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#### **Influence of Suspended Sediment Front on Nutrients and Phytoplankton Dynamics** 1 off the Changjiang Estuary: A FVCOM-ERSEM Coupled Model Experiment 2 3 Jianzhong Ge<sup>1, 2\*</sup>, Ricardo Torres<sup>3</sup>, Changsheng Chen<sup>4</sup>, Jie Liu<sup>5</sup>, Yi Xu<sup>1</sup>, Richard 4 Bellerby<sup>1,6</sup>, Fang Shen<sup>1</sup>, Jorn Bruggeman<sup>3</sup>, Pingxing Ding<sup>1</sup> 5 <sup>1</sup>State Key Laboratory of Estuarine and Coastal Research, East China Normal University, 6 Shanghai, China, 200062 7 <sup>2</sup>Institute of Eco-Chongming, No.20 Cuiniao Road, Chenjiazhen, Shanghai, China, 202162 8 9 <sup>3</sup>Marine Ecosystems Models & Predictions, Plymouth Marine Laboratory, Prospect Place, The Hoe, Plymouth, PL1 3DH, UK 10 11 <sup>4</sup>School for Marine Science and Technology, University of Massachusetts-Dartmouth, New Bedford, MA 02744, United States 12 13 <sup>5</sup>Depeartment of Biological Sciences, University of Bergen, Postbox 7803, N- 5020, Bergen, Norway 14 <sup>6</sup>Norwegian Institute for Water Research, Thormølensgate 53D, N-5006, Bergen, Norway 15 16

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### 22 Abstract

High-turbidity water is a common feature in the estuary and inner shelf. Sediment suspension
 functions as a modulator that directly influences the interactions among nutrients, phytoplankton

and other related ecosystem variables. A physical-biological coupling model system was applied

- to examine the impact of sediment front on interactions among on suspended sediment, vertical
- 27 mixing, nutrients and phytoplankton over the inner shelf off the high-turbidity, phosphate-limited
- 28 Changjiang Estuary. The physical model was the Finite-Volume Community Ocean Model
- 29 (FVCOM) and the biological model was the European Regional Seas Ecosystem Model
- 30 (ERSEM). Results revealed that in the nearshore region the growth of phytoplankton over the
- 31 spring-summer seasons was limited by suspended sediments and intensified vertical mixing
- 32 during the autumn-winter seasons extended the sediment-induced suppression extended offshore
- to restrict the phytoplankton growth over the shelf. Nutrients were diluted by spreading of
- 34 freshwater discharge and significantly decreased off the suspended sediment front due to the 35 depletion by the offshore phytoplankton growth. The simulation results showed that although the
- diatom phytoplankton dominated the Chlorophyll a (*Chl-a*) concentration, the non-diatom group
- had a more contribution to the biomass. The relatively high phytoplankton biomass was found
- over the offshore deep underwater valley area as results of remote advection by the Taiwan
- 39 Warm Current and weak turbulent mixing.
- 40

# 41 **1 Introduction**

Estuaries are highly productive system in which physical, biogeochemical and ecological 42 processes strongly interact. Dissolved and particulate matters (organic and inorganic), including 43 nitrogen (N), phosphorus (P), silicon (Si) and carbon (C), enter the estuary through upstream 44 rivers, point discharges from human wastewater, and non-point fluxes from the land along the 45 coast as well as submarine groundwaters (Sadat-Noori, et al., 2016; Moore, 2006, 2010). In an 46 estuary, the growth of phytoplankton is controlled by the abiotic and biotic components in the 47 nutrient cycles of the ecosystem (Klausmeier et al., 2008). Nutrient loadings from various 48 sources make an estuary become a nutrient-rich ecosystem, where abundant supplies of nutrients 49 50 greatly impact biological productivity.

Suspended sediment is a common feature in estuaries. In a high-turbidity region, 51 52 sediment resuspension has a great impact on nutrient cycles and phytoplankton community (Kang et al., 2013; Franklin et al., 2018). On the biological level, resuspension activates the 53 nutrient exchange between the water column and bottom benthic layer, and hence enhances the 54 non-diatom plankton growth in the water column (Chen et al, 2004; Boynton et al., 2018). On a 55 physical level, suspended particulate sediments reduce light penetration in the water column. In a 56 high-turbidity environment, the phytoplankton growth is light limited. Increasing light 57 attenuation coefficient limit the photosynthesis process and thus primary production in a 58 nutrient-rich region (Ji et al., 2002; Chen et al., 2004; Vanderploeg et al., 2007; Donohue and 59 Molinos, 2009; Sobolev et al., 2009; Huettel et al., 2014; Hu, et al., 2016; He et al., 2017; 60 Niemistö, et al., 2018). 61

Previous observational and modeling studies have already confirmed the inhibition of
suspended sediment on phytoplankton growth in the suspended sediment plume in the
Changjiang Estuary (Zhu et al., 2009; Wang et al., 2019). In this estuary, however, a sediment

plume usually co-occurs with a dissolved nutrient plume as well as a low-salinity plume (Chen et 65 al. 2003b; Ge et al, 2015). The sediment and dissolved nutrient plumes are located near the river 66 mouth, whereas the low-salinity plume extends offshore over the shelf. Driven mainly by 67 sediment loading and freshwater discharge in the upstream river, the sediment plume exhibits 68 large temporal and spatial changes (Ge et al., 2015, 2018), so that the transition between 69 suspended sediment/dissolved nutrient and low-salinity plumes varies significantly with time and 70 in space. In this multi-plume estuarine system, the plankton dynamics are much complex, which 71 are manifested through the fully nonlinear interaction of physical, biological and chemical processes 72 under varying suspended sediment environment (May et al., 2003; Barbosa et al., 2010; Shi et al., 73 74 2017). Field campaigns with few sampling sites and over a short time coverage are unable to resolve the fine structures of physical and biogeochemical components, especially within the 75 frontal zones. Some simplified numerical models were developed to simulate the phytoplankton 76 growth and nutrient cycles in turbid estuaries (May et al., 2003; de Swart et al., 2009). These 77 models, however, usually failed to resolve the short-term highly-varying temporal and spatial 78 scales, which integrate to form the mid- and long-term variability of the pelagic planktonic 79 ecosystem. Assessing the impact of sediment plume on the ecosystem in a high-turbidity 80 estuarine environment requires an integrated three-dimensional numerical model capable of 81 comprehensively capturing the interactions between sediment and biogeochemical processes, 82 which was demonstrated by previous modeling studies in Great Lakes (Ji et al., 2002; Chen et 83 84 al., 2004), but was scarce in estuaries. Since sediment resuspension in the high-turbidity estuaries is highly controlled by tidal currents regulating with winds and waves, the leading physical 85 drivers, such as tides, waves and dominant estuarine and costal currents, must be integrated into 86 the model. 87

In this study, we applied a fully physical and biogeochemical coupled model to examine 88 89 the impact of sediment plume on the variability of nutrients and phytoplankton in the mega turbid and eutrophic Changjiang (Yangtze) Estuary and inner shelf of the East China Sea (ECS). 90 One novelty of this study is to highlight the development of a realistic, comprehensive 91 92 biogeochemical model system consisting of the interaction among surface waves, sediments, tides, coastal/oceanic currents over finer temporal and spatial scales. The other novelty lies in 93 the exploration of ecosystem responses to high nutrient loading in a high-turbidity estuarine 94 environment. 95

This paper is organized as follows. In Section 2, the study area and massive collection of 96 97 observational data in the Changjiang Estuary and adjacent inner shelf of the ECS are introduced, following with the description of the coupled physical and biogeochemical model. In Section 3, 98 99 the validation results of the model via observational data are presented. In Section 4, the effect of the sediment plume on the offshore nutrient and phytoplankton dynamics is evaluated. In Section 100 5, the dynamics controlling the physical-and-biogeochemical interaction in the Changjiang 101 Estuary and inner shelf of ECS are discussed. Finally, in Section 6, major findings are 102 103 summarized and conclusions are drawn.

- 104 **2. Study area, data and model**
- 105 2.1 Study area

The region of the Changjiang Estuary and inner shelf of the ECS is a typical estuarineshelf coastal zone with a large input of freshwater and sediment from the upstream river (Fig.

- 108 1a). Over the period of 1950-2010, the annually-averaged freshwater volume entering this
- estuary was  $\sim$ 896 km<sup>3</sup>, with a sediment load of  $\sim$ 390 Mt (CWRC, 2011). The freshwater
- discharge and sediment loading were measured at the Datong hydrological station locating ~620
- 111 km upstream from the ECS. This station is considered as a boundary site between the river and 112 estuary. The freshwater discharge varies significantly with seasons. The average freshwater
- estuary. The freshwater discharge varies significantly with seasons. The average freshwater discharge is  $\sim 40 \text{ km}^3$ /s in the wet season and  $\sim 13 \text{ km}^3$ /s in the dry season (Luan et al., 2016).
- discharge is  $\sim 40 \text{ km}^3$ /s in the wet season and  $\sim 13 \text{ km}^3$ /s in the dry season (Luan et al., 2016). Mixing of freshwater with the oceanic water forms a low-salinity plume. This plume changes
- seasonally in terms of its spreading areas and pathway, flowing either into the ECS shelf during
- summer or further away along the Zhejiang coast during winter (Chen et al., 1994; Ge et al.,
- 117 2015b). In summer, the large freshwater discharge makes the low-salinity plume enter the ESC
- shelf region, where it interacts with the large-scale regional oceanic circulations, such as the
- 119 Yellow Sea Coastal Current (YSCC), the intrusion of Taiwan Warm Current (TWC, a sub-
- 120 stream of the Kuroshio Current) and even Kuroshio (Chen et al., 1994 and 2008) (Fig. 1a).



121

Figure 1. The locations of the Changjiang River, the Datong hydrological station and the East China Sea (a). The panel (b) is an enlarged view of the area bounded by the dashed rectangle in panel (a). The pink and red dots indicate the survey sites taken in winter and summer, respectively. The black dots indicate the survey sites taken in both winter and summer.

The abundant sediment loading from the upstream Changjiang River makes this estuary and its adjacent inner shelf of the ECS become a high-turbidity environment. The high-turbidity are clearly appeared in remote sensing images and measured in the field surveys (Siswanto et al., 2011; Sokolowski et al., 2014; Ge et al., 2015). The suspended sediment concentration varied significantly with time and in space, with a range of ~0.1-1.0 g/L near the surface inside the estuary and of < 0.2 g/L in the offshore region (Ge et al., 2015).

In the recent years, with the rapid economic growth and corresponding increase in
 fertilizer use in agriculture, the Changjiang Estuary has experienced a dramatic increase period in

nutrient loading. The environmental monitoring recorded at the Datong station from 1960 to

- 135 2000 indicated a sharp rise in nutrients, with the concentration of dissolved inorganic nitrogen
- (DIN) and phosphate (DIP) increasing from ~20  $\mu$ mol/l to ~130  $\mu$ mol/l, and from ~0.2  $\mu$ mol/l to
- $\sim 1.6 \,\mu$ mol/l, respectively (Li et al., 2007). This evidence was also reported early by Chen et al.
- 138 (2003b) based on other data sources. In contrary to DIN and DIP, however, the dissolved
- inorganic silicate (DSi) dropped from  $\sim 120 \ \mu \text{mol/l}$  to  $\sim 70 \ \mu \text{mol/l}$ , resulting from fixation of DSi within hundreds of reservoirs built along the Changjiang river (Friedl and Wüest, 2002;). These
- 140 within hundreds of reservoirs built along the Changjiang river (Friedr and Wuest, 2002,). These 141 nutrient changes have caused severe eutrophication problems in the area and a dramatic change
- in the ratio of N: Si in the Changjiang Estuary and inner shelf (Zhou et al., 2008; Jiang et al.,
- 143 2010; Zhu et al., 2014; Zhou et al., 2017).

