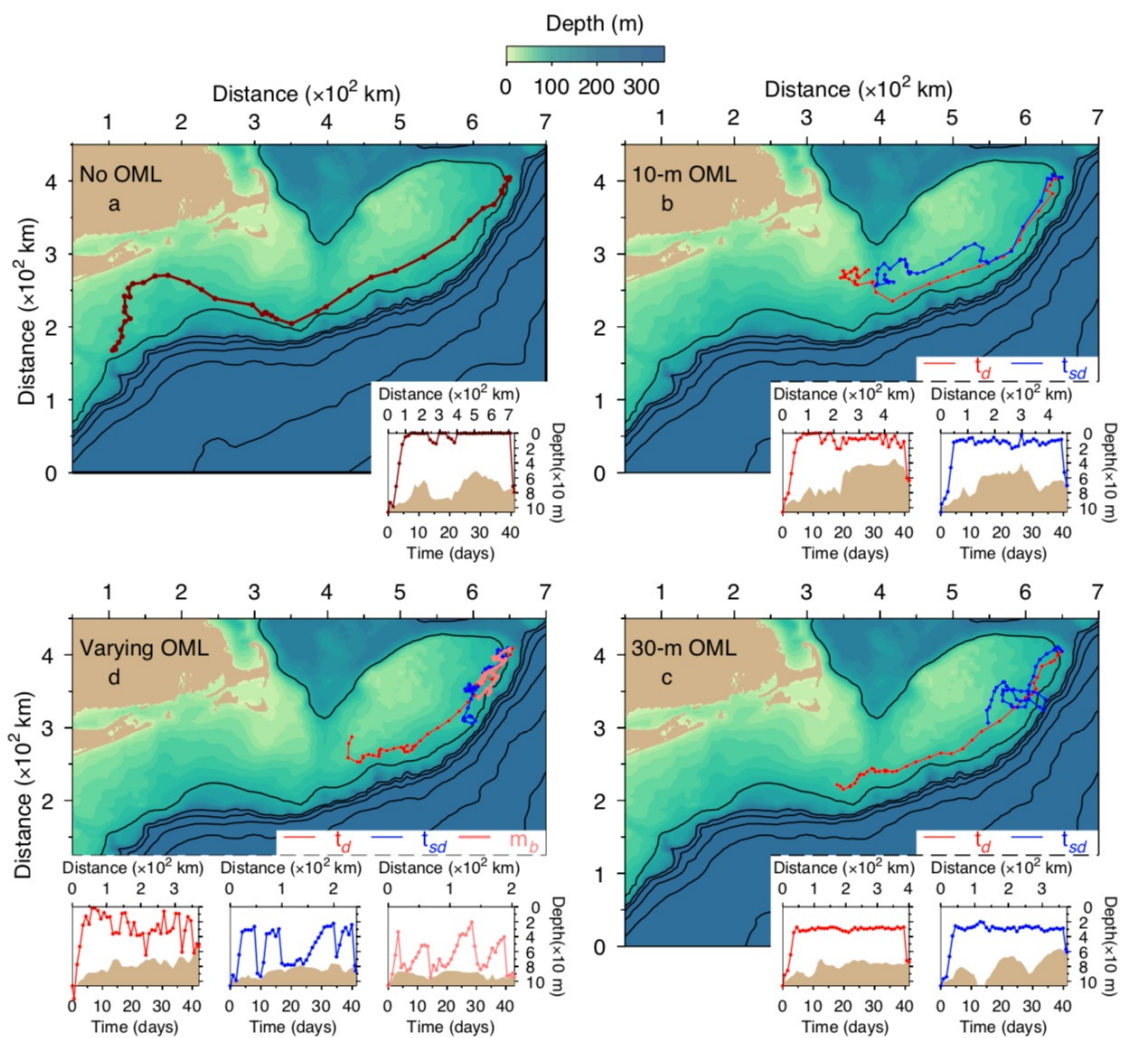


Figure 14

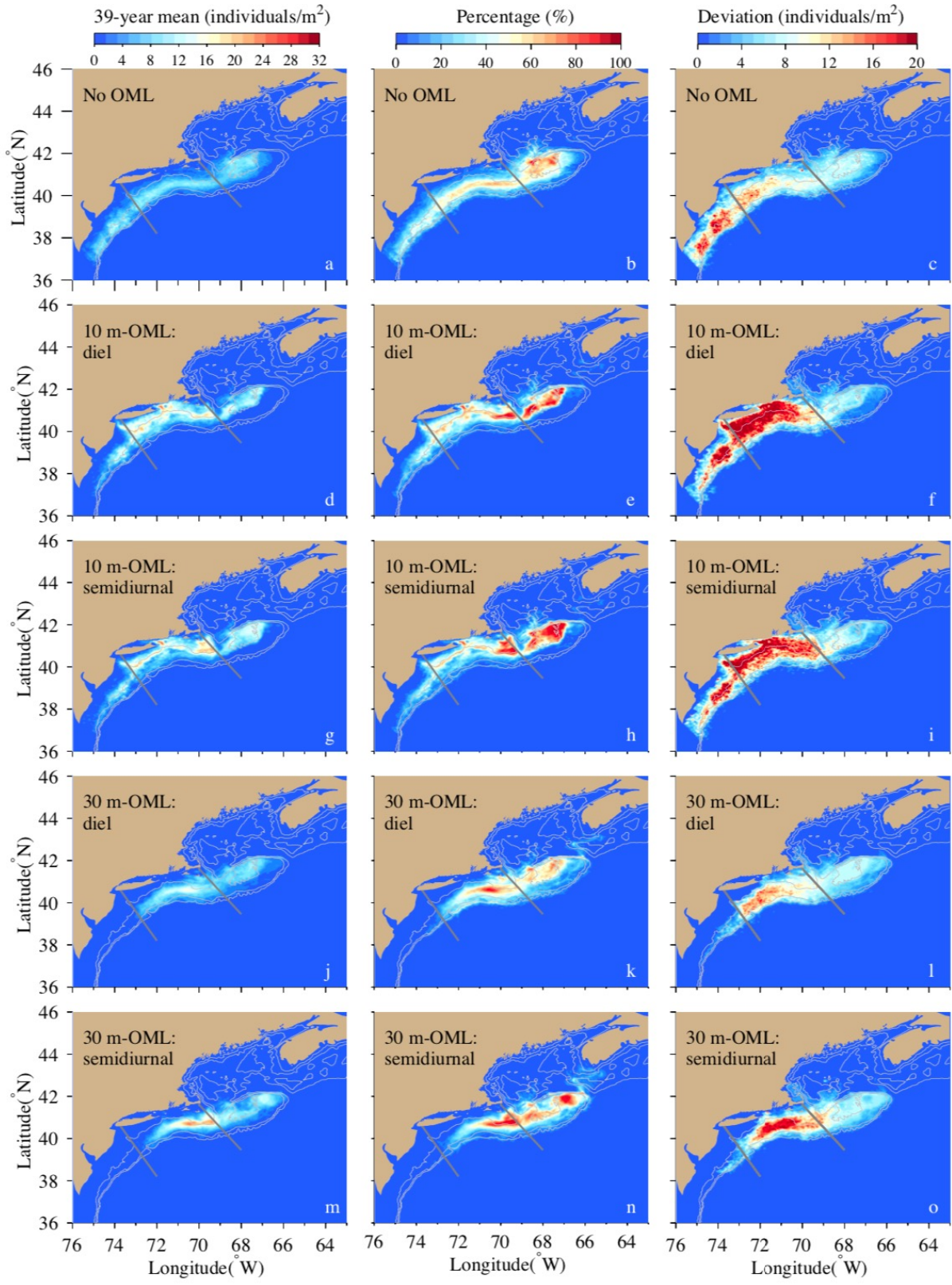


