| 1 | Optimization of Biological Production for Indian Ocean upwelling zones: Part – I: |
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| 2 | Improving Biological Parameterization via a variable Compensation Depth |
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- 21 Abstract
- 22

Biological modeling approach adopted by the Ocean Carbon-cycle Model Inter-comparison 23 24 Project (OCMIP-II) provided amazingly simple but surprisingly accurate rendition of the annual mean carbon cycle for the global ocean. Nonetheless, OCMIP models are known to have 25 seasonal biases which are typically attributed to their bulk parameterization of 'compensation 26 depth'. Utilizing the criteria of surface Chl-a based attenuation of solar radiation and the 27 minimum solar radiation required for production, we have proposed a new parameterization for a 28 spatially and temporally varying 'compensation depth' which captures the seasonality in the 29 production zone reasonably well. This new parameterization is shown to improve the seasonality 30 of CO_2 fluxes, surface ocean p CO_2 , biological export and new production in the major upwelling 31 32 zones of the Indian Ocean. The seasonally varying compensation depth enriches the nutrient 33 concentration in the upper ocean yielding more faithful biological exports which in turn leads to 34 an accurate seasonality in the carbon cycle. The export production strengthens by $\sim 70\%$ over the 35 western Arabian Sea during monsoon period and achieves a good balance between export and 36 new production in the model. This underscores the importance of having a seasonal balance in 37 model export and new productions for a better representation of the seasonality of carbon cycle 38 over upwelling regions. The study also implies that both the biological and solubility pumps play 39 an important role in the Indian Ocean upwelling zones.

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Keywords: Indian Ocean upwelling zones, Carbon cycle, Seasonal cycle - CO₂ flux and Oceanic
pCO₂, Biogeochemical model parameterization, Export production - New production balance,
Solubility and Biological pump.

44

1. Introduction

46 The Indian Ocean is characterized by the unique seasonally reversing monsoon wind systems which act as the major physical drivers for the coastal and open ocean upwelling processes. The 47 major upwelling systems in the Indian Ocean are (1) the western Arabian Sea (WAS; Ryther and 48 49 Menzel, 1965; Smith et al., 2001; Sarma, 2004; Wiggert et al., 2005, 2006; Murtugudde et al., 2007; McCreary et al., 2009; Prasanna Kumar et al., 2010; Naqvi et al., 2010; Roxy et al., 2015) 50 (2) the Sri Lanka Dome (SLD; Vinayachandran et al., 1998, 2004), (3) Java and Sumatra coasts 51 (SC; Murtugudde et al., 1999; Susanto et al., 2001; Osawa et al., 2010; Xing et al., 2012) and (4) 52 the Seychelles-Chagos thermocline ridge (SCTR; Murtugudde et al., 1999; Dilmahamod et al., 53 54 2016, Figure 1). The physical and biological processes and their variability over these key 55 regions are inseparably tied to the strength of the monsoon winds and associated nutrient dynamics. The production and its variability over these coastal upwelling systems are a key 56 57 concern for the fishing community, since they affect the day-to-day livelihood of the coastal populations (Harvell et al., 1999; Roxy et al., 2015; Praveen et al., 2016) and are important for 58 the Indian Ocean rim countries due to their developing country status. 59

Arabian Sea is a highly productive coastal upwelling system characterized by phytoplankton 60 blooms both in summer (Prasanna Kumar et al., 2001; Naqvi et al., 2003; Wiggert et al., 2005) 61 62 and winter (Banse and McClain, 1986; Wiggert et al., 2000; Barber et al., 2001; Prasannakumar et al., 2001; Sarma, 2004). The Arabian Sea is known for the second largest Tuna fishing region 63 in the Indian Ocean (Lee et al., 2005). The Somali and Omani upwelling regions experience 64 phytoplankton blooms that are prominent with Net Primary Production (NPP) exceeding 435 g C 65 m⁻² yr⁻¹ (Liao et al., 2016). On the other hand productivity over the SLD (Vinayachandran and 66 Yamagata, 1998) is triggered by open ocean Ekman suction with strong Chl-a blooms during the 67

summer monsoon (Murtugudde et al., 1999; Vinayachandran et al., 2004). Similarly, the SC 68 upwelling is basically due to the strong alongshore winds and its variation is associated with the 69 impact of equatorial and coastal Kelvin waves (Murtugudde et al., 2000, Valsala and Rao, 2016). 70 The interannual variability associated with the Java-Sumatra coastal upwelling is strongly 71 coupled with ENSO (El-Niño Southern Oscillation) through the Walker cell and Indonesian 72 73 throughflow (Susanto et al., 2001; Valsala et al., 2011) and peaks in July through August with a potential new production of 0.1 Pg C yr⁻¹ (Xing et al., 2012). The SCTR productivity has a large 74 spatial and interannual variability. The warmer upper ocean condition associated with El Niño 75 76 reduces the amplitude of subseasonal SST variability over the SCTR (Jung and Kirtman., 2016). The Chl-a concentration peaks in summer when the southeast trade winds induce mixing and 77 initiate the upwelling of nutrient-rich water (Murtugudde et al., 1999; Wiggert et al., 2006; 78 Vialard et al., 2009; Dilmahamod et al., 2016). 79