Under a nutrient-rich environment, the high primary production and phytoplankton 144 bloom were often observed in the Changjiang Estuary and it adjacent inner shelf of the ECS 145 (Furuya et al., 1996, 2003), with a Chlorophyll-a (*Chl-a*) concentration of  $>10 \,\mu g \, L^{-1}$  off the 146 (Chen et al., 2003b; Zhu et al., 2009) and even reached ~20  $\mu$ g L<sup>-1</sup> when the harmful algal bloom 147 occurred (Chen et al., 2003b). The diatom remains to be a predominant species contributing to 148 the total phytoplankton *Chl-a* (>90%). Recent surveys reported that dinoflagellates have become 149 dominant component in the phytoplankton community, particularly in the high resuspended 150 sediment plume (Chen et al., 2003b) and in eutrophic onshore waters influenced by the low-151

salinity plume (Jiang et al., 2015).

# 153 2.2 The cruise data

Massive physical and biogeochemical data were collected from multiple field campaigns. 154 Observational sites covered the river channel, estuary and inner shelf of the ECS (Fig. 1b). 155 Responding to different patterns of Changjiang River plume's extensions in summer and winter 156 around the inner shelf, 14 more sampling sites were placed in the south of the Changjiang 157 Estuary during winter plus additional 12 sites in the east of the inner shelf in summer (Fig. 1b). A 158 total of 90 and 92 sites were visited during the summer and winter cruises, respectively. Physical 159 and biogeochemical variables were measured at these sites. The physical variables contained 160 water velocities from a vessel-mounted Acoustic Doppler Current Profiler (ADCP), temperature 161 and salinity from sensors of Conductivity-Temperature-Depth (CTD). The biogeochemical 162 variables included pH, nutrients including DIN, DIP, ammonia (NH4), and DSi, Chl-a, and 163 suspended particulate matter (SPM). Cruises periods covered every winter and summer over the 164 period of 2015 to 2017. 165

# 166 2.3 The satellite data

Both surface sediment concentration and phytoplankton *Chl-a* can be calculated from 167 satellite remote sensing data (O'Reilly et al., 1998; Shen et al., 2010). In this study the satellite 168 data were used for model validation and interpretation. The data came from two sources. One 169 170 was the 4-km-resolution-mapped daily Chl-a field from Moderate Resolution Imaging Spectroradiometer Aqua satellite (MODIS-Aqua, https://oceancolor.gsfc.nasa.gov/data/aqua/), 171 which covered the period from January 1 2005 to December 31 2016. The other one was the 500-172 m-resolution-mapped hourly sky-view observational data from the Multi-channel Geostationary 173 Ocean Color Imager (GOCI) satellite, which covered the East Asian seas and recorded every 8 174 hours per day (Cho et al., 2012, 2014). Surface Chl-a concentrations were retrieved and 175

calculated using the ocean chlorophyll 2 algorithm (OC2) (O'Reilly et al., 1998; Ryu et al., 2012)

through the GOCI Data Processing System (GDPS version 1.4.1) from 2012 to 2016, in which

the Korea Ocean Satellite Center (KOSC) standard was applied in atmospheric correction. For

public access, the daily product contained three snapshots taking at 10:30 am, 11:30 am and
12:30 am, respectively.

Generally, in coastal optically complex waters, *Chl-a* algorithms suffer from the 181 contamination by colored particulate and dissolved substances. Using the semi-empirical 182 radiative transfer (SERT) algorithm with physical based empirical coefficients (Shen et al., 183 2010), the GOCI optical reflectance data were used to estimate surface sediment concentrations 184 in the Changjiang Estuary (Choi et al., 2012; He et al., 2013; Ge et al., 2015b), as well as Chl-a 185 (Choi et al., 2014; Piwowarczyk et al., 2016; Sun et al., 2018). Previous studies have already 186 confirmed the reliability of GOCI production for the Chl-a under the interference of colored 187 dissolved organic matters (Lamquin et al., 2012; Hu et al., 2012). 188

189 2.4 Hydrodynamic model

Main hydrodynamic forcings and components in the Changjiang Estuary include the 190 191 astronomical tide, river discharge, wind, waves, coastal/oceanic currents, and the sediment. The core system of the physical numerical model is the Finite-Volume Community Ocean Model 192 (FVCOM). FVCOM is an unstructured-grid community ocean model, in which governed 193 194 equations were discretized and solved using the finite-volume integration algorithm (Chen et al., 195 2003a) and gradually upgraded to include multiple hydrodynamic and ecosystem modules (Chen et al., 2013). The non-overlapping triangular grid configured in FVCOM provided accurate 196 197 geometrical fitting for irregular coastlines while guaranteeing flexibility for refining the grid in steep bathymetry, islands, or regions of interest (Chen et al, 2003a, Chen et al, 2006, Qi, et al., 198 2018). An accurate finite-volume second-order Ruge-Kutta algorithm guaranteed volume and 199 mass conservations of the momentum fluxes over each iteration. Horizontal diffusion was 200 parameterized based on Smagorinsky's formation (Smagorinsky, 1963) and vertical turbulent 201 mixing was calculated using the 2.5-level Mellor and Yamada turbulence model (Chen et al., 202 2013). 203

FVCOM is a fully current-wave coupled model (Chen et al., 2013). An unstructured-grid version of the surface wave model-SWAN was developed on the platform of FVCOM and named "FVCOM-SWAVE" (Qi et al., 2009). The FVCOM-SWAVE implemented to FVCOM as a wave module and coupled with hydrodynamic component of FVCOM. This coupled model been applied for the coastal wave and inundation applications with inclusion of wave-current interactions (Wu et al., 2010; Ge et al., 2013; Beardsley et al., 2013; Chen et al, 2013; Qi et al., 2018).

FVCOM also has incorporated a sediment module, covering from suspended sediment, to bed-load transport and layered bed-soil dynamics (Chen et al., 2006). This model has the ability of supporting unlimited sediment classes for non-cohesive and cohesive sediment dynamics. Additionally. the wave and sediment have been fully coupled with the hydrodynamic kernel with their physical interactions (Wu et al., 2010; Ge et al., 2018). The two-way coupling was considered in this modeling. This means individual process (such as waves) can impact on the other two processes (e.g. current and sediment).

#### 218 2.5 Biogeochemical Model

The biogeochemistry and ecosystem dynamics are simulated by the European Regional 219 Seas Ecosystem Model (ERSEM ver. 15.06), which is a generic and well established lower-220 trophic level marine food web and biogeochemical cycling model (Butenschön et al., 2016). It 221 resolves the ecosystem dynamics with nutrients and carbon cycles in the low trophic levels. It 222 223 divides the phytoplankton, zooplankton and benthos into function groups, and calculates the biomass for the group individually. The model calculates various state variables, including 224 pelagic and benthic living organisms, dissolved and particulate nutrients, dissolved oxygen and 225 carbonates. ERSEM considers the functional groups following the lower trophic food-web chain 226 from primary producers of phytoplankton, consumers like zooplankton, to decomposers of 227 bacteria. For phytoplankton, ERSEM could consider up to four types, according their group 228 sizes. They include pico-, nano-, and microphytoplankton, and diatoms. ERSEM accounts for up 229 to three types of zooplanktons in the system, from meso-, and microzooplankton and 230 heterotrophic nanoflagellates. For decomposers, only one type of heterotrophic bacteria is 231 considered. Chl-a is determined separately in the model based on the quantification formulation 232 (Geider et al., 1997). ERSEM is also equipped with a comprehensive benthic model (Ebenhoh et 233 al., 1995) and a fully resolved carbonate system. It provides active nutrients and mass exchanges 234 in the water-sediment interface. The ERSEM considers both dissolved and particulate organic 235 236 matter, including labile dissolved organic matter, semi-labile organic matter, semi-refractory organic matter, small particulate organic matter, medium size particulate organic matter, large 237 particulate organic matter. The particulate organic matter for nitrogen, phosphate and silicate are 238 all considered in autochthonous compounds, but not external/forced fluxes (Butenschön, et al., 239 2016). All the biogeochemical representation and mathematical formulations in ERSEM were 240

described in detail in Butenschön et al. (2016).

# 242 **3. Coupling of physical and biogeochemical models**

243 3.1 Physical-biogeochemical model coupling

Coupling between FVCOM and ERSEM were utilized through FABM (Framework for 244 Aquatic Biogeochemical Models; http://fabm.net) (Bruggeman and Bolding, 2014). The aim of 245 this coupling is to provide FVCOM with a mechanism to easily incorporate new and existing 246 biogeochemical models: FABM is a domain-independent programming framework with support 247 for any number of processes, prognostic variables, diagnostic variables, and advanced features 248 such as surface- and bottom layers (sea ice biota, benthos, sediment) and multiple feedbacks to 249 physics. It comes with a comprehensive library of existing biogeochemical models, including 250 descriptions of suspended sediment, redox chemistry (BROM) and pelagic and benthic 251 ecosystems (NPZD, ERGOM, ERSEM). 252

FABM runs as part of its "host model", in this case, it is FVCOM. In a coupled FVCOM-253 FABM simulation, the ERSEM can be run either offline or online simultaneously with FVCOM 254 255 with variable exchanges through FVCOM-FABM coupler (Bruggeman and Bolding, 2014). FABM itself provides information to biogeochemical processes, including source terms and 256 residual vertical velocities (e.g., sinking rates) of prognostic variables, and the value of 257 diagnostics. It doesn't handle features related to hydrodynamics or the spatial domain, e.g., 258 transport of biogeochemical variables, surface boundary conditions (dilution/concentration due 259 to precipitation/evaporation, open boundary conditions), rivers, reading restart files, and saving 260

- output. It also doesn't manage the memory for spatially explicit fields, e.g., the physical
- 262 environment and the biogeochemical variables themselves. All these features are implemented in
   263 the FVCOM-FABM coupler.
- Additionally, a python-based open-source utility PyFVCOM is used to easily manipulate and analyze the results from FVCOM-FABM-ERSEM simulation (Cazenave and Bedington, 2018).
- 3.2 Integration of model system and configuration

The model system for the Changjiang Estuary (Fig. 2) consisted of a shelf-scale East China Sea model and coastal-estuarine scale model for the Changjiang Estuary that incorporated waves and sediment dynamics. Wave-current-sediment interactions were activated only in the coastal-estuarine scale model, and coupling approach was extensively described in Ge et al. (2013). Differing from Ge et al. (2013), the upgraded version included the coupling of ERSEM.



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Figure 2. Framework of the FVCOM-based coupling model system for physical and biogeochemical dynamics,
 including regional FVCOM model for the East China Sea and adjacent seas, coastal-estuarine FVCOM model for
 the Changjiang Estuary, the surface wave model FVCOM-SWAVE, sediment model FVCOM-SED, the
 biogeochemistry and ecosystem model FVCOM-ERSEM.

The model grids for the regional and coastal domains were shown in Fig. 3. The regional model covers the adjacent shelf seas, including the Bohai Sea, Yellow Sea and the East China Sea (Fig.4 a), with a mesh resolution of ~3km along the coastal region and Kuroshio path and meandering region (Chen et al., 2008; Ge et al., 2013). The large domain provides the boundary conditions of tide, ocean circulation and waves to the higher resolution coastal domain.