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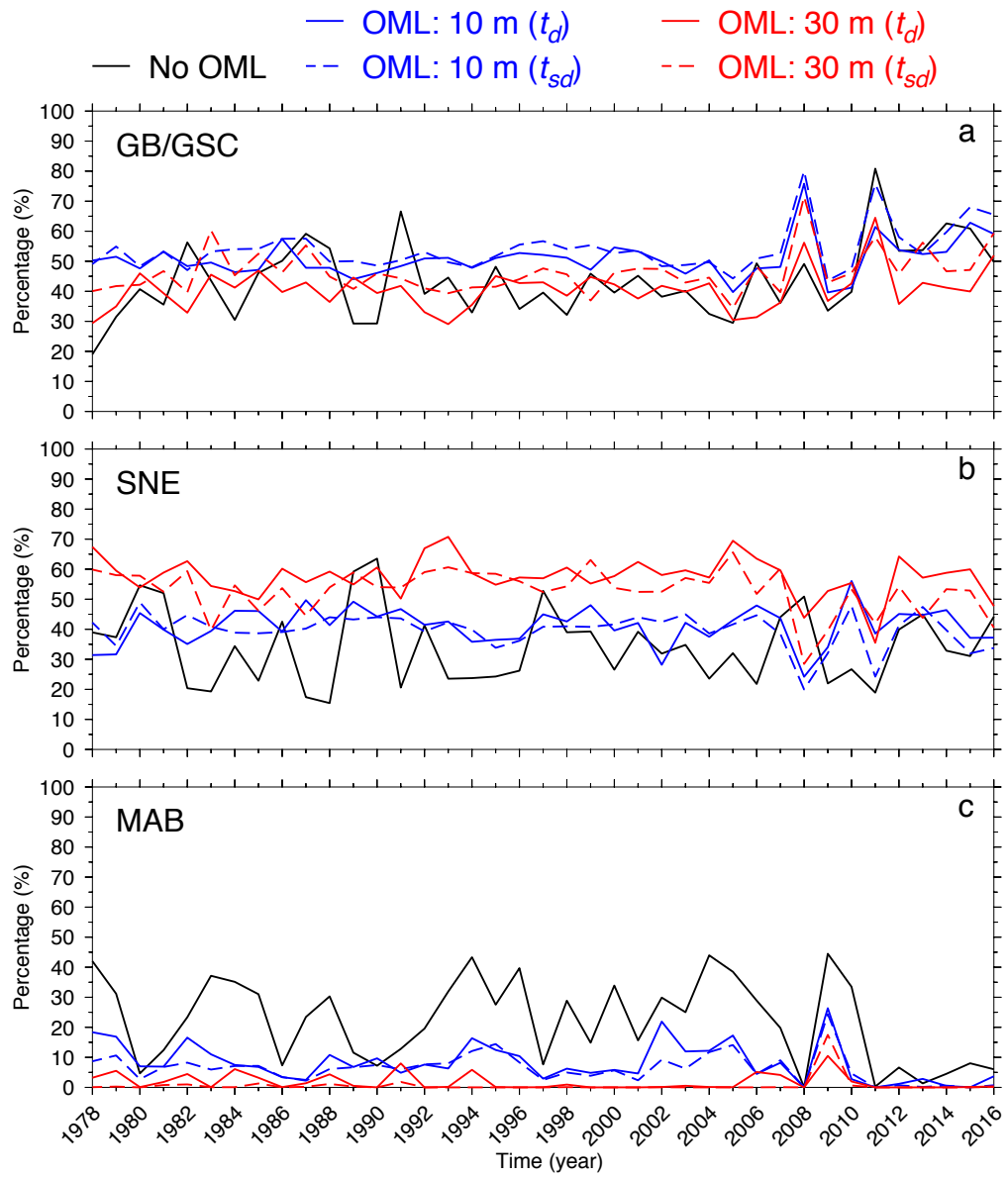