80 Understanding the biological production and variability in the upwelling systems is important because it gives us crucial information regarding marine ecosystem variability (Colwell, 1996; 81 Harvell et al., 1999). The observations also provide vital insights into physical and biological 82 interactions of the ecosystem (Naqvi et al., 2010) as well as the biophysical feedbacks 83 (Murtugudde et al., 2001), although limitations of sparse observations often force us to depend 84 on models to examine the large spatio-temporal variability of the ecosystem (Valsala et al., 85 2013). Simple to intermediate complexity marine ecosystem models have been employed by 86 several of the previous studies (Sarmiento et al., 2000; Orr et al., 2001; Matsumoto et al., 2008). 87 However, the representation of marine ecosystem with proper parameterizations in models has 88 always been a daunting task. This is an impediment to the accurate representation of biological 89

primary and export productions in models (Friedrichs et al., 2006, 2007) and these issues also
impact the modeling of upper trophic levels (Lehodey et al., 2010).

Biological production can be quantified with a better understanding of primary production 92 which depends on water temperature, light and nutrient availability (Brock et al., 1993; Moisan 93 94 et al., 2002) and this became the key reason for parameterizing the production in models as one 95 or more combinations of these terms (Yamanaka et al., 2004). Any of these basic parameters can be tweaked to alter production in models. For example, the availability of nutrients and light 96 determines the phytoplankton growth (Eppely et al., 1972) or growth rate (Boyd et al., 2013). 97 98 Stoichiometry and carbon-to-Chl-a ratios are other important factors to be considered in 99 modeling (Christian et al., 2001, Wang et al., 2009) but we will not consider them in this study.

100 The Ocean Carbon-cycle Model Intercomparison Project (OCMIP) greatly improved our understanding of global carbon cycle (Najjar and Orr, 1998). OCMIP-II further introduced a 101 simple phosphate-dependent production term in biological models for long-term simulations of 102 103 the carbon cycle in response to anthropogenic climate change with an accurate annual mean state (Najjar and Orr, 1998; Orr et al., 2001; Doney et al., 2004). However, the OCMIP – II model 104 105 simulations come with a penalty of strong seasonal biases when compared with observations (Orr et al., 2003). In this protocol, the community compensation depth (hereinafter Zc) is defined 106 107 as the depth at which photosynthesis equals entire community respiration and the irradiance at which this balance is achieved is the compensation irradiance (E_{com}). Note that Zc is clearly 108 different from the conventional euphotic zone depth (Morel, 1988). At Zc, the Net Community 109 110 Production is zero and above Zc the NPP exceeds the community respiration and the ecosystem 111 will grow (Smetacek and Passow, 1990; Gattuso et al., 2006; Sarmiento and Gruber, 2006; Regaudix-de-Gioux and Duarte, 2010; Marra et al., 2014). Therefore the region above Zc 112

113 represents the production zone in this modeling approach. However, Zc was held constant in time and space in OCMIP-II models (Najjar and Orr, 1998; Matsumoto et al., 2008) because the 114 OCMIP-II protocol takes the minimalistic approach to biology and simplifies the model 115 calculations with a very limited set of state variables suitable for long term simulations when 116 casted in coarse resolution models (Orr et al., 2005). Though in reality Zc varies in space and 117 time (Najjar and Keeling, 1997) just as the euphotic zone depth does as documented in ship 118 measurements (Qasim, 1977, 1982). The variation in Zc indicates the seasonality of the 119 production zone itself. 120

121 Most of the biophysical models prescribe a constant value for Zc, e.g., a default value of Zc = 75 m in OCMIP –II protocol (Najjar and Orr, 1998) and Zc = 100 m in Minnesota Earth 122 System Model (Matsumoto et al., 2008). Depending on the latitude, Zc varies between 50m and 123 100m in the real world (Najjar and Keeling, 1997). In our study we have attempted a novel 124 125 biological parameterization scheme for spatially and temporally varying Zc in the OCMIP-II framework by representing the production as a function of solar radiation (Parsons et al., 1984) 126 and prescribed Chl-a. In this hypothesis, a spatially and temporally varying Zc is estimated from 127 the vertical attenuation of insolation by the surface Chl-a. The depth at which the insolation 128 reaches the compensation irradiance (chosen as 10 Wm⁻²) is taken as Zc. Phosphorous is the 129 basic currency which limits the production within this varying Zc. This spatially and temporally 130 varying Zc represents the seasonality in the production zone which is lacking in the original 131 OCMIP-II protocol. 132