The coastal-estuarine domain for the Changjiang Estuary was discretized as the refined mesh with a resolution of  $\sim$ 1-3 km in the river mouth and coastal region (Fig. 3b). The upstream

- river boundary extended to the Datong station, which was the closest hydrological station to the
- tidal-limit measuring runoff rate, sediment flux and biological variables' concentrations. The
- 287 coastal-estuarine domain covered the high-turbidity region in the estuary, including the
- Hangzhou Bay and the Jiangsu offshore region. The lateral open boundary was placed at
- 124.5°E, while the north and south lateral boundaries were located at 34.2°N and 28.25°N,
- respectively, with the purpose of eliminating the noise from the boundaries in our estuarine
- region of interest.



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Figure 3. Unstructured meshes for the shelf-scale East China Sea model (a) and coastal-estuarine scale Changjiang
 Estuary model (b). Two individual locations (Site A and B) and one section (blue dashed line) from the river mouth
 to the offshore region are included in (b).

The Connecting between the regional and coastal domains was performed with a one-way nesting method. These two model meshes shared a common-grid layer with identical bathymetry, horizontal and vertical coordinates (blue lines in Fig. 3). This common-grid nesting method guaranteed the conservation of mass and momentum during regional-to-coastal nesting (Chen et al. 2013; Ge et al., 2013; Qi et al., 2018), thus this coupling of physical and biogeochemical dynamics through FABM was only active in the coastal-estuarine scale domain (Fig. 2).

The simulation covered the period from January 1 1999 to December 31 2016. The atmospheric conditions were driven by ERA Interim data from European Centre for Medium-Range Weather Forecasts (ECMWF). Surface wind velocities at 10 m height, shortwave, longwave, latent and sensible radiation flux were provided at a 6-hour interval. Lateral boundary conditions were specified with a hybrid forcing of astronomical tide and subtidal oceanic

- 307 currents. The tidal boundary condition for the regional domain includes eight major tide
- harmonic constituents, M<sub>2</sub>, S<sub>2</sub>, K<sub>2</sub>, N<sub>2</sub>, K<sub>1</sub>, O<sub>1</sub>, P<sub>1</sub>, and Q<sub>1</sub>, retrieved from TPXO 7.2 Global Tidal
- 309 Solution (Egbert and Erofeeva 2002). Volume transport along the lateral boundary is interpolated
- 310 from daily HYCOM/NCODA Global 1/12° Analysis data (GLBa0.08 branch). The upstream
- river boundary was driven by the daily freshwater and sediment discharges collected at the
- 312 Datong station (www.cjh.com.cn ).
- Satellite-derived sea surface temperature (SST) data were assimilated in the model with a nudging algorithm (Chen et al., 2013). Daily SST data from the Advanced Very High-Resolution
- Radiometer (AVHRR) L4 at 0.25° resolution was assimilated in the model over the period from
- 1999 to 2004, while the data from GHRSST/JPL (https://podaac.jpl.nasa.gov/GHRSST) at
- 317 0.011° resolution over the period from 2005 to 2016. The wave model followed the
- configuration described in Ge et al. (2013). The sediment model shared the same parameters as
- 319 Ge et al. (2015). The spatially non-uniform distribution of the critical shear stress for erosion,
- adopted from Ge et al. (2015), were also applied in this configuration.
- Since the main purpose of our modeling study was to examine the nutrients and
   phytoplankton dynamics in the pelagic system and the benthic ecosystem is not a dominant
   contributor, only the pelagic (non-benthic) dynamics were considered here.
- To avoid zero values of the carbonate system variables at the river nodes, the dissolved oxygen (DO), carbonate bio-alkalinity, total alkalinity, and carbonate total dissolved inorganic carbon (DIC) were encoded by constant values of 200 mm al  $O(m^3 - 2.50)$  mm al/lag 20(5.0)
- carbon (DIC) were specified by constant values of 300 mmol O/m<sup>3</sup>, 2.50  $\mu$ mol/kg, 2065.0
- $\mu$ mol/kg, and 2200.0 mmol C/m<sup>3</sup>, respectively. Nutrient concentrations at the lateral boundaries
- were interpolated from the World Ocean Atlas 2013 ver. 2
- 329 (https://www.nodc.noaa.gov/OC5/woa13/). The carbonate system variables (DIC and total
- alkalinity) were determined from the Global Ocean Data Analysis Project version 2
- 331 (GLODAPv2, Olsen et al., 2016) and imposed as monthly climatology. The parameters and
- biogeochemical coefficients used in ERSEM simulation were listed in Table 1.

Symbol	Definition		Value	Unit	
pCO2a	mole fraction of carbon dioxide in air		385	10-6	
a0w	adsorption coefficient of clear water		0.015		m <sup>-1</sup>
b0w	backscatter coefficient of clear water		0.00135		m <sup>-2</sup>
pEIR_eow	photosynthetically active fraction of shortwave radiation		0.5		-1
EPSESSa	specific shortwave absorption coefficient of silt		0.00004		$m^2 mg^{-1}$
EPSESSb	specific shortwave backscatter coefficient of silt		0.00001		$m^2 mg^{-1}$
Nlp	initialization concentration of phosphate phosphorus		0.4	mmol P m <sup>-3</sup>	
N3n	initialization concentration of nitrate nitrogen		8	mmol N m <sup>-3</sup>	
N4n	initialization concentration of ammonium nitrogen		0.1	mmol N m <sup>-3</sup>	
N5s	initialization concentration of silicate		4.5	mmol Si m <sup>-3</sup>	
<i>O2o</i>	initialization concentration of oxygen		300	mmol O <sub>2</sub> m <sup>-3</sup>	
ОЗс	total dissolved inorganic carbon		2130	mmol C m <sup>-3</sup>	
Labile (R1), semi-labile (R2) and refractory (R3) dissolved organic carbon		R1	R2	R3	
С	initialization carbon concentration	10	12	12	mg C m <sup>-3</sup>
n	initialization nitrogen concentration	0.14	-	-	mmol N m <sup>-3</sup>

**Table 1.** Variable definitions and values used in FVCOM-ERSEM coupling.

р	initialization phosphorus concentration	0.01 -			-	mmol P m <sup>-3</sup>
Small (R4), medium (R6) and large (R8) particulate organic carbon				R6	R8	
iopABS	specific shortwave absorption	0.00001 0.00001			0.0000	m <sup>2</sup> m <sup>-1</sup> C
iopBBS	specific shortwave backscatter	0.00001	6 0.	000016	0.0000 16	$m^2 m^{-1} C$
rm	sinking velocity	1		5	10	m d <sup>-1</sup>
С	initialization carbon concentration	7.2		17	0.17	mg C m <sup>-3</sup>
n	initialization nitrogen concentration	0.1		0.24	0.0024	mmol N m <sup>-3</sup>
р	initialization phosphorus concentration	0.007		0.02	0.0002	mmol P m <sup>-3</sup>
S	initialization silicate concentration	-		0.1	0.001	
	benthic dissolved (Q1), particulate (Q6) organic, refractory (Q7) matter	Q1		Q6	Q7	
remin	remineralisation rate	0.1		0.05	0.01	d-1
pN3	nitrate fraction of remineralised nitrogen (remainder is ammonium)	0.9		0.9	0.9	Dimensionless
С	initialization carbon concentration	18.9		3052	30520	mg C m <sup>-2</sup>
n	initialization nitrogen concentration	0.6		3.5	80	mg N m <sup>-2</sup>
р	initialization phosphorus concentration	0.0035		0.5	8	mg P m <sup>-2</sup>
resuspension	enable resuspension	TRUE		-	-	Dimensionless
vel_crit	critical shear velocity for resuspension	0.2		-	-	m s <sup>-1</sup>
S	initialization silicate concentration	149	149 -		-	mg Si m <sup>-2</sup>
pel_nitq10	q <sub>10</sub> temperature coefficient of pelagic nitrification	2				Dimensionless
pel_nitISWph	ph impact on pelagic nitrification (0:off,1:on)		-		Dimensionless	
pel_nitsN4N3	specific nitrification rate of pelagic nitrification	0.5				1 d <sup>-1</sup>
pel_nitchN30	michaelis-menten constant for cubic oxygen dependence of pelagic nitrification	2700				$(\text{mmol O2 m}^{-3})^3$
pel_nitchN4n	michaelis-menten constant for cubic ammonium dependence of pelagic nitrification	0.5			(mmol N m <sup>-3</sup> ) <sup>3</sup>	
Ref_temp	reference temperature	10			degree C	
	Diatoms (P1), Nanoflagellates (P2), Picoplankton (P3), and dinoflagellates (P4) phytoplankton functional groups	P1 P2 P3		P4		
sum	maximum specific productivity at reference temperature	1.375	1.375 1.625		1.125	1d-1
<i>q10</i>	q10 temperature coefficient	2	2 2 2		2	Dimensionless
STS	specific rest respiration at reference temperature	0.04	0.04 0.04		0.035	1d-1
pu_ea	excreted fraction of phytoplankton's primary production	0.2	0.2	0.2	0.2	Dimensionless
pu_ra	respired fraction of phytoplankton's primary production	0.2	0.2	0.2	0.2	Dimensionless
qnlc	minimum nitrogen to carbon ratio	0.0042	0.005	0.006	0.0042	mmol N mg <sup>-1</sup> C
qplc	minimum phosphorus to carbon ratio	0.0001	0.0001 0.0002 0.0 25		0.0001	mmol P mg <sup>-1</sup> C
хqср	threshold for phosphorus limitation (relative to redfield ratio)	1	1 1		1	Dimensionless
xqcn	threshold for nitrogen limitation (relative to redfield ratio)	1 1		1	1	Dimensionless
xqp	maximum phosphorus to carbon ratio (relative to redfield ratio)	2	2	1.5	2.7	Dimensionless
xqn	maximum nitrogen to carbon ratio (relative to redfield ratio)	1.075	1.075	1.05	1.1	Dimensionless
qun3	nitrate affinity	0.0025	0.004	0.006	0.002	$(m^3 mg^{-1} C)d^{-1}$
qun4	ammonium affinity	0.0025	0.004	0.007	0.002	$(m^3 mg^{-1} C)d^{-1}$