Figure 16

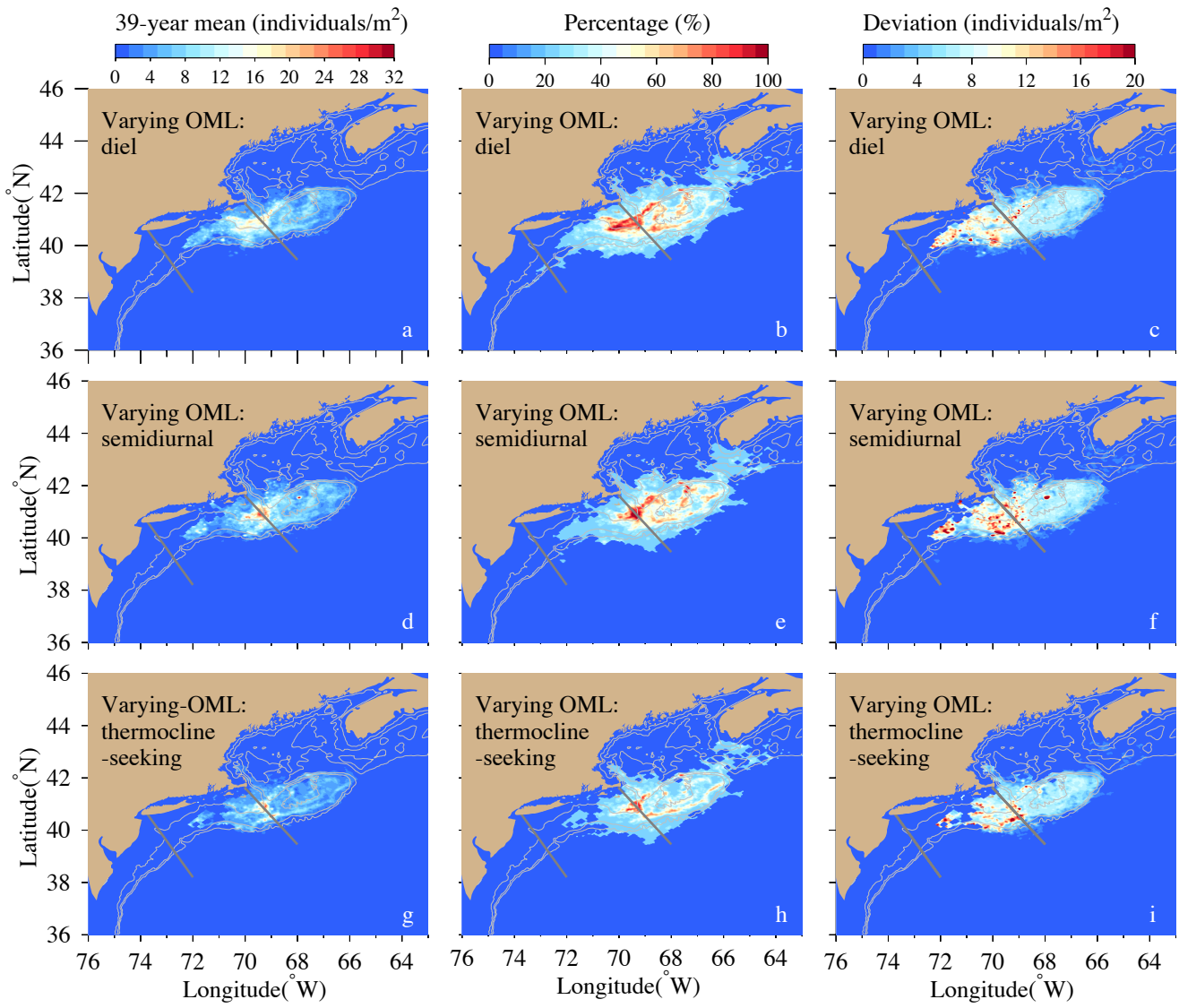


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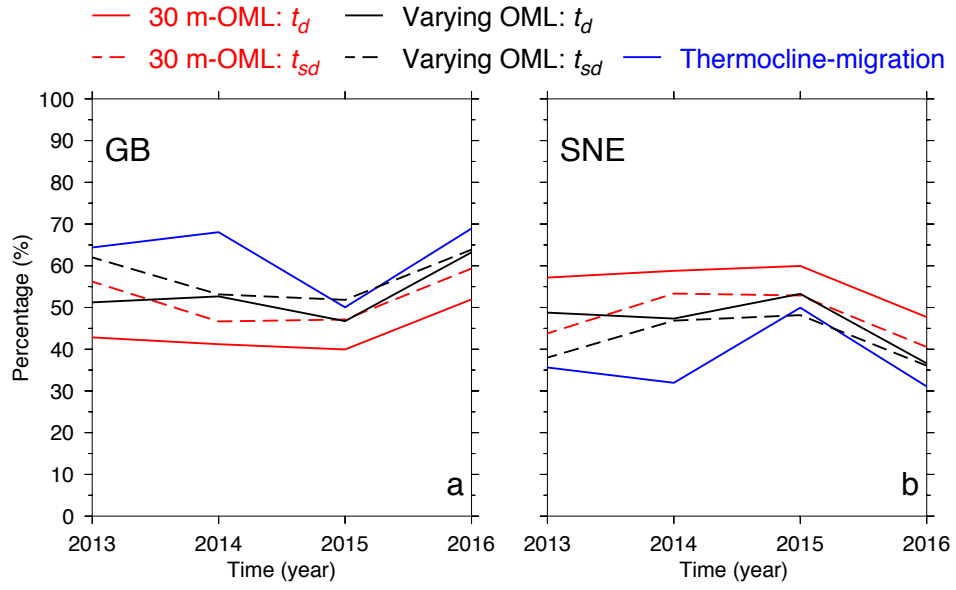