Regions of sustained upwelling like the eastern equatorial Pacific are well understood in terms of the role of upwelling in increasing the surface water pCO_2 to drive an outgassing of CO_2 into the atmosphere (Feely et al., 2001; Valsala et al., 2014). The Indian Ocean on the other hand 136 experiences only seasonal upwelling which is relatively weak in the deep tropics but stronger off 137 the coasts of Somalia and Oman and in the SLD (Valsala et al., 2013). The relative importance of the solubility vs. biological pump is not well understood. Our focus here on implementing 138 139 seasonality in Zc of OCMIP models nonetheless leads to new insights on the impact of improved biological production on surface water pCO₂ and air-sea CO₂ fluxes. The improvements due to 140 the effect of a variable Zc over the Indian Ocean and the sensitivity experiments where 141 upwelling is muted strongly imply that the biological pump may play as much of a role as the 142 solubility pump in determining surface pCO₂ and CO₂ fluxes over the Indian Ocean. 143

The paper is organized as follows. Model, Data, and Methods are detailed in Section 2. The spatially inhomogeneous Zc derived out of the new parameterization and its impact on simulated seasonality of biology and carbon cycle are detailed in Section 3. Further results and discussion are followed in Section 4 and a conclusion is given in Section 5.

148

149 **2. Model, Data, and Methods**

150 **2.1. Model**

The study utilizes the Offline Ocean Tracer Transport Model (OTTM; Valsala et al., 2008) coupled with OCMIP biogeochemistry model (Najjar and Orr, 1998). OTTM does not compute currents and stratifications (i.e., temperature and salinity) on its own. It is capable of accepting any ocean model or data-assimilated product as physical drivers. The physical drivers prescribed include 4-dimensional currents (u,v), temperature, salinity, and 3-dimensional mixed layer depth, surface freshwater and heat fluxes, surface wind stress and sea surface height. The resolution of the model setup is similar to the parent model from which it borrows the physical drivers. With 158 the given input of Geophysical Fluid Dynamics Laboratory (GFDL) reanalysis data (Chang et. 159 al., 2012), the zonal and meridional resolutions are 1° with 360 grid points longitudinally and 1° at higher latitudes but having a finer resolution of 0.8° in the tropics, with 200 latitudinal grid 160 points. The model has 50 vertical levels with 10m increment in the upper 225m and stretched 161 vertical levels below 225m. The horizontal grids are formulated in spherical coordinates and 162 vertical grids are in z levels. The model employs a B-grid structure in which the velocities are 163 resolved at corners of the tracer grids. The model uses a centered-in-space and centered-in-time 164 (CSCT) numerical scheme along with an Asselin-Robert filter (Asselin, 1972) to control the 165 166 ripples in CSCT.

167 The tracer concentration (C) evolves with time as

168
$$\frac{\partial C}{\partial t} + U.\nabla_{\rm H}C + W\frac{\partial C}{\partial z} = \frac{\partial}{\partial z}K_{\rm z}\frac{\partial}{\partial z}C + \nabla_{\rm H}.(K_{\rm h}\nabla_{\rm H}C) + J + F$$
 (1)

where $\nabla_{\rm H}$ is the horizontal gradient operator, U and W are the horizontal and vertical velocities respectively. K_z is the vertical mixing coefficient, and K_h is the two-dimensional diffusion tensor. J represents any sink or source due to the internal consumption or production of the tracer. F represents the emission or absorption of fluxes at the ocean surface. Here, the source and sink terms are provided through the biogeochemical model. Vertical mixing is resolved in the model using K- profile parameterization (KPP) (Large et al., 1994).

In addition to KPP, the model uses a background vertical diffusion reported by Bryan and Lewis (Bryan and Lewis, 1979). For horizontal mixing, model incorporates Redi fluxes (Redi, 1982) and GM fluxes (Gent and Mcwilliams, 1990) which represent the eddy-induced variance in the mean tracer transport. A weak Laplacian diffusion is also included in the model for computational stability where the sharp gradient in concentration occurs.

180 2.2. Biogeochemical model

181 The biogeochemical model used in the study is based on the OCMIP - II protocol as stated above. The main motivation of OCMIP-II protocol is to employ a minimalistic approach 182 183 to simulate the ocean carbon cycle with a nutrient restoration approach to calculate the oceanic 184 biological production (Najjar et al., 1992; Anderson and Sarmiento, 1995). The present version of the model has four prognostic variables coupled with the circulation field, viz., inorganic 185 phosphate (PO_4^{3-}), dissolved organic phosphorus (DOP), dissolved inorganic carbon (DIC) and 186 alkalinity (ALK). The basic currency for the biological model is phosphorous because of the 187 availability of a more extensive phosphate database and to eliminate the complexities associated 188 with nitrogen fixation and denitrification. 189

190 The production of organic phosphorus in the model using the nutrient restoring approach is given191 by

192
$$J_{prod} = \frac{1}{\tau} ([PO_4] - [PO_4]^*)$$
 (2)

193
$$[PO_4] > [PO_4]^*; Z < Zc$$

 $194 \quad J_{prod} = 0 \tag{3}$

195
$$[PO_4] \le [PO_4]^*; Z > Zc$$

196 Where $[PO_4]^*$ is the observed phosphate concentration and $\tau=30$ days is the restoration time scale 197 (Najjar et. al., 1992).