qurp	phosphate affinity	0.003	0.004	0.006	0.002	(m <sup>3</sup> mg <sup>-1</sup> C)d <sup>-1</sup>
snplux	specific tendency of luxury uptake of nutrients towards maximum quota	1	1 1 1		1	1 d <sup>-1</sup>
use_Si	if phytoplankton use silicate	TRUE	FALSE	FALSE	FALSE	Dimensionless
qsc	maximum silicate to carbon ratio	0.0118	-	-	-	mmol Si mg <sup>-1</sup> C
chs	michaelis-menten constant for silicate limitation	0.2	-	-	-	mmol m <sup>-3</sup>
sdo	1.1 of minimal specific lysis rate	0.05	0.05	0.055	0.045	1d-1
alpha	initial slope of pi-curve	4	5	6	3	$((mg C m^2 mg^{-1} Chl) W^{-1})d^{-1}$
beta	photoinhibition parameter	0.07	0.1	0.12	0.06	$\begin{array}{c} ((mg \ C \ m^2 \ mg^{-1} \ Chl) \\ W^{-1}) d^{-1} \end{array}$
phim	maximum effective chlorophyll to carbon photosynthesis ratio	0.06	0.025	0.015	0.045	mg Chl mg <sup>-1</sup> C
uB1c_O2	oxygen produced per unit of carbon fixed	0.11	0.11	0.11	0.11	mmol O <sub>2</sub> mg <sup>-1</sup> C
urB1_O2	oxygen consumed per unit of carbon respired	0.1	0.1	0.1	0.1	mmol O <sub>2</sub> mg <sup>-1 C</sup>
iopABS	specific shortwave absorption	0.007	0.0041	0.023	0.008	m <sup>2</sup> mg <sup>-1</sup> Chl
iopBBS	specific shortwave backscatter	0.00048	0.003	0.003	0.0004 8	m <sup>2</sup> mg <sup>-1</sup> Chl
resm	maximum nutrient-limitation-induced sinking velocity	5	0	0	5	m d <sup>-1</sup>
С	initialization carbon concentration	8	5.9	5.9	5.9	mg C m <sup>-3</sup>
п	initialization nitrogen concentration	0.1114	0.1114 0.0926 0.092		0.0926	mmol N m <sup>-3</sup>
р	initialization phosphorus concentration	0.009	0.009 0.0036 0.0036		0.0036	mmol P m <sup>-3</sup>
Chl	initialization chlorophyll a concentration	0.4	0.4 0.3 0.3		0.3	mg m <sup>-3</sup>
Pls	initialization silicate concentration of diatoms		0.1	mmol Si m <sup>-3</sup>		
P2calcify	calcify of nanophytoplankton		TR	UE		Dimensionless
L2sedL2	sinking velocity of calcite		1	0		m d <sup>-1</sup>
L2c0	initialization carbon concentration of calcite		0.	05		mg C m <sup>-3</sup>
bL2remin	remineralisation rate of benthic calcite		0.	05		1d-1
bL2c	initialization carbon concentration of benthic calcite	0.05			mg C m <sup>-2</sup>	
Mesozooplankton (Z4), microzooplankton (Z5), nanoflagellates (Z6)		Z4	Z	5	Z6	
q10	Q10 temperature coefficient	2	2 2		2	Dimensionless
minfood	Michaelis-Menten constant to perceive food	12	1	2	12	mg C m <sup>-3</sup>
chuc	Michaelis-Menten constant for food uptake	36	3	2	28	mg C m <sup>-3</sup>
sum	Maximum specific uptake at reference temperature	1	1.	25	1.5	1 d <sup>-1</sup>
pu	Assimilation efficiency	0.6	0	.5	0.4	Dimensionless
pu_ea	Fraction of unassimilated prey that is excreted (not respired)	0.5	0	.5	0.5	Dimensionless
pe_R1	Dissolved fraction of excreted/dying matter	0.5	0	.5	0.5	Dimensionless
srs	Specific rest respiration at reference temperature	0.015	0.	02	0.025	1d <sup>-1</sup>
sd	Basal mortality	0.05	0.	05	0.05	1d-1
sdo	Maximum mortality due to oxygen limitation	0.2	0.	25	0.3	1d <sup>-1</sup>
chro	Michaelis-Menten constant for oxygen limitation	7.81	7.	81	7.81	Dimensionless
qpc	Phosphorus to carbon ratio	0.000786	5 0.0	01	0.001	mmol P mg <sup>-1</sup> C
qnc	Nitrogen to carbon ratio	0.0126	0.0	167	0.0167	mmol N mg <sup>-1 C</sup>
R1R2	Labile fraction of produced dissolved organic carbon	1	1		1	1d-1
xR1p	Transfer of phosphorus to DOM, relative to POM	1.2	1.	.2	1.2	Dimensionless
xR1n	Transfer of nitrogen to DOM, relative to POM	1	]	l	1	Dimensionless

urB1_O2	Oxygen consumed per carbon respired	0.1 0.1 0.1			mmol O2 mg <sup>-1</sup> C
gutdiss	fraction of prey calcite that dissolves after ingestion	0.5 0.5 0.5			Dimensionless
c0	Background carbon concentration	0.0033	0.0033	mg C m <sup>-3</sup>	
с	Initialization carbon concentration	1.2	7.2	2.421	mg C m <sup>-3</sup>
Z4pu_eaR	Fraction of unassimilated detritus that is excreted (not respired)	0.9			Dimensionless
Z4Minprey	Food threshold for overwintering state of mesozooplankton	300			mg C m <sup>-2</sup>
Z4repw	Specific overwintering respiration of mesozooplankton	0.0025			1d <sup>-1</sup>
Z4mort	Specific overwintering mortality of mesozooplankton	0.0025			1d <sup>-1</sup>
stempp	Specific excretion rate of excess phosphorus		0.5	0.5	1d <sup>-1</sup>
stempn	Specific excretion rate of excess nitrogen		0.5	0.5	1 d <sup>-1</sup>
n	Initialization nitrogen concentration		0.12	0.0505	mmol N m <sup>-3</sup>
р	Initialization phosphorus concentration		0.0113	mmol P m <sup>-3</sup>	
	bacteria		<b>B</b> 1		
iswBlim	Nutrient limitation of bacteria (1:minimum of inorganic and organic availability,2:additive availability)		2		Dimensionless
q10	Q10 temperature coefficient of bacteria		2		Dimensionless
chdo	Michaelis-Menten constant for oxygen limitation of bacteria		0.31		Dimensionless
chn	Michaelis-Menten constant for nitrate limitation of bacteria		0.5	mmol N m <sup>-3</sup>	
chp	Michaelis-Menten constant for phosphate limitation of bacteria	0.1			mmol P m <sup>-3</sup>
sd	Specific mortality of bacteria at reference temperature		0.05		1d <sup>-1</sup>
sum	Maximum specific uptake of bacteria at reference temperature		2.2	1d <sup>-1</sup>	
pu	Efficiency of bacteria at high oxygen levels		0.6	Dimensionless	
puo	Efficiency of bacteria at low oxygen levels	0.2			Dimensionless
srs	Specific rest respiration of bacteria at reference temperature	0.1			Dimensionless
sR1	Maximum turn-over rate of DOM of bacteria	1			1d-1
qpc	Maximum phosphorus to carbon ratio of bacteria	0.0019			mmol P mg <sup>-1</sup> C
qnc	Maximum nitrogen to carbon ratio of bacteria	0.0167			mmol N mg <sup>-1</sup> C
ur_o2	Oxygen consumed per carbon respired of bacteria	0.1			mmol O2 mg <sup>-1</sup> C
sR1N1	Mineralisation rate of labile dissolved organic phosphorus of bacteria	0			1d <sup>-1</sup>
sR1N4	Mineralisation rate of labile dissolved organic nitrogen of bacteria	0			1d <sup>-1</sup>
fsink	Scavenging rate for iron of bacteria		0.00007	1d-1	
c0	Background carbon concentration of bacteria	0.01			mg C m <sup>-3</sup>
rR2	Fraction of semi-labile DOC available to bacteria	0.0075			Dimensionless
rR3	Fraction of semi-refractory DOC available to bacteria	0.0025			Dimensionless
frR3	Fraction of activity respiration converted to semi- refractory DOC	0.3			Dimensionless
c	Initialization carbon concentration of bacteria	15.7			mg C m <sup>-3</sup>
n	Initialization nitrogen concentration of bacteria	0.26			mmol N m <sup>-3</sup>
р	Initialization phosphorus concentration of bacteria	0.029			mmol P m <sup>-3</sup>

#### 335 3.3 Univariate verification

To validate the model, the remote sensing data of *Chl-a* concentration from satellites 336 (MODIS-Aqua and GOCI) were used to perform model-data comparisons. Two sites located 337 offshore of the sediment plume were selected for this comparison (Fig. 3): site A (122.89°E, 338 31.46°N) and site B (122.80°E, 29.92°N). As shown in Fig. 4, the daily MODIS-Aqua Chl-a data 339 for 2005-2016 (red dots) were combined with the GOCI *Chl-a* data for 2012-2016 (blue dots) 340 and then used to verify the model-simulated Chl-a data (solid line). Two distinct blooms were 341 evident from the observational Chl-a in spring and in autumn, respectively. These blooms were 342 captured by the model-simulated results. The peaks of MODIS and simulated Chl-a 343 concentrations both reached  $\sim 10 \text{ mg/m}^3$  during the spring bloom. The root-mean-square-error 344 between them were 2.2 mg/m<sup>3</sup> and 2.47 mg/m<sup>3</sup> at Site A and B, respectively, indicating this 345 coupled model system successfully reproduced the magnitude and seasonal variations of Chl-a in 346 the offshore regions. Besides, the comparison between GOCI data and simulated *Chl-a* shown 347 that short-term temporal variations could also be captured by the model simulation. Although the 348 validation was conducted on a daily basis, it showed the variations over the time scale from 349



seconds to minutes as the model was integrated for iteration.

351

**Figure 4.** Comparisons between model-simulated and satellite remote-sensing-derived surface *Chl-a* concentrations

at Site A and Site B. The red and blue dots indicate the Chlorophyll-a concentration values from MODIS and GOCI
 satellites, respectively.

#### 355 3.3 Multivariate verification

375

In addition to useful independent and univariate assessments, the comparison between 356 model-simulated results and observations for all common variables simultaneously provides a 357 better evaluation of the overall performance of the model (Allen and Somerfield, 2009; Stow et 358 al., 2009). To determine the model's capability of revealing the realistic dynamics in the offshore 359 region, Principle Component Analysis (PCA) was applied to reduce the dimensionality of the 360 dataset and to identify the significant underlying components (Allen and Somerfield, 2009). PCA 361 was performed on both instantaneous observed values and daily averages of variables at 362 measurement sites from surface to bottom layers for salinity, temperature, Chl-a, SiO<sub>3</sub>, NO<sub>3</sub>, 363 PO<sub>4</sub>, SPM, NH<sub>4</sub>, and pH. 364

Both cumulative proportion of the first three eigenvectors of observed and simulated 365 variables were more than 90%, and the first two eigenvectors aggregated more than 85% (Table 366 2). This meant that the first principal component (PC1) and second principal component (PC2) 367 were able to reproduce the main factors that described the variability of the system as captured in 368 the observed variables. PC1 of observed and simulated variability accounted for 64.8% and 369 370 62.5%, respectively, both observed and simulated results indicated the very similar contribution of salinity, temperature, SiO<sub>3</sub>, NO<sub>3</sub> and PO<sub>4</sub>. PC2 was 22.8% and 23.4%, respectively, with the 371 temperature to be the most essential. PC3 of the observed variability was mainly controlled by 372 the combination of NH<sub>4</sub> and suspended sediment, while that of model simulated variability was 373 under the mixed effect of NH4 and Chl-a. 374

Table 2. PCA Analysis of the modeling and measuring variables.								
	observation				simulation			
Eigenvector	PC1	PC2	PC3	PC1	PC2	PC3		
pН	-0.039	0.054	0.149	-0.361	-0.067	0.016		
NH4	-0.021	-0.070	0.891	-0.202	0.142	0.502		
PO <sub>4</sub>	0.402	-0.227	-0.046	0.273	-0.384	-0.231		
NO <sub>3</sub>	0.446	0.017	0.012	0.470	0.109	0.152		
SiO <sub>3</sub>	0.543	0.048	-0.093	0.462	0.148	0.139		
Chl-a	0.006	0.149	0.058	-0.159	0.125	0.687		
salinity	-0.581	-0.167	-0.116	-0.495	-0.283	-0.159		
temperature	-0.035	0.935	0.070	-0.153	0.835	-0.391		
SPM	0.059	-0.116	0.389	0.160	-0.020	0.039		
variance	0.65	0.23	0.04	0.62	0.23	0.07		
variance%	64.89	22.84	3.75	62.49	23.46	7.28		
cumulative%	64.89	87.73	91.48	62.49	85.94	93.23		

The ordination values in the PCA demonstrated the similarity between the main modes of variability in both observed and modeled variables, and the seasonal changes associated with the controls on the pelagic ecosystem and environmental conditions (Fig. 5). The samples of the two

datasets in summer were in the positive direction of PC2 axis and the samples in winter were in

the negative direction of PC2 axis because of the dominating control of temperature in PC2. The

observation PCA points had a more compact distribution than the simulated results. This implied
 that the model reproduced the observed variability to some degree. The high positive values on
 PC2 axis were possibly related with the overestimation of river temperature. In general, both
 patterns from PCA showed some similarities, indicating the model revealed the major modes of

seasonal effects of the biogeochemical variables, at least on winter/summer seasons.