Figure 18

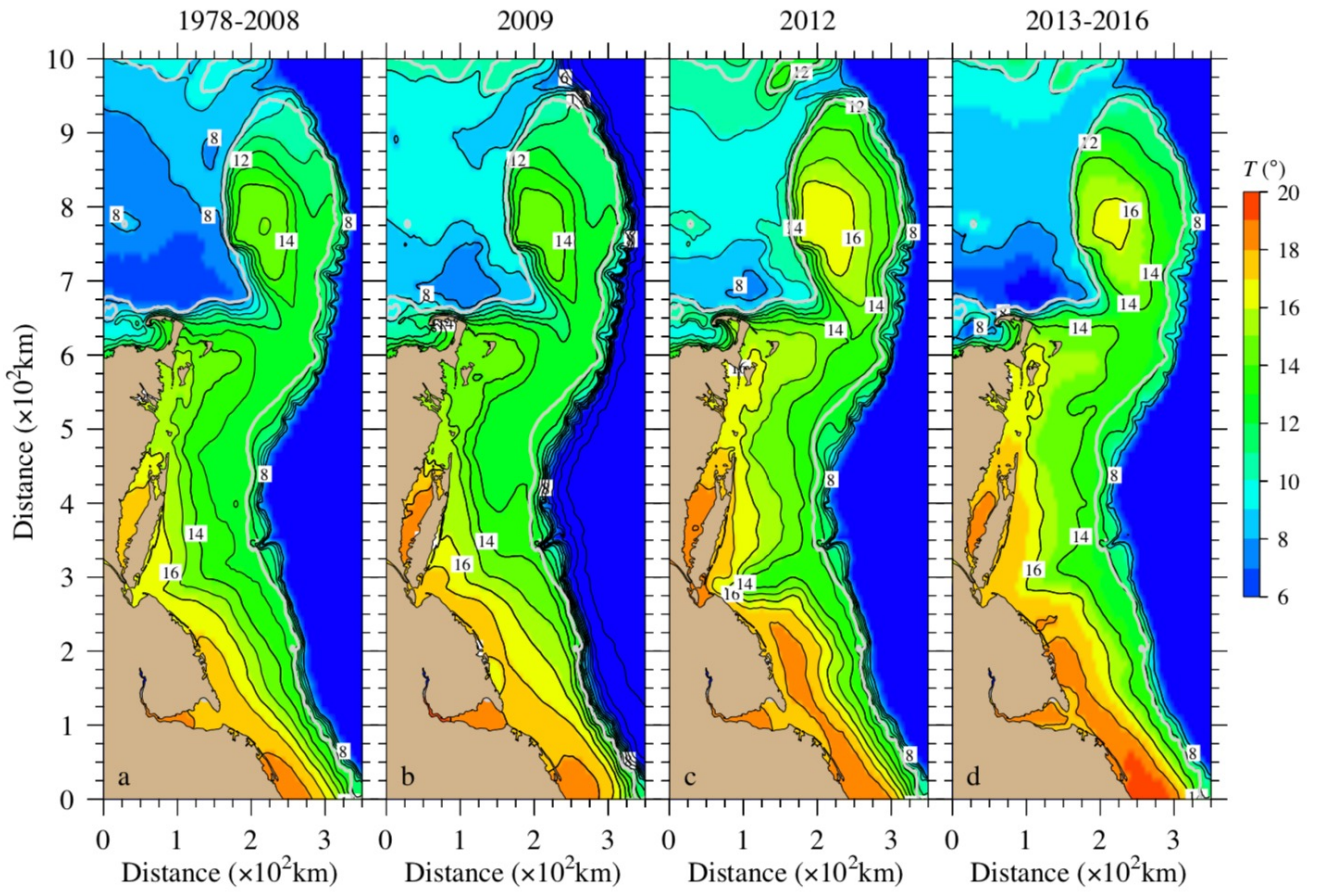


Figure 19