198 The vertically integrated new production (g C m^{-2} yr⁻¹) in the model is defined as

199 New production =
$$\int_{zc}^{0} -J_{prod} dz$$
 (4)

200 The export production (g C m^{-2} yr⁻¹) in the model is calculated as given below

201 Export production =
$$(1 - \sigma) \int_0^{zc} J_{prod} dz$$
 (5)

202 Air-sea CO_2 flux in the model is estimated by,

$$203 F = K_w \Delta p C O_2 (6)$$

where K_w is gas transfer velocity and $\triangle pCO_2$ is the difference in partial pressure of carbon dioxide between the ocean and atmosphere.

 pCO_2 is calculated in the model by using DIC and ALK and is given by,

207
$$pCO_2 = \frac{[DIC]}{K_0} \frac{[H^+]^2}{[H^+]^2 + K_1[H^+] + K_1K_2}$$
 (7)

Where $[H^+]$ is calculated using Newton-Raphson iterative method (Press et. al., 1996, Najjar and Orr, 1998). K_0 is the solubility constant of CO₂ and K_1 , K_2 are the dissociation constant for carbonic acid respectively (Sarmiento and Gruber, 2006, Weiss, 1974, Mehrbach et al., 1973, Dickson and Millero, 1987, Najjar and Orr, 1998).

Details of all parameters in the biogeochemical model and calculations of solubility and biological pump are provided in Appendix-A. The design and validation of the physical model is reported by Valsala et al., (2008, 2010) and biogeochemical model by Najjar and Orr (1998).

215

216 **2.3. Data**

217 For validating the model results, observational datasets of CO_2 flux and p CO_2 are taken from Takahashi et al., (2009). Satellite-derived Net Primary Production (NPP) data were taken 218 from Sea-viewing Wide Field of view Sensor (SeaWiFS) Chl-a product, calculated using 219 220 Vertically Generalized Production Model (VGPM) (Behrenfeld and Falkowski, 1997). The NPP data is scaled to export production (EP) by multiplying with an e-ratio (e = 0.37) representative 221 of Indian Ocean upwelling zones (Sarmiento and Gruber, 2006, Laws et al., 2000; Falkowski et 222 al., 2003). The initial conditions for PO₄ are taken from the World Ocean Atlas (Garcia et al., 223 2014). Initial conditions for DIC and ALK are taken from the Global Ocean Data Analysis 224 Project (GLODAP; Key et al., 2004) dataset. The dissolved Organic Phosphorous (DOP) is 225 initialized with a constant value of 0.02 µmol kg⁻¹ (Najjar and Orr, 1998). The data sources and 226 citations are provided in the Acknowledgement. 227

228

229 **2.4. Methods**

A spin-up for 50 years from the given initial conditions is performed with the climatological physical drivers. As the initial conditions are provided from a mean state of observed climatology this duration of spin-up is sufficient to reach statistical equilibrium in the upper 1000 m (Le Quere et al., 2000). Atmospheric pCO₂ has been set to a value from the 1950s in the spin-up run for calculating the air-sea CO₂ exchange. A seasonal cycle of atmospheric pCO₂ has been prescribed as in Keeling et al. (1995).

After the spin-up, an interannual simulation for 50 years from 1961 to 2010 has been carried out with the corresponding observed atmospheric pCO_2 described in Keeling et al., (1995). The first five years of the interannual run were looped five times through the physical

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fields of 1961 repeatedly for a smooth merging of the spin-up restart to the interannual physical variables. Since the study is focused on bias corrections to the seasonal cycle of pCO_2 and DIC with a variable Zc, a model climatology for carbon cycle has been constructed from 1990 to 2010, which includes the anthropogenic increase of oceanic DIC in the climatological calculation and is comparable with the Takahashi et al. (2009) observations.

Additional two sensitivity experiments have been performed separately by providing 244 annual mean currents or temperatures as drivers over selected regions of the basin in order to 245 segregate the role of varying Zc in improving the seasonality of carbon cycle. The aim of these 246 sensitivity experiments is to understand how successful the new parameterization for Zc is in 247 248 capturing the carbon cycle variability related to the upwelling episodes even though the seasonal 249 cycle in physics is suppressed. The model driven with annual mean currents suppresses the effect of upwelling by muting the Ekman divergence over the region of interest. On the other 250 251 hand, the model forced with annual mean temperatures suppresses the cooling effect of upwelling. A smoothing technique with linear interpolation $(u = u(1 - x) + \overline{u}x)$ is applied to 252 253 the offline-data in order to blend the annual mean fields (\bar{u}) provided to the selected region with the rest of the domain (u) in order to reduce a sudden transition at the boundaries. Here x 254 represents an index which varies between 0 and 1 within a distance of 10^{0} from the boundaries of 255 the region of interest to the rest of the model domain. 256