386

Figure 5. Principle Component Analyses (PCA) based on observed (a) and simulated (b) *Chl-a*, nitrate, silicate,
 phosphate and ammonia concentrations as well as temperatures, salinities, pH levels and SPM values. The simulated
 values were the nearest equivalent to the measurements in temporal and spatial coordinates. Data were normalized
 before the analysis was performed. PC1 and PC2 indicate the first and second principal components, respectively.

#### **4. Nutrient and phytoplankton simulations**

# 392 4.1 *Chl-a* and sediment front

To determine the seasonal variability of phytoplankton under the condition with 393 sediments, we calculated the seasonal means of surface Chl-a overlapped with the sediment 394 concentration (Fig. 6). Four seasons were defined as follow, March-May as spring, June-August 395 as summer, September-November as autumn and December-February as winter. At this level of 396 aggregation, the model results clearly showed two distinct zones separated by the waters with 397 high-concentration suspended sediment and with high phytoplankton Chl-a concentration, 398 respectively. The computational domain contained several significant high-turbidity zones, 399 including the Changjiang Estuary, Hangzhou Bay and the coastal region off Jiangsu Province. 400 The model also reproduced the seasonal variation of sediment dynamics in the offshore regions 401 of the Changjiang Estuary and Jiangsu. The sediment front in the Changjiang Estuary was 402 confined within 122.5°E during spring and summer (Fig. 6a-b), but extended to 123°E in autumn 403

and winter (Fig. 6c-d). In the Jiangsu offshore region, there was a similar pattern, but with further offshore extension in autumn and winter, which reached  $\sim 124.3^{\circ}$ E in winter.



Figure 6. Seasonal-averaged surface distributions of the suspended sediment concentration (contour, unit: g/L) and
 phytoplankton *Chl-a* concentration (colored image, unit: mg/m<sup>3</sup>) during spring (a), summer (b), autumn (c) and
 winter (d).

406

However, the extension of this sediment front is not uniquely controlled by the remote
 horizontal sediment advection. Local resuspension dynamics in combination with vertical

412 stratification were responsible for the behavior and extension of the sediment plume. In spring

- and summer, in the offshore region, under strong stratification condition, it was due to the off-
- shore extension of the low-salinity river plume (Wu et al., 2011; Ge et al., 2015). The advection
- 415 of riverine derived SSC was the major driving force since the ambient concentrations were low.
- 416 Surface heating in these two seasons also played an important role in intensifying stratification.
- This limited the upward vertical transport of sediment in the offshore region and led to low SSC in the upper water column. With sufficient nutrient supply, the increase in light availability in the
- 416 In the upper water column, with sufficient nutrient suppry, the increase in light availability in the 419 surface layers under a low SSC condition made a more favor environment for the phytoplankton
- 420 growth in the offshore region compared to the nearshore and estuarine region.

In autumn and winter, the physical conditions changed. Due to decreasing freshwater discharge and colder weather-induced surface cooling, the offshore destratification occurred. Vertical mixing increased the local resuspension of sediment throughout the water column (Luo et al., 2018), producing the high broad-scale sediment concentration at the surface (Fig. 6d). Under lower water temperature and light penetration, the phytoplankton growth in the offshore region was greatly limited.

427 4.2 Diatom and non-diatom phytoplankton group

Multiple species of phytoplankton jointly contributed to the total Chl-a concentration in 428 the observations and model results. In the shelf of the ECS, diatoms and dinoflagellates are the 429 major groups and generally represent up to 90 % of the total biomass in the spring bloom (Chen 430 et al., 2003; Zhu et al., 2009). While the single specie Skeletonema costatum dominates the 431 diatom group, dinoflagellates are represented by hundreds of species, covering sizes from 432 microphytoplankton (> 20  $\mu$ m), nanophytoplankton (2–20  $\mu$ m), to picophytoplankton (< 2  $\mu$ m). 433 Nonetheless, the dominant specie is Prorocentrum donghainese. In order to facilitate the 434 comparison with observations and known dynamics in the region, we have aggregated the model 435 results for microphytoplankton, nanophytoplankton and picophytoplankton from FVCOM-436 ERSEM modeling results as a non-diatom phytoplankton group. 437

To determine the major phytoplankton group that contributes to the increase of *Chl-a* in 438 spring and summer (Fig. 6a-b), the surface carbon-based (not *Chl-a*-based) biomass of diatom 439 and non-diatom phytoplankton were shown in Fig. 7. The overall diatom percentage in the 440 phytoplankton community was about 43% during spring. The results showed that diatom carbon 441 distribution (Fig. 7a) matched well with the pattern of the *Chl-a* concentration during spring 442 season (Fig. 6a), indicating the diatom had a major contribution to the *Chl-a* concentration. 443 However, the diatom biomass was weaker than the non-diatom group, including dinoflagellate 444 and other small-size species. The distribution pattern of diatom followed the *Chl-a* distribution, 445 which were confined offshore but closed to the sediment frontal boundary along the estuary, 446 447 indicating light availability limits the diatom growth. Non-diatom biomass was higher offshore of the sediment front in the shelf region, particularly in the northeastern region, where the non-448 diatom concentration reached 200 mg  $C/m^3$ , while the *Chl-a* concentration remained low (50-75) 449 mg  $C/m^3$ ) in spring due to nutrient limitation. 450

In summer, the diatom percentage in total biomass of phytoplankton declined to 37%, especially on the offshore side of the sediment front (Fig. 7c). The wide diatom biomass regions shrank to a narrow band along the sediment front on the offshore side with a maximum concentration declining from ~175 to ~80 mg C/m<sup>3</sup>, resulting from the diatoms' dependency on light and nutrient physiologically. On the other hand, non-diatom had high biomass both along the sediment front and even in the out region, with a maximum concentration of more than 200

457 mg C/m<sup>3</sup> (Fig. 7d).



458

Figure 7. Seasonal-averaged surface distributions of the suspended sediment concentration (unit: g/L) (contours) and biomass (colored image; unit: mg C/m<sup>3</sup>) of diatom (a, c) and dinoflagellate (b, d) (colored image) during spring

461 (a-b) and summer (c-d).

462

463

#### 4.3 Dissolved Nutrients

Taking the nitrogen flux from the Changjiang Estuary for example, the annual DIN flux 464 was about  $1.25 \times 10^{11}$  mol yr<sup>-1</sup>; the particulate nitrogen, however, was only about  $0.9 \times 10^{10}$  mol 465 yr<sup>-1</sup> (Gao et al., 2012). The content of PN was less than 10% of total nitrogen. The particulate 466 nutrient was only a small component in the whole nutrient based on the measurements in the 467 Changjiang River (Gao et al., 2012). Among the various phosphorus forms, particulate 468 phosphorus represented <50%, and is closely related to the concentration of suspended sediment 469 concentration (Liu et al., 2016). Nonetheless, the dissolved inorganic nutrients made the major 470 contribution to the phytoplankton growth and ecosystem cycle. Therefore, the temporal and 471 spatial variabilities of nutrients were determined through their dissolved inorganic forms. 472

While dissolved nutrients were mainly dominated by the Changjiang River source in the 473 study area, they experienced various seasonal behaviors. Taking the DIN and DIP as examples, 474 they behaved conservatively in the high-turbidity area, undergoing only physical dilution from 475 the river channel to the river mouth (Fig. 8-9). Phytoplankton growth was quite weak in this area, 476 and thus no significant nitrate was consumed. In the offshore region, DIN was consumed in the 477 offshore side region of the sediment front during spring and summer (Fig. 8a-b), causing a sharp 478 479 decrease over there. However, since the DIN was not the limiting factor in this system, the concentrations in the offshore region of the sediment front were still high, 40-70 mmol N/m<sup>3</sup> in 480 summer (Fig. 8b) and 10-30 mmol N/m<sup>3</sup> in winter (Fig. 8d). With the eastward extension of the 481 sediment front in autumn and winter, the DIN reached the offshore region as a result of 482 advection (Fig. 8c-d). 483

DIP, on the other hand, was the limiting dissolved nutrient in the ecosystem of the 484 Changjiang Estuary and adjacent regions (Gao et al., 2005; Zhou et al., 2017). In addition to 485 riverine sources, local mixing, pelagic and benthic bacterial mediated degradation were also the 486 significant sources of DIP, causing DIP varying with the stratification status. DIP produced by 487 these dynamics had the similar distribution patterns as the surface sediment. Phytoplankton 488 growth consumed most of DIP in the water column during spring, summer and autumn, forming 489 the same boundary of DIP and SSC at the sea surface in these three seasons (Fig. 9a-c). Outside 490 the sediment front, the concentration of DIP was quite low, indicating DIP was depleted due to 491 the growth of phytoplankton. In winter, the concentration of DIP increased in the offshore area 492 due to vertical mixing and weak growth of phytoplankton (Fig. 9d), and similar behavior was 493 seen in DIN in the offshore area (Fig. 8d). In winter, vertical mixing provided more dissolved 494 nutrients to the upper column (Fig. 8d and Fig. 9d) and supported the growth of phytoplankton 495 496 offshore in Zhejiang, where the suspended sediment concentration was relatively low.





Figure 8. Seasonal-averaged surface distributions of suspended sediment concentration (contour, unit: g/L) and DIN 499 (colored image, unit: mmol N/m<sup>3</sup>) during spring (a), summer (b), autumn (c) and winter (d).





Figure 9. Seasonal-averaged surface distributions of suspended sediment concentration (contour, unit: g/L) and DIP
 (colored image, unit: mmol P/m<sup>3</sup>) during spring (a), summer (b), autumn (c) and winter (d).

503 Vertical mixing in the offshore region played an important role in the seasonal regulation 504 of sediment, phytoplankton and dissolved nutrients as mentioned above. Thus, a vertical section 505 from the river mouth to the offshore region was selected to investigate the seasonal variability of 506 the vertical distribution (blue dashed line in Fig. 3). On this transect, the SSC in the upper layer

of the offshore region increased from summer to winter (Fig. 10e, i and m) due to the decrease in 507 stratification and higher resuspension driven by the increasing vertical mixing (Fig. 10h, 1 and p). 508 The highest concentration of *Chl-a* in the low-turbidity upper column represented the spring 509 510 bloom of phytoplankton (Fig. 10b). The DIP distribution showed a distinct low-value zone due to

the consumption by phytoplankton in the upper column (Fig. 10c). 511

In summer, the stratification was strongest (Fig. 10h), resulting in the lowest sediment 512 concentrations in the upper water column (Fig. 10e). Under this weak-mixing condition, the DIP 513 in the upper column was depleted after the spring bloom and not replenished via vertical mixing 514 (Fig. 10g), this dissolved nutrient limiting decreased the phytoplankton growth at the surface. 515

Under these conditions, a subsurface Chl-a maximum developed (Fig. 10f). 516



517 518

Figure 10. Seasonal-averaged vertical distributions of the suspended sediment concentration (first column, unit: 519 g/L), phytoplankton (second column, unit: mg/m<sup>3</sup>), DIP (third column, unit: mmol P/m<sup>3</sup>), and temperature (forth 520 column, unit: degree) on the selected section during spring (a-d), summer (e-h), autumn (i-l) and winter (m-p).