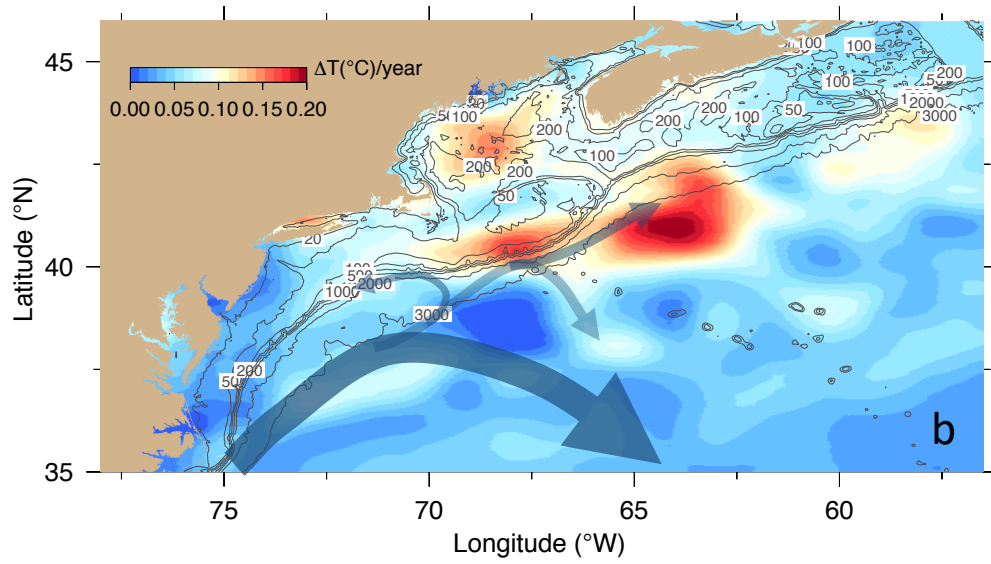
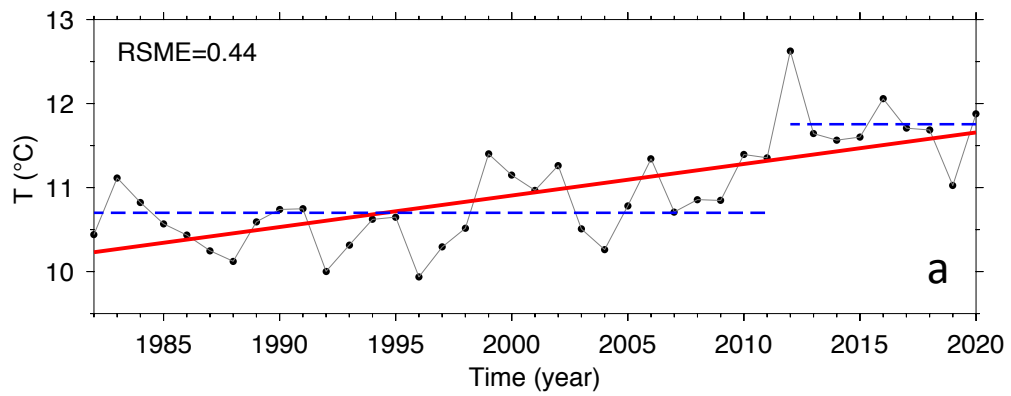