257

258 2.5. Community compensation depth (Zc) parameterization

259 The OCMIP – II protocol separates the production and consumption zones by a depth termed
260 as compensation depth (Zc); the depth at which photosynthesis is large enough to balance the

community respiration (i.e., both the autotrophic and heterotrophic respiration). At the 261 262 community compensation depth, the Net Community Production (NCP) is zero i.e., NCP = NPP - $R_h = 0$, where NPP is Net Primary Production (i.e., NPP = GPP - R_a), GPP is gross primary 263 production, and R_h and Ra are the heterotrophic and autotrophic respirations, respectively 264 (Smetacek and Passow, 1990; Najjar and Orr, 1998; Gattuso et al., 2006; Regaudix-de-Gioux 265 and Duarte, 2010; Marra et al., 2014). The light intensity at Zc is compensation irradiance (E_{com}), 266 the irradiance at which the gross community primary production balances respiratory carbon 267 losses for the entire community (Gattuso et al., 2006; Regaudix-de-Gioux and Duarte, 2010). We 268 define a spatially and temporally varying compensation depth (hereinafter varZc) as a depth 269 where compensation irradiance (attenuated by surface Chl-a, Jerlov et al., 1976) reaches a 270 minimum value of 10 W m⁻². In this way, the varZc has both spatio-temporal variability of light 271 272 as well as Chl-a. The Chl-a is given as monthly climatology as constructed from satellite data. Observations show that the primary production reduces rapidly to 20% or less of the surface 273 value below a threshold of 10 W m⁻² (Parsons et al., 1984; Ryther, 1956; Sarmiento and Gruber, 274 275 2006). Moreover higher ocean temperatures (those in the tropics) enhance the respiration rates resulting in high compensation irradiance (Parsons et al., 1984; Ryther, 1956; Lopez-Urrutia et 276 al., 2006; Regaudix-de-Gioux and Duarte, 2010). A study by Regaudix-de-Gioux and Duarte 277 (2010) reported the mean value of compensation irradiance over the Arabian Sea as 0.4 ± 0.2 278 mol photon $m^{-2} day^{-1}$ which is close to 10 W $m^{-2} day^{-1}$. 279

Figure 2 compares the scatter of average relative photosynthesis within varZc as a function of solar radiation for the Indian Ocean (see Appendix-B). This encapsulates the corresponding curve from the observations for the major phytoplankton species in the ocean such as diatoms, green algae and dinoflagellates (Ryther et al., 1956; Parsons et al., 1984; Sarmiento and Gruber, 2006). The model permits 100% production of organic phosphorus for radiation above 50 W m⁻². However the availability of phosphate concentration in the model acts as an additional limit for production which indirectly represents the photoinhibition at higher irradiance; for example, in the oligotrophic gyres.

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289 3. Results and Discussions

The inclusion of seasonality in Zc by way of parameterizing varZc leads to a remarkable spatio-temporal variability in Zc (Figure 3). Zc over the Arabian Sea varies from 10 m to 25 m during December to February (DJF) and deepens down to 45 m during March to May (MAM) due to the increase in the surface solar radiation. During the monsoon season i.e., June to September (JJAS), Zc again shoals to 10 m - 35 m due to the attenuation of solar radiation by the increased biological production (Chl-a). During October to November (OCT-NOV), Zc slightly deepens as compared to JJAS.

The Bay of Bengal Zc deepens from 35 m to 40 m during DJF and further deepens to 50 m during MAM when the solar radiation is maximum and biological production is minimum (Prasannakumar et al., 2002). Further reduction of Zc can be seen through JJAS as a result of a reduction in solar radiation during monsoon cloud cover. Zc during OCT-NOV is 35 m on average.

The equatorial Indian Ocean can be seen as a belt of 40 m - 45 m Zc throughout the season except for JJAS. During JJAS, a shallow Zc is seen near the coastal Arabian Sea (around 10 m to 304 35 m) presumably due to the coastal Chl-a blooms. Deep Zc off the coast of Sumatra (~ 40 m to 305 50 m) is found during JJAS. Java-Sumatra coastal upwelling is centered on September to November (Susanto et al., 2001) and upwelling originates at around 100 m depth (Valsala and
Maksyutov, 2010; Xing et al., 2012).