521 With stronger vertical mixing from autumn to winter, and additional remineralization of 522 detritus from dead phytoplankton, zooplankton and other biological components, the 523 concentration of DIP reached its peak in winter. The whole water column became well mixed 524 with DIP concentrations reaching ~0.8 mmol P/m<sup>3</sup>.

In summary, the interactions between the physical and biogeochemical processes modulated the seasonal variabilities of dissolved nutrients and phytoplankton around the offshore area. The sediment dynamics confined the light penetration in the water column, which dominated the boundary of phytoplankton growth. Vertical mixing in the different seasons led to different dissolved nutrient supplies regimes, with river supplies becoming less important in autumn and winter.

Although dissolved nutrients were the major components of nutrient states, it should be noted that besides the dissolved form, the particulate nutrients, both organic and inorganic form, also contributed to the nutrient cycle, especially for the phosphorus (Liu et al., 2016). The particulate nutrients had much more complex dynamics, such as interaction with suspended sediment particles. Since most of available measurements for nutrients in this study were based on dissolved form, the only dynamics of dissolved nutrients were examined. The particulate nutrients should be resolved if one needs to understand full dynamics of nutrients on all forms.

538

# 539 **5. Discussion**

540

5.1 Interpretation from remote sensing data

Remote sensing from satellite platforms has been recognized as a useful method to 541 retrieve surface SPM and Chl-a (O'Reilly et al., 1998; Shen et al., 2010). Using the semi-542 empirical radiative transfer (SERT) algorithm with physical based empirical coefficients (Shen et 543 al., 2010), the snapshot for the Changjiang Estuary and its adjacent regions at 07:16 am May 16, 544 2016 in the spring season illustrated the offshore extent of the SSC in Fig. 11a. It appeared that 545 turbid water covered the whole coastal region with the typical sediment concentration of ~500 546 mg/L. In the offshore region, the concentrations declined to 20-50 mg/L, where light penetrated 547 into the water column and promote phytoplankton growth. 548

*Chl-a* was mostly observed in the offshore region (Fig. 11b), where relatively low-549 turbidity water presented (Fig 11a). In the turbidity maximum and coastal high-turbidity regions, 550 the concentration of *Chl-a* was low due to light limitation. The front between the high-, and low-551 turbidity water acted like a boundary for phytoplankton growth, which demonstrated the 552 sediment's modulation of phytoplankton dynamics. The location of modeled high Chl-a in spring 553 matched the spot where historical spring bloom was observed (Gao et al., 2005; Zhu et al., 2009; 554 Guo et al., 2014). The distribution of Chl-a concentration showed a narrow band off the 555 sediment front in May. 556

557 The snapshots of SPM and *Chl-a* from remote sensing also revealed the existence of 558 small-scale structures along the sediment front. These fine structures suggested the existence of 559 eddies along the low-salinity front. Chen et al. (2008) proposed that such eddies were produced 560 by baroclinic instabilities due to the salinity-induced density gradient. These eddies also led to 561 some isolated patches of high-turbidity water, causing detachment occurred, then resulting in the 562 horizontal mixing of coastal high-turbidity water and offshore low-turbidity water. it was partially responsible for the supply of dissolved nutrients that supported that narrow diatom patch along the front. The snapshot from remote sensing showed that sediment-phytoplankton interactions exhibited much more complicated temporal and spatial variations on tidal or windevent scales. This was a mechanism presented in the model and it was responsible for the supply of dissolved nutrients that supported that narrow diatom patch along the front over a short time

scale. This eddy-induced water exchange between inner and outer regions of the front was

- revealed also in previous studies (Chen et al., 2008; Ge et al., 2015a). However, it should be noted that these structures have been smoothened during the temporally-averaging on the
- 571 seasonal scale shown in Fig. 6-9.





**Figure 11.** Distributions of the GOCI-retrieved surface sediment concentration (a) and *Chl-a* (b) around the Changjiang Estuary from the snapshot at 15:30PM, May 16th, 2016.

# 575 5.2 Influence of ocean circulation

Besides the regulation effect that riverine sediment or locally resuspended sediment had on phytoplankton and nutrient dynamics, the oceanic flow also played an important role in the offshore dynamics of the pelagic ecosystem. The Taiwan Warm Current (TWC) is the strongest oceanic intrusion in the offshore region of the Changjiang Estuary, influencing the physical and biogeochemical conditions.

581 In summer, the TWC reached the Changjiang Estuary, with seasonal-averaged velocities of ~0.1m/s in the bottom layer (~40-50m) (Fig. 12a). This current carried high-temperature water 582 from the south to the north. Additionally, the intrusion of TWC and Kuroshio were recognized as 583 a water mass with relatively higher DIP concentration in the bottom column, compared with 584 adjacent shelf water (Chen et al., 2003b, Zhou et al., 2017). These higher-DIP water from 585 intrusion and its induced upwelling partly supported the dinoflagellate bloom in the coastal 586 region (Zhou et al., 2019). Along the selected dashed line in Fig. 12a, the SSC was relatively 587 low, less than 10 mg/L in the offshore area (Fig. 12b), indicating the SPM had s weak effect on 588 the biogeochemical process in this area. While there was low DIP concentration of less than 0.1 589 mmol/ $m^3$  in the surface water, the TWC contributed to higher DIP in bottom (> 0.6 mmol/ $m^3$ ), 590 thus vertically stratified DIP distribution was seen from Fig. 12c with additive effect from 591 summer stratification in water column discussed in Section 4. Due to low SPM, the light reached 592

subsurface layer (~25-30m), and enabled the development of a subsurface *Chl-a* maximum

where the nutrients were adequate for phytoplankton growth (Fig. 12c), which matches well with

595 previous measurements in Chen et al. (2003b). Previous observations also indicated that the DIP 596 from the TWC played a critical role in the maintenance of dinoflagellate blooms here (Zhou et

<sup>597</sup> al., 2017).



598



601 5.3 Zooplankton impact

602 Phytoplankton biomass are not the only functional groups depending on nutrients and 603 light. There are four phytoplankton, three zooplankton and one bacteria functional groups in 604 ERSEM model, so the competition (for resources, i.e. nutrients) and predation also plays a role. 605 Low summer diatoms biomass generally reflects both higher predation pressure from 606 zooplankton and competition from smaller phytoplankton and bacteria which have higher growth 607 rates and smaller sedimentation rates. In fact, zooplankton predation together with nutrient 608 depletion contributes to the demise of the diatom spring bloom.

The model results showed that the total zooplankton biomass (mesozooplankton, microzooplankton and heterotrophic flagellates) on the seasonal scale mainly followed the distribution of phytoplankton, namely higher concentrations during spring and summer, and lower concentrations in autumn and winter (Fig. 13). Zooplankton dominated the larger offshore areas of the sediment front, reaching the outer shelf where non-diatom biomass was adequate to be predated. Meanwhile, low zooplankton concentration (<70 mg C/m<sup>3</sup>) was found within the coastal region in all seasons.





Figure 13. Seasonal-averaged surface distributions of the suspended sediment concentration (contour) and total
biomass of zooplankton (colored image) during spring (a), summer (b), autumn (c) and winter (d).

### 5.4 Stability of seasonal structures

The model simulation covered the time period from 1999 to 2016. Besides the 621 atmospheric and oceanic variabilities that caused the model fluctuations, the major influencing 622 factor for the Changjiang Estuary and adjacent region could be the construction of the Three 623 Gorges Dam. It became operational in 2003, and caused very strong sedimentation impact on the 624 lower reach of the Changjiang river. The sediment concentration into the estuary has significant 625 decreased, as well as the annual sediment load (Luan et al., 2016; Yang et al., 2015). However, 626 the decrease in sediment concentration mainly occurred in the river channel from Datong station 627 to river mouth. The offshore region, particularly around the sediment front region in this study, 628 did not show significant variation trend based on continuous remote sensing from 2003 to 2010 629 (Shen et al., 2013). It mainly indicated a local temporal oscillation. The annual freshwater flux 630 before and after the construction of the Three Gorges Dam were of the same amplitude (Luan et 631 632 al., 2016).

633 Since the freshwater discharge and offshore sediment concentration did not change 634 significantly, the seasonality of dissolved nutrients and phytoplankton has remained relatively 635 stable before and after the operation of Three Gorges Dam. The magnitude of nutrient and 636 phytoplankton concentration surely has some oscillation, which probably was still caused by 637 increasing use of fertilizers.

5.5 Stoichiometry in sediment and nutrients

Nutrients entering the estuarine environment, with an additional contribution from 639 submarine groundwater discharges, can undergo rapid adsorption into suspended sediments with 640 the potential to conversely have desorption processes during transportation and resuspension, or 641 642 even deposit into sea bed (Wang et al., 2018; Liu et al., 2017, 2018). It should be noted that the absorption and desorption of nutrients in the suspended particulate matters were not included in 643 the ERSEM. Nutrients were treated as dissolved and particulate states in the biogeochemical 644 model. Based on previous experiments, the nitrate has weak capacity on absorption and 645 desorption in sediment, whereas the phosphate has notable absorption and desorption processes 646 with suspended sediment. In future studies, the sediment's biogeochemical process should be 647 included in whole nutrient cycle, particularly the adsorption/desorption behavior for phosphorus, 648 given its great importance in high-turbidity environments such as the Changjiang Estuary and 649 adjacent coastal regions. 650

# 651 6. Summary

In this study, a coupled physical-biogeochemical model system, based on FVCOM and 652 ERSEM, was developed to examine the impact of the sediment front on the nutrients and 653 phytoplankton dynamics over the inner shelf of the ECS off the high-turbidity Changjiang 654 Estuary. The model system successfully revealed the variation of *Chl-a* in the offshore region, 655 reproducing the major variability modes of various physical and biogeochemical variables 656 obtained from the seasonal field campaigns as seen in PCA. The model results revealed that the 657 suspended sediments played a role in regulating the growth of phytoplankton. Influenced by the 658 seasonal variations in the extension of low-salinity water, which carried nutrients and sediment, 659 the phytoplankton followed the sediment from more onshore locations during spring and 660 summer to further offshore locations during autumn and winter. Results demonstrated that 661 diatoms were major contributors to Chl-a concentration, but lower biomass contribution to total 662

663 phytoplankton carbon, and non-diatoms (i.e. dinoflagellates etc.) dominated out of the sediment 664 front, and extended to the outer shelf region.