Figure 20

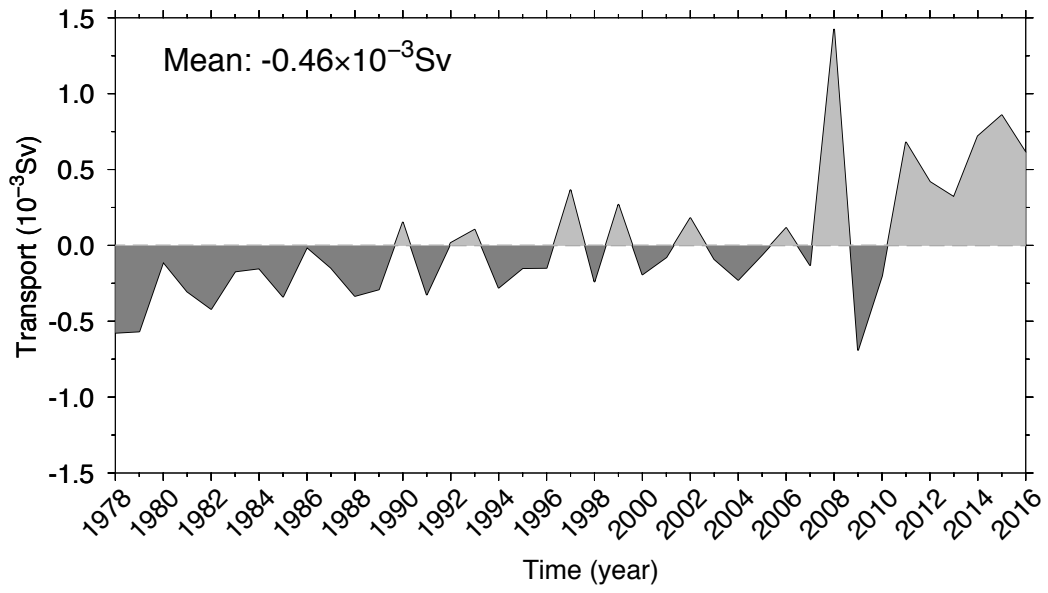


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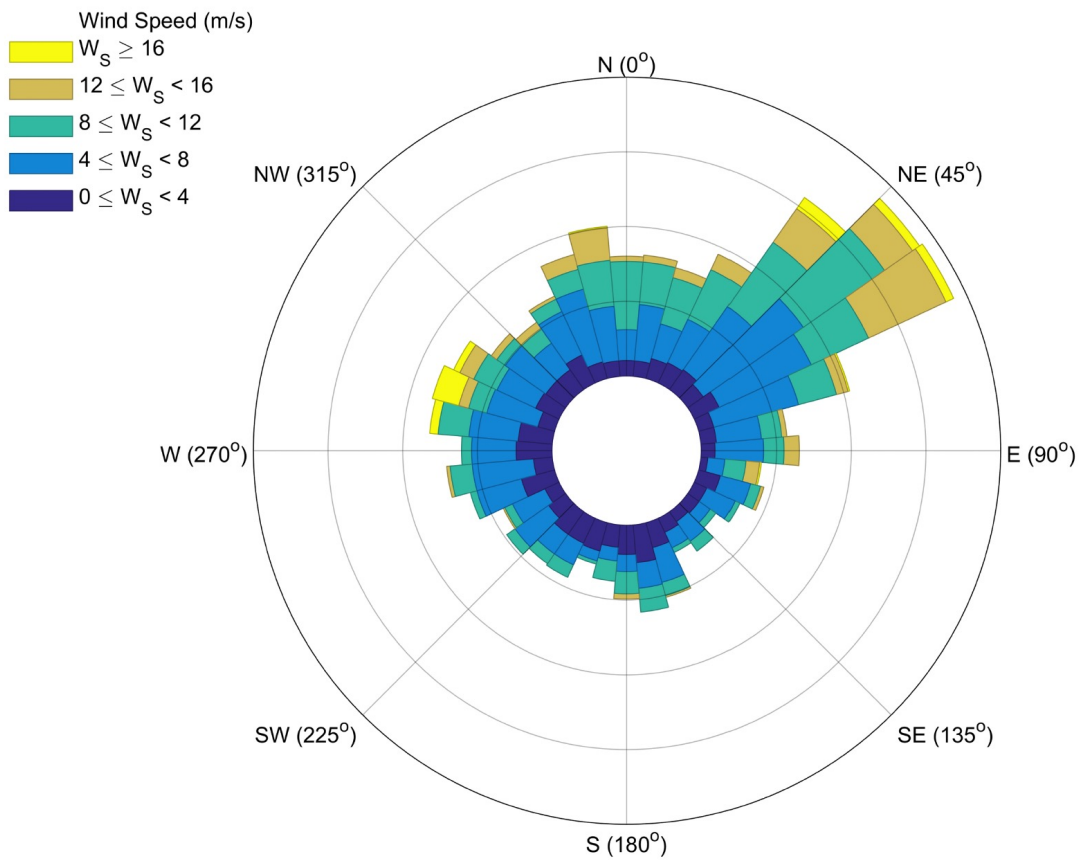