Southward of 10°S in the oligotrophic gyre region, Zc varies from 40 m to more than 60 m 308 throughout the year. A conspicuous feature observed while parameterizing the solar radiation 309 310 and Chl-a dependent Zc is that its maximum value never crosses 75 m especially in the Indian Ocean which is the value specified in OCMIP-II models. The cutoff depth of 75 m in OCMIP-II 311 is obtained from observing the seasonal variance in oxygen data (Najjar and Keeling, 1997) as an 312 indicator of the production zone. However, our results show that parameterizing a production 313 zone based on solar radiation and Chl-a predicts a production zone and its variability that is 314 315 largely less than 75 m. The relevance of varZc in the seasonality of the modeled carbon cycle is illustrated as follows. 316

317

318 **3.1. Simulated seasonal cycle of pCO₂ and CO₂ fluxes**

The annual mean biases in simulated CO_2 fluxes and pCO_2 were evaluated by comparing with Takahashi et al., (2009) observations (Figure 4). The model biases are significantly reduced with the implementation of varZc compared to that of the constant Zc (hereinafter constZc). A notable reduction in pCO_2 bias (by ~ 10µatm) is observed along the WAS (Figure 4d).

In order to address the role of the new biological parameterization of a variable Zc, we have extended our study by choosing four key regions where the biological production and CO_2 fluxes are prominent in the Indian Ocean with additional sensitivity experiments (see Introduction and references therein). The regions (boxes shown in Figure 1) we considered are, (1) Western Arabian Sea (WAS; 40°E:65°E, 5°S:25°N) (2) Sri Lanka Dome (SLD; 81°E:90°E, 328 0°:10°N) (3) Seychelles-Chagos Thermocline Ridge (SCTR; 50°E:80°E, 5°S:10°S) and (4)
329 Sumatra Coast (SC; 90°E:110°E, 0°:10°S; Figure 1). The seasonal variations of Zc over these
330 selected key regions are shown in Figure 5. A detailed analysis of CO₂ fluxes, pCO₂, biological
331 export and new productions and the impact of varZc simulations in improving the strength of
332 biological pump and solubility pump for these key regions are presented below.

333

334 **3.2. Western Arabian Sea (WAS)**

335 The WAS Zc has a double peak pattern over the annual cycle. During the February-March 336 period, Zc deepens down to a maximum of 43.85 ± 2.3 m into March and then shoals to $25.75 \pm$ 1.5 m (Figure 5) during the monsoon period (uncertainty represents the interannual standard 337 deviations of monthly data from 1990-2010). This shoaling of Zc depth during the monsoon 338 339 indicates the potential ability of the present biological parameterization to capture the wind-340 driven upwelling related production in the WAS. During the post-monsoon period, the second 341 deepening of Zc occurs during November with a maximum depth of 34.91 ± 2.2 m. The ability to 342 represent the seasonality of the production zone renders a unique improvement in CO₂ flux variability especially in the WAS in comparison to the OCMIP-II experiments (Orr et al, 2003; 343 Figure 6a). 344

OCMIP –II simulations with a constZc of 75 m underestimate the CO₂ flux when compared to the observations of Takahashi et al. (2009). This underestimation is clearly visible during the monsoon period. Our simulations with the varZc resulted in a better seasonality of CO₂ flux when compared with Takahashi et al. (2009) observations (Figure 6a). The improvement due to the varZc scheme is able to represent the seasonality of CO₂ flux better especially during the monsoon period when wind-driven upwelling is dominant. Obviously, the relative role of thebiological and solubility pumps have to be deciphered in this context.

The CO₂ flux during July from observations, constZc, and varZc simulations are 3.09 mol m⁻² 352 yr⁻¹, 1.82 ± 0.4 mol m⁻² yr⁻¹ and 3.10 ± 0.5 mol m⁻² yr⁻¹, respectively. Southwesterly wind-driven 353 354 upwelling over the WAS especially off the Somali coast (Smith and Codispoti, 1980; Schott, 1983; Smith, 1984) and Oman (Bruce, 1974; Smith and Bottero, 1977; Swallow, 1984; Bauer et 355 al., 1991), pulls nutrient-rich subsurface waters closer to the surface while the available turbulent 356 energy due to the strong winds lead to mixed layer entrainment of the nutrients resulting in a 357 strong surface phytoplankton bloom (Krey and Babenerd, 1976; Banse, 1987; Bauer, 1991; 358 Brock et al, 1991). This regional bloom extends over 700 km offshore from the Omani coast due 359 to upward Ekman pumping driven by strong, positive wind-stress curl to the northwest of the low 360 level jet axis and the offshore advection (Bauer et al., 1991; Brock et al., 1991; Brock and 361 362 McClain, 1992a, b; Murtugudde and Busalacchi, 1999, Valsala, 2009) resulting in strong outgassing of CO₂ flux and an enhanced pCO₂ in the WAS (Valsala and Maksyutov, 2013; 363 Sarma et al., 2002). The seasonal mean CO_2 flux during the southwest monsoon period (JJAS) 364 for constZc and varZc simulations are 1.44 ± 0.2 mol m⁻² yr⁻¹ and 2.31 ± 0.4 mol m⁻² yr⁻¹, 365 respectively. The biological parameterization of varZc considerably improves the average CO₂ 366 flux during the monsoon period by 0.86 ± 0.1 mol m⁻² yr⁻¹. The annual mean CO₂ flux from 367 observations, constZc and varZc simulations are 0.94 mol m⁻² yr⁻¹, 0.80 \pm 0.17 mol m⁻² yr⁻¹ and 368 1.07 ± 0.2 mol m⁻² yr⁻¹, respectively. The annual mean CO₂ flux is improved by 0.27 ± 0.05 mol 369 $m^{-2} yr^{-1}$. 370