665 DIN and DIP had different behaviors on the seasonal scale. Nitrate behavior could 666 generally be approximated by physical dilution and transport of freshwater discharge, and

remained high offshore of the sediment front. Whereas phosphate, acting as the limiting nutrient

668 in the ecosystem, showed more local dynamics and reached lower concentrations offshore of the

- front. It's implied from the vertical profiles that vertical mixing largely controlled sediment re-
- 670 suspension, nutrient mixing and phytoplankton distribution in the offshore water column.
- 671

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681 682	References
683	Allen, J. I., & Somerfield, P. J. (2009). A multivariate approach to model skill assessment.
684	<i>Journal of Marine Systems</i> , 76(1), 83–94. http://doi.org/10.1016/j.jmarsys.2008.05.009
685	Baretta-Bekker, J. G., Baretta, J. W., & Koch Rasmussen, E. (1995). The microbial food web in
686	the European Regional Seas Ecosystem Model. <i>Netherlands Journal of Sea Research</i> ,
687	33(3-4), 363–379. http://doi.org/10.1016/0077-7579(95)90053-5
688	Beardsley, R. C., Chen, C., & Xu, Q. (2013). Coastal flooding in Scituate (MA): A FVCOM
689	study of the 27 December 2010 nor'easter. <i>Journal of Geophysical Research-Oceans</i> ,
690	118(11), 6030–6045. http://doi.org/10.1002/jgrc.20443
691	Blackford, J. C., Allen, J. I., & Gilbert, F. J. (2004). Ecosystem dynamics at six contrasting sites:
692	a generic modelling study. <i>Journal of Marine Systems</i> , 52(1-4), 191–215.
693	http://doi.org/10.1016/j.jmarsys.2004.02.004
694	Boynton, W. R., Ceballos, M. A. C., Bailey, E. M., Hodgkins, C. L. S., Humphrey, J. L., &
695	Testa, J. M. (2018). Oxygen and Nutrient Exchanges at the Sediment-Water Interface: a
696	Global Synthesis and Critique of Estuarine and Coastal Data. <i>Estuaries and Coasts</i> ,
697	41(2), 1–33. http://doi.org/10.1007/s12237-017-0275-5
698	Bruggeman, J., & Bolding, K. (2014). A general framework for aquatic biogeochemical models.
699	<i>Environmental Modelling and Software</i> , 61(C), 249–265.
700	http://doi.org/10.1016/j.envsoft.2014.04.002
701	Butenschon, M., Clark, J., Aldridge, J. N., Allen, J. I., Artioli, Y., Blackford, J., et al. (2016).
702	ERSEM 15.06: a generic model for marine biogeochemistry and the ecosystem dynamics
703	of the lower trophic levels. <i>Geoscientific Model Development</i> , 9(4), 1293–1339.
704	http://doi.org/10.5194/gmd-9-1293-2016-supplement
705	Cazenave, P. W. and Bedington, M. (2018). PyFVCOM (version 2.1.3) [software]. Plymouth,
706	Devon, United Kingdom: Plymouth Marine Laboratory.
707	https://doi.org/10.5281/zenodo.1422462
708	Chen, C., Beardsley, R. C., and Limeburner, R. (1994). Comparison of winter and summer
709	hydrographic observations in the Yellow and East China Seas and adjacent Kuroshio
710	during 1986. <i>Continental Shelf Research</i> . 14, 909-929.

- Chen, C. H. Liu, R. C. Beardsley. (2003a). An unstructured, finite-volume, three-dimensional,
   primitive equation ocean model: application to coastal ocean and estuaries. *Journal of Atmospheric and Oceanic Technology*. 20, 159-186.
- Chen, C., J. Zhu, R. C. Beardsley, and P. S. Franks. (2003b). Physical-Biological Sources for the
   Dense Algal Bloom over the Western Shelf of the East China Sea. *Geophysical Research Letter*. 30(10), 1515, 22-1:4.
- Chen, C., L. Wang, R. Ji, J. W. Budd, D. J. Schwab, D. Beletsky, G. L. Fahnenstiel, H. A.
  Vanderploeg, B. J. Eadie, and J. Cotner (2004). Impacts of suspended sediment on the
  ecosystem in Lake Michigan: A comparison between the 1998 and 1999 plume events. *Journal of Geophysical Research*. 109(C10S05):18,
- Chen, C, R. C. Beardsley and G. Cowles (2006). An unstructured grid, finite-volume coastal
   ocean model (FVCOM) system. Special Issue entitled "Advance in Computational
   Oceanography", *Oceanography*. vol. 19, No. 1, 78-89.
- Chen, C. P. Xue, P. Ding, R. C. Beardsley, Q.Xu, X. Mao, G. Gao, J. Qi, C. Li, H. Lin, G.
  Cowles and M. Shi (2008). Physical mechanisms for the offshore detachment of the
  Chanjiang diluted water in the East China Sea. *J. Geophys. Res.*, 113. C02002, doi:
  10.1029/2006JC003994.
- Chen, C., R.C. Beardsley, G. Cowles, J. Qi, Z. Lai, G. Gao, D. Stuebe, Q. Xu, P. Xue, J. Ge, R.
  Ji, S. Hu, R. Tian, H. Huang, L. Wu, H. Lin, Y. Sun and L. Zhao (2013), An unstructured
  grid, finite-volume community ocean model FVCOM user manual, SMAST/UMASSD
  Technical Report 13-0701, School of Marine Science and Technology, University of
  Massachusetts-Dartmouth, New Bedford, MA, USA
- Choi, J. K., Y. J. Park, J. H. Ahn, H. S. Lim, J. Eom, and J. H. Ryu (2012), GOCI, the world's
  first geostationary ocean color observation satellite, for the monitoring of temporal
  variability in coastal water turbidity, *J. Geophys. Res.*, 117, C09004,
  doi:10.1029/2012JC008046.
- Choi, J. K., J. E. Min, J. H. Noh, T. H. Han, S. Yoon, Y. J. Park, J. E. Moon, J. H. Ahn, S. M.
  Ahn, and J. H. Park (2014), Harmful algal bloom (HAB) in the East Sea identified by the
  Geostationary Ocean Color Imager (GOCI), *Harmful Algae*, 39, 295-302,
- CWRC (Changjiang Water Resources Commission) (2011). Changjiang Sediment Bulletin.
   available at:. http://www.cjh.com.cn/pages/nsgb.html (in Chinese).
- Donohue, I., & Garcia Molinos, J. (2009). Impacts of increased sediment loads on the ecology of
  lakes. *Biological Reviews*, 84(4), 517–531. http://doi.org/10.1111/j.1469185X.2009.00081.x
- de Swart, H. E., Schuttelaars, H. M., & Talke, S. A. (2009). Initial growth of phytoplankton in
  turbid estuaries: A simple model. *Continental Shelf Research*, 29(1), 136–147.
  http://doi.org/10.1016/j.csr.2007.09.006
- Ebenhoh, W., Kohlmeier, C., Radford, P.J. (1995). The benthic bio-logical model in the
  European Regional Seas Ecosystem Model. *Netherlands Journal of Sea Research*, 33,
  423–452.

# Egbert, G. D., & Erofeeva, S. Y. (2002). Efficient inverse modeling of barotropic ocean tides. *Journal of Atmospheric and Oceanic Technology*, 19(2), 183–204.

- Friedl, G. & Wüest, A. (2002). Disrupting biogeochemical cycles Consequences of damming
   *Aquat. sci.* 64: 55. https://doi.org/10.1007/s00027-002-8054-0
- Furuya, K., Kurita, K., and Odate, T. (1996). Distribution of phytoplankton in the East China Sea
  in the winter of 1993, *J. Oceanogr.*, 52, 323–333, https://doi.org/10.1007/BF02235927,
- Furuya, K., Hayashi, M., Yabushita, Y., and Ishikawa, A. (2003). Phytoplankton dynamics in the
   East China Sea in spring and summer as revealed by HPLC-derived pigment signatures,
   *Deep-Sea Res. Pt. II*, 50, 367–387, https://doi.org/10.1016/S0967-0645(02)00460-5.
- Gao, X., & Song, J. (2005). Phytoplankton distributions and their relationship with the
   environment in the Changjiang Estuary, China. *Marine Pollution Bulletin*, 50(3), 327–
   335. http://doi.org/10.1016/j.marpolbul.2004.11.004
- Ge, J., Ding, P., Chen, C., Hu, S., Fu, G., & Wu, L. (2013). An integrated East China Sea–
  Changjiang Estuary model system with aim at resolving multi-scale regional–shelf–
  estuarine dynamics. *Ocean Dynamics*, 63(8), 881–900. http://doi.org/10.1007/s10236013-0631-3
- Ge, J., Chen, C., & Ding, P. (2015). Estimation of critical shear stress for erosion in the
   Changjiang Estuary: A synergy research of observation, GOCI sensing and modeling.
   *Journal of Geophysical Research-Oceans*, 120(1), 8439–8465.
   http://doi.org/10.1002/2015JC010992
- Ge, J., Ding, P., & Chen, C. (2015). Low-salinity plume detachment under non-uniform summer
   wind off the Changjiang Estuary. *Estuarine, Coastal and Shelf Science*, 156(C), 61–70.
   http://doi.org/10.1016/j.ecss.2014.10.012
- Ge, J., Zhou, Z., Yang, W., Ding, P., Chen, C., Wang, Z. B., & Gu, J. (2018). Formation of
   Concentrated Benthic Suspension in a Time-Dependent Salt Wedge Estuary. *Journal of Geophysical Research-Oceans*, 123(11), 8581–8607.
   http://doi.org/10.1029/2018JC013876
- Geider R.J., MacIntyre H.L., Kana T.M. (1997). A dynamic model of phytoplankton growth and
   acclimation: responses of the balanced growth rate and the chlorophyll a:carbon ratio to
   light, nutrient-limitation and temperature. *Mar Ecol Prog Ser* 148:187-200
- Guo, S., Feng, Y., Wang, L., Dai, M., Liu, Z., Bai, Y., & Sun, J. (2014). Seasonal variation in the
   phytoplankton community of a continental-shelf sea: the East China Sea. *Marine Ecology Progress Series*, 516, 103–126. http://doi.org/10.3354/meps10952
- He, Q., Qiu, Y., Liu, H., Sun, X., Kang, L., Cao, L., et al. (2017). New insights into the impacts
  of suspended particulate matter on phytoplankton density in a tributary of the Three
  Gorges Reservoir, China. *Scientific Reports*, 7(1), 1–11. http://doi.org/10.1038/s41598017-13235-0
- Hu, B., Wang, P., Zhang, N., Wang, C., & Ao, Y. (2016). Photoproduction of dissolved organic carbon and inorganic nutrients from resuspended lake sediments. *Environmental Science and Pollution Research*, 23(21), 1–10. http://doi.org/10.1007/s11356-016-7327-4
- Huettel, M., Berg, P., & Kostka, J. E. (2014). Benthic Exchange and Biogeochemical Cycling in
   Permeable Sediments. *Annual Review of Marine Science*, 6(1), 23–51.
   http://doi.org/10.1146/annurev-marine-051413-012706

- Jiang T, Yu ZM, Song XX, Cao XH, Yuan YQ. (2010). Long-term ecological interactions
   between nutrient and phytoplankton community in the Changjiang estuary. *Chin J Oceanol Limnol*; 28: 887–98.
- Jiang, Z., Chen, J., Zhou, F., Shou, L., Chen, Q., Tao, B., et al. (2015). Controlling factors of
  summer phytoplankton community in the Changjiang (Yangtze River) Estuary and
  adjacent East China Sea shelf. *Continental Shelf Research*, 101, 1–14.
  http://doi.org/10.1016/j.csr.2015.04.009
- Ji, R., C. Chen, J. Budd, D. Schwab, D. Beletsky, D. Fahanenstial, T. H. Johengen, H.
  Lavrentyev, B. Eadies, J. Cotner, W. Gardner and M. Bundy (2002). A coupled
  biological and physical model study of the ecosystem in Lake Michigan Part II: Influence
  of suspended sediment. *Ecological Modeling*. 152, 169-190.
- Klausmeier C.A., Litchman E., Daufresne T. and Levin S. (2008). Phytoplankton stoichiometry.
   *Ecol. Res.*, 23, 479–485.
- Lamquin, N., C. Mazeran, D. Doxaran, J. H. Ryu, and Y. J. Park (2012), Assessment of GOCI
   radiometric products using MERIS, MODIS and field measurements, *Ocean Science Journal*, 47(3), 287-311
- Liu, S.-M., Qi, X. H., Li, X., Ye, H. R., Wu, Y., Ren, J. L., et al. (2016). Nutrient dynamics from
  the Changjiang (Yangtze River) estuary to the East China Sea. *Journal of Marine Systems*, 154(Part A), 15–27. http://doi.org/10.1016/j.jmarsys.2015.05.010
- Luan, H. L., Ding, P.-X., Wang, Z. B., Ge, J.-Z., & Yang, S.-L. (2016). Decadal morphological
  evolution of the Yangtze Estuary in response to river input changes and estuarine
  engineering projects. *Geomorphology*, 265(C), 12–23.
  http://doi.org/10.1016/j.geomorph.2016.04.022
- Luo, Z., Zhu, J., Wu, H., & Li, X. (2017). Dynamics of the sediment plume over the Yangtze
  Bank in the Yellow and East China Seas. *Journal of Geophysical Research: Oceans*, 122,
  10,073–10,090. https://doi.org/10.1002/2017JC013215
- Niemistö, J., Kononets, M., Ekeroth, N., Tallberg, P., Tengberg, A., & Hall, P. O. J. (2018).
  Benthic fluxes of oxygen and inorganic nutrients in the archipelago of Gulf of Finland,
  Baltic Sea Effects of sediment resuspension measured in situ. *Journal of Sea Research*,
  135, 95–106. http://doi.org/10.1016/j.seares.2018.02.006
- Qi, J., Chen, C., Beardsley, R., Perrie, W., Cowles, G., & Lai, Z. (2009). An unstructured-grid
   finite-volume surface wave model (FVCOM-SWAVE): Implementation, validations and
   applications. *Ocean Modelling*, 28(1-3), 153–166.
- Qi, J., Chen, C., & Beardsley, R. C. (2018). FVCOM one-way and two-way nesting using
   ESMF\_Development and validation. *Ocean Modelling*, 124, 94–110.
   http://doi.org/10.1016/j.ocemod.2018.02.007
- Olsen, A., Key, R. M., van Heuven, S., Lauvset, S. K., Velo, A., Lin, X., et al. (2016). The
  Global Ocean Data Analysis Project version 2 (GLODAPv2) an internally consistent
  data product for the world ocean. *Earth System Science Data*, 8(2), 297–323.
  http://doi.org/10.5194/essd-8-297-2016