Figure 22

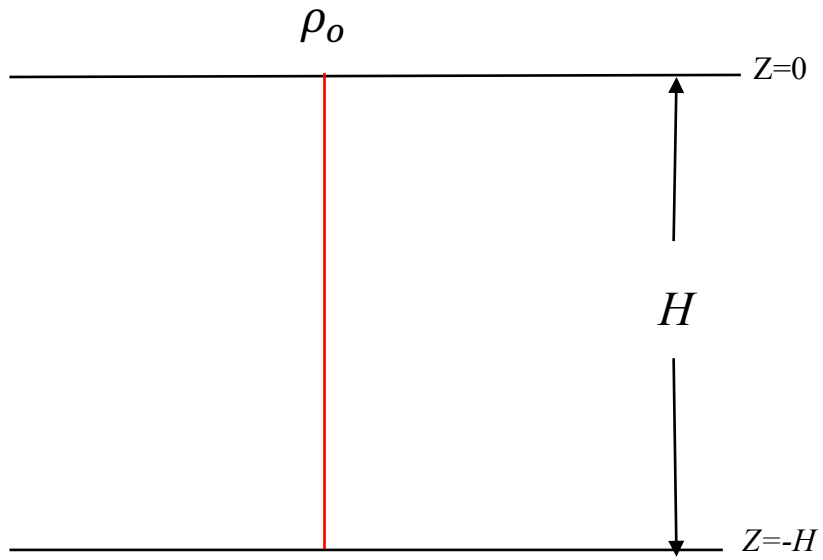


Figure A1

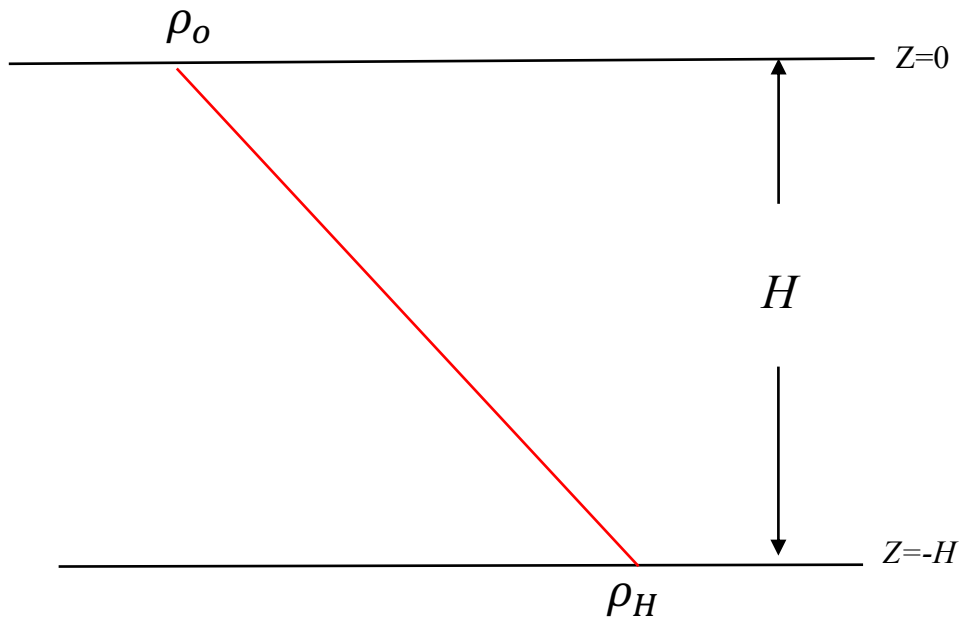