371 Seasonality in pCO₂ also shows a remarkable improvement during the southwest monsoon 372 period (Figure 6b). The pCO₂ with constZc is considerably lower at a value of $385.22 \pm 3.5 \mu$ atm 373 during June compared to observational values of 392.83 µatm. However, varZc simulation performs better in terms of pCO₂ variability. The peak value of pCO₂ reaches up to 405.42 ± 5.8 374 μ atm. The seasonal mean pCO₂ during the Southwest monsoon period from observations, 375 376 constZc, and varZc simulations are 397.58 μ atm, 389.18 ± 3.6 μ atm and 399.95 ± 5.0 μ atm, respectively. The improvement in pCO₂ by varZc simulation is 10.76 ± 1.3 µatm when compared 377 378 with the constZc simulation. This inherently infers that constZc simulation fails to capture the pCO₂ driven by upwelling during the Southwest monsoon while the varZc simulation is 379 demonstrably better in representing this seasonal increase. The annual mean pCO₂ from 380 381 observations, constZc, and varZc simulations are 394.69 μ atm, 389.62 \pm 3.9 μ atm and 391.19 \pm 4.7 µatm, respectively. However, it is worth mentioning that there are parts of the year where the 382 constZc performs better compared to varZc. For instance, during MAM as well as in November, 383 the constZc simulation yielded a better comparison with the observed pCO_2 whereas varZc 384 simulation yields a reduced magnitude of pCO₂. This may well indicate the biological vs. 385 solubility pump controls on pCO₂ during the intermonsoons. The role of mesoscale variability in 386 387 the ocean dynamics may also play a role (Valsala and Murtugudde, 2015). Nevertheless, during the most important season (JJAS) when the pCO_2 , CO_2 fluxes, and biological production are 388 389 found to be dominant in the Arabian Sea, the varZc produces a better simulation.

The improvements shown by the use of varZc in the simulation of CO₂ flux and pCO₂ can be elicited by further analysis of the model biological production. Figure 7 shows the comparison of model export production and new production with observational export production from satellite-derived NPP for constZc and varZc simulations. The model export production in the constZc simulation is much weaker when compared to varZc simulation. The varZc simulation has improved the model export production. Theoretically, the new and export productions in the model should be in balance with each other (Eppley and Peterson, 1979). The constZc export
production is much weaker than new production and it is not in balance. In contrast, the varZc
simulation yields a close balance among them.

Compared with the observational export production which peaks in August at a value of 399 154.78 g C m⁻² yr⁻¹, the varZc simulated export and new productions peak at a value of 160.44 \pm 400 20.4 g C m⁻² yr⁻¹ and 167.18 \pm 24.0 g C m⁻² yr⁻¹, respectively, but in July. A similar peak can be 401 observed in constZc simulated new production as well, with a value of 178.19 ± 28.0 g C m⁻² yr⁻ 402 ¹. This apparent shift of one month during JJAS in the model export production as well as in the 403 new production is noted as a caveat in the present set up which will need further investigation. 404 Arabian Sea production is not just limited by nutrients but also the dust inputs (Wiggert et al., 405 2006). The dust-induced primary production in the WAS, especially over the Oman coast is 406 noted during August (Liao et al., 2016). The mesoscale variability in the circulation and its 407 408 impact on production and carbon cycle are also a limiting factor in this model as noted above.

409 The seasonal mean export production during the southwest monsoon period from satellitederived estimate is 123.57 g C m⁻² yr⁻¹, whereas for constZc and varZc simulations it is 84.81 \pm 410 16.0 g C m⁻² yr⁻¹ and 147.19 \pm 23.8 g C m⁻² yr⁻¹, respectively. The new biological 411 parameterization strengthens the model export production by 62.38 ± 7.8 g C m⁻² yr⁻¹ for the 412 southwest monsoon period, which is over a 70% increase. This indicates a considerable impact 413 of the biological pump in the model simulated CO₂ flux and pCO₂ over the WAS. For constZc 414 simulation, the computed new production is slightly higher (150.84 \pm 27.9 g C m⁻² yr⁻¹) than that 415 of varZc (133.03 \pm 19.5 g C m⁻² yr⁻¹). The annual mean export production from observations, 416 constZc and varZc simulations are 94.31 g C m⁻² yr⁻¹, 77.41 \pm 15.1 g C m⁻² yr⁻¹ and 122.54 \pm 25.2 417 $g C m^{-2} yr^{-1}$, respectively. 418

To understand how the varZc parameterization strengthens the export production in the model, we have analyzed the phosphate profiles. It appears that the varZc parameterization allows more phosphate concentration (Figure 8a, b) in the production zone and thereby increases the corresponding biological production (Figure 8c, d). The net export production in the model during JJAS is consistent with the satellite data (Figure 7b). However, in the constZc case, the exports are rather 'flat' throughout the season with the imperfect representation of seasonal biological export. The Table 1-4 summarize all the values discussed here.

The impact of varZc in the biological and solubility pumps is computed as per Louanchi et 426 427 al., (1996, see Appendix A). The varZc parameterization has strengthened the biological as well as the solubility pump in the model and thereby modifying the phosphate profiles and achieves a 428 seasonal balance in export versus new production (Figure 9a). During the monsoon period, the 429 varZc simulation increases the strength of the solubility and biological pumps by 10.43 ± 1.3 g C 430 $m^{-2} vr^{-1}$ and 106.52 ± 9 g C m⁻² vr⁻¹, respectively (see Table 5 and 6). Similarly, the annual mean 431 strength of solubility pump and the biological pump is increased by 3.29 ± 0.6 g C m⁻² yr⁻¹ and 432 81.18 ± 9.92 g C m⁻² yr⁻¹, respectively. This supports the fact that the varZc parameterization 433 434 basically modifies the biological and solubility pumps in the model simulation and thereby improves the seasonal cycle of CO_2 flux and pCO_2 . 435

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437 **3.3 Sri Lanka Dome (SLD)**

The seasonal variation in Zc for SLD has a similar pattern as that of WAS. Zc deepens to its maximum during March up to 45.23 ± 0.3 m and reaches its minimum during the following monsoon period at 30.79 ± 1.5 m (Figure 5). The similarities of varZc between WAS and SLD indicates that they both are under similar cycles of solar influx and biological production. The
SLD Chl-a dominates only up to July (Vinayachandran et al., 2004) which explains why
production with varZc increases earlier compared to WAS which occurs during August-October.

The seasonality in CO_2 flux and pCO_2 were compared with Takahashi et al., (2009) 444 observations (Figure 10). varZc results in a slight improvement in CO₂ flux when compared with 445 constZc (Figure 10a). However, both constZc and varZc simulations underestimate the 446 magnitude of CO₂ flux when compared with observations. The seasonal mean CO₂ flux during 447 the monsoon period is $1.79 \text{ mol m}^{-2} \text{ yr}^{-1}$ from observations, which means SLD region is a source 448 of CO₂. But the mean values of constZc and varZc simulations yield flux values of -0.008 ± 0.2 449 mol m⁻² yr⁻¹ and 0.24 \pm 0.2 mol m⁻² yr⁻¹, respectively. The constZc simulation misrepresents the 450 SLD region as a sink of CO₂ during monsoon period which is opposite to that of observations. 451 The varZc simulation corrects this misrepresentation to a source albeit at a smaller magnitude by 452 0.24 ± 0.09 mol m⁻² yr⁻¹ for the monsoon period. Compared to observations, the varZc case 453 underestimates the magnitude of JJAS mean by $1.55 \text{ mol m}^{-2} \text{ yr}^{-1}$. 454

The annual mean CO₂ fluxes for constZc and varZc simulations are -0.02 \pm 0.1 mol m⁻² yr⁻¹ 455 and 0.10 ± 0.2 mol m⁻² yr⁻¹, respectively. The varZc parameterization leads to an improvement of 456 $0.13 \pm 0.1 \text{ mol m}^{-2} \text{ yr}^{-1}$ in the annual mean CO₂ flux when compared with constZc simulation. 457 The observational annual mean of CO_2 flux is 0.80 mol m⁻² yr⁻¹ which is highly underestimated 458 by both simulations. This indicates a regulation of biological production of the region by varZc 459 which makes this region a source of CO₂ during monsoon. The role of the solubility pump may 460 also be underestimated due to the biases in the physical drivers and the lack of mesoscale eddy 461 462 activities in these simulations (Prasanna Kumar et al., 2002; Valsala and Murtugudde, 2015).

463 The seasonality of pCO_2 (Figure 10b) especially in the monsoon period is significantly improved. The mean pCO_2 during the monsoon season from observation over the SLD region is 464 382.44 µatm. The seasonal mean pCO₂ during monsoon period for constZc and varZc 465 simulations are 371.67 ± 6.04 µatm and 379.24 ± 8.9 µatm, respectively. The annual mean pCO₂ 466 from observations, constZc, and varZc simulations are 380.21 μ atm, 370.76 \pm 6.1 μ atm and 467 374.94 ± 9.6 µatm, respectively. varZc simulations improve the JJAS mean pCO₂ by 7.56 ± 2.8 468 μ atm and the annual mean pCO₂ by 4.18 ± 3.5 μ atm, which is reflected in CO₂ flux as well. This 469 is likely due to the impact of new biological parameterization in