Piwowarczyk, L., H. Bishop, K. Saia, S. Crosby, F. T. Mudymba, N. I. Hashi, and A. Raj (2016), 834 Application of GOCI Satellite Data to Ocean Modeling, J. Coastal Res., 15(1), 1409-835 1414. 836 Ryu, J.-H., Han, H.-J., Cho, S., Park, Y.-J., & Ahn, Y.-H. (2012). Overview of geostationary 837 ocean color imager (GOCI) and GOCI data processing system (GDPS). Ocean Science 838 Journal, 47(3), 223–233. http://doi.org/10.1007/s12601-012-0024-4 839 Sadat-Noori, M., Santos, I. R., Tait, D. R., & Maher, D. T. (2016). Fresh meteoric versus 840 841 recirculated saline groundwater nutrient inputs into a subtropical estuary. Science of the Total Environment, the, 566-567(C), 1440-1453. 842 http://doi.org/10.1016/j.scitotenv.2016.06.008 843 Shen, F., W. Verhoef, Y. Zhou, M. S. Salama, and X. Liu (2010), Satellite Estimates of Wide-844 Range Suspended Sediment Concentrations in Changjiang (Yangtze) Estuary Using 845 846 MERIS Data, Estuar. Coast., 33(6), 1420-1429, doi:10.1007/s12237-010-9313-2 Shen, F., Zhou, Y., Li, J., He, Q., & Verhoef, W. (2013). Remotely sensed variability of the 847 848 suspended sediment concentration and its response to decreased river discharge in the Yangtze estuary and adjacent coast. Continental Shelf Research, 69(C), 52-61. 849 http://doi.org/10.1016/j.csr.2013.09.002 850 Stow, C. A., Jolliff, J., McGillicuddy, D. J., Jr., Doney, S. C., Allen, J. I., Friedrichs, M. A. M., 851 et al. (2009). Skill assessment for coupled biological/physical models of marine systems. 852 Journal of Marine Systems, 76(1-2), 4-15. http://doi.org/10.1016/j.jmarsys.2008.03.011 853 Sun, X., F. Shen, D. Liu, R. G. Bellerby, Y. Liu, and R. Tang (2018), In Situ and Satellite 854 Observations of Phytoplankton Size Classes in the Entire Continental Shelf Sea, China, J. 855 Geophys. Res. Oceans, 123(5), 3523-3544, 856 857 Siswanto, E., Tang, J., Yamaguchi, H., Ahn, Y.-H., Ishizaka, J., Yoo, S., et al. (2011). Empirical ocean-color algorithms to retrieve chlorophyll-a, total suspended matter, and colored 858 dissolved organic matter absorption coefficient in the Yellow and East China Seas. 859 Journal of Oceanography, 67(5), 627-650. http://doi.org/10.1007/s10872-011-0062-z 860 Sokoletsky, L., Yang, X., & Shen, F. (2014). MODIS-based retrieval of suspended sediment 861 concentration and diffuse attenuation coefficient in Chinese estuarine and coastal waters. 862 In R. J. Frouin, D. Pan, & H. Murakami (Eds.), (Vol. 9261, p. 926119). Presented at the 863 SPIE Asia Pacific Remote Sensing, SPIE. http://doi.org/10.1117/12.2069205 864 O'Reilly, J. E., S. Maritorena, B. G. Mitchell, D. A. Siegel, K. L. Carder, S. A. Garver, M. 865 Kahru, and C. McClain, (1998). Ocean color chlorophyll algorithm for SeaWiFS. Journal 866 of Geophysical Research, 103(C11): 24937-24953. 867 Olsen A, Key RM, van Heuven S, Lauvset SK, Velo A, Lin X, Schirnick C, Kozyr A, Tanhua T, 868 Hoppema M, Jutterstro m S, Steinfeldt R, Jeansson E, Ishii M, Pe rez FF, Suzuki T 869 (2016). The Global Ocean Data Analysis Project version 2 (GLODAPv2)-aninternally 870 consistent data product for the world ocean. Earth Syst Sci Data 8:297-323. 871 https://doi.org/10.5194/essd-8-297-2016 872 Moore, W. S. (2010). The Effect of Submarine Groundwater Discharge on the Ocean. Annual 873 Review of Marine Science, 2(1), 59-88. http://doi.org/10.1146/annurev-marine-120308-874 081019 875

Moore, W.S., J.O. Blanton, S. Joye (2006). Estimates of Flushing Times, Submarine 876 Groundwater Discharge, and Nutrient Fluxes to Okatee River, South Carolina, J. 877 Geophys. Res., 111, C09006, doi:10.1029/2005JC003041,. 878 Moore, W.S. (2006). The role of submarine groundwater discharge in coastal biogeochemistry. 879 Journal of Geochemical Exploration, 88, 389-393,. 880 Hu, C., L. Feng, and Z. P. Lee (2012), Evaluation of GOCI sensitivity for At-Sensor radiance 881 and GDPS-Retrieved chlorophyll-a products, Ocean Science Journal, 47(3), 279-285, 882 Li, M., Xu, K., Watanabe, M., & Chen, Z. (2007). Long-term variations in dissolved silicate, 883 nitrogen, and phosphorus flux from the Yangtze River into the East China Sea and 884 impacts on estuarine ecosystem. Estuarine, Coastal and Shelf Science, 71(1-2), 3-12. 885 http://doi.org/10.1016/j.ecss.2006.08.013 886 887 Liu, J., Su, N., Wang, X., & Du, J. (2017). Submarine groundwater discharge and associated nutrient fluxes into the Southern Yellow Sea: A case study for semi-enclosed and 888 oligotrophic seas-implication for green tide bloom. Journal of Geophysical Research-889 890 Oceans, 122(1), 139–152. http://doi.org/10.1002/2016JC012282 Liu, J., Du, J., Wu, Y., & Liu, S. (2018). Nutrient input through submarine groundwater 891 discharge in two major Chinese estuaries: the Pearl River Estuary and the Changjiang 892 River Estuary. Estuarine, Coastal and Shelf Science, 203, 17–28. 893 http://doi.org/10.1016/j.ecss.2018.02.005 894 Vanderploeg, H. A., Johengen, T. H., Lavrentyev, P. J., Chen, C., Lang, G. A., Agy, M. A., et al. 895 (2007). Anatomy of the recurrent coastal sediment plume in Lake Michigan and its 896 897 impacts on light climate, nutrients, and plankton. Journal of Geophysical Research, 112(C3), C03S90. http://doi.org/10.1029/2004JC002379 898 899 Wang, X., Baskaran, M., Su, K., & Du, J. (2018). The important role of submarine groundwater discharge (SGD) to derive nutrient fluxes into River dominated Ocean Margins – The 900 East China Sea. Marine Chemistry, 204, 121–132. 901 http://doi.org/10.1016/j.marchem.2018.05.010 902 903 Wang, Y., Wu, H., Gao, L., Shen, F., & Liang, X. S. (2019). Spatial Distribution and Physical Controls of the Spring Algal Blooming Off the Changjiang River Estuary. Estuaries and 904 Coasts, 42(4), 1–18. http://doi.org/10.1007/s12237-019-00545-x 905 Wu, L., Chen, C., Guo, P., Shi, M., Qi, J., & Ge, J. (2011). A FVCOM-based unstructured grid 906 wave, current, sediment transport model, I. Model description and validation. Journal of 907 Ocean University of China, 10(1), 1-8. http://doi.org/10.1007/s11802-011-1788-3 908 Wu, H., Zhu, J., Shen, J., & Wang, H. (2011). Tidal modulation on the Changjiang River plume 909 in summer. Journal of Geophysical Research, 116(C8), C08017. 910 http://doi.org/10.1029/2011JC007209 911 Yang, S. L., Xu, K. H., Milliman, J. D., Yang, H. F., & Wu, C. S. (2015). Decline of Yangtze 912 River water and sediment discharge: Impact from natural and anthropogenic changes. 913 Scientific Reports, 1-14. http://doi.org/10.1038/srep12581 914 Zhu, Z.-Y., Ng, W.-M., Liu, S.-M., Zhang, J., Chen, J.-C., & Wu, Y. (2009). Estuarine 915 916 phytoplankton dynamics and shift of limiting factors: A study in the Changjiang (Yangtze

River) Estuary and adjacent area. Estuarine, Coastal and Shelf Science, 84(3), 393-401. 917 http://doi.org/10.1016/j.ecss.2009.07.005 918 919 Zhu, Z.-Y., Wu, Y., Zhang, J., Du, J.-Z., & Zhang, G.-S. (2014). Reconstruction of anthropogenic eutrophication in the region off the Changjiang Estuary and central Yellow 920 Sea From decades to centuries. Continental Shelf Research, 72(C), 152–162. 921 922 http://doi.org/10.1016/j.csr.2013.10.018 Zhou, M.-J., Shen, Z.-L., & Yu, R.-C. (2008). Responses of a coastal phytoplankton community 923 924 to increased nutrient input from the Changjiang (Yangtze) River. Continental Shelf Research, 28(12), 1483-1489. http://doi.org/10.1016/j.csr.2007.02.009 925 Zhou, Y., Zhang, Y., Li, F., Tan, L., & Wang, J. (2017). Nutrients structure changes impact the 926 competition and succession between diatom and dinoflagellate in the East China Sea. 927 Science of the Total Environment, the, 574(C), 499–508. 928 929 http://doi.org/10.1016/j.scitotenv.2016.09.092 Zhou, Z.-X., Yu, R.-C., Sun, C., Feng, M., & Zhou, M.-J. (2019). Impacts of Changjiang River 930 931 Discharge and Kuroshio Intrusion on the Diatom and Dinoflagellate Blooms in the East China Sea. Journal of Geophysical Research-Oceans, 8(1), 39–14. 932 http://doi.org/10.1029/2019JC015158 933

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