Figure A2

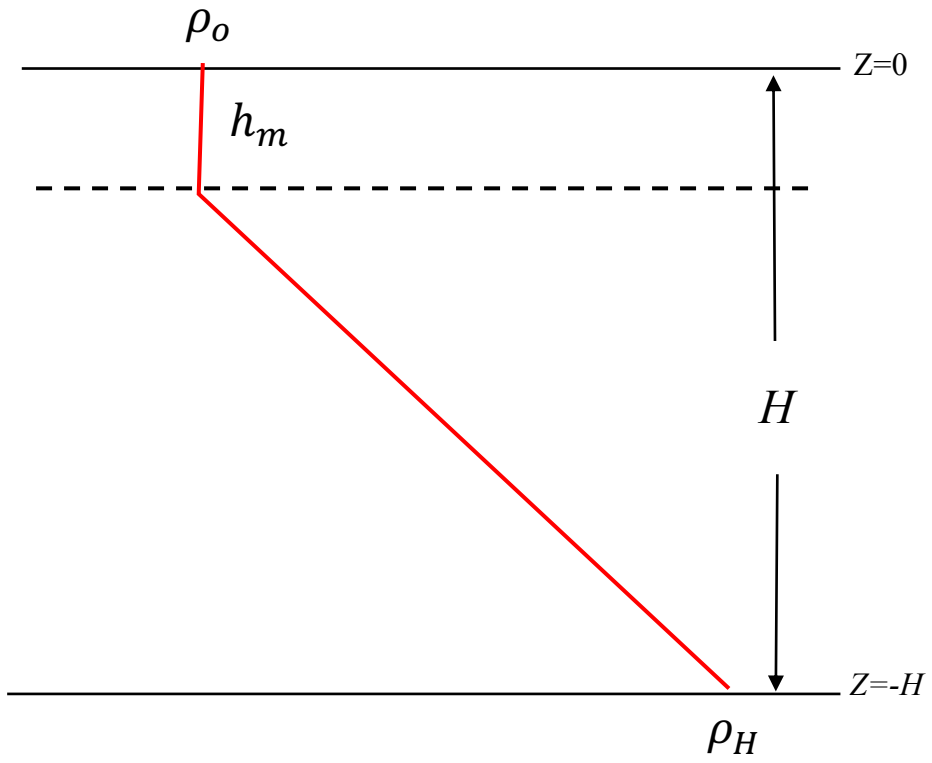


Figure A3

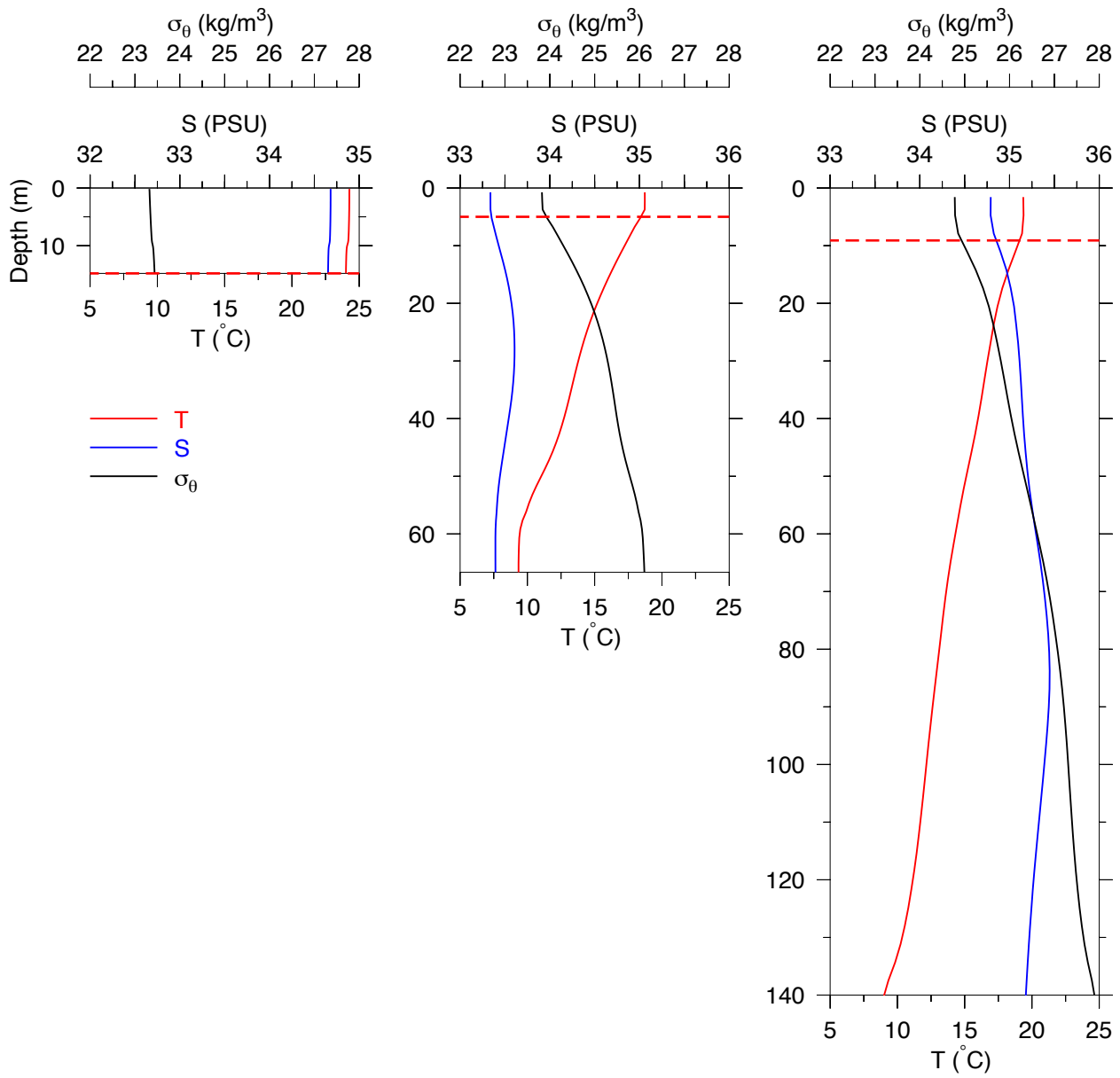


Figure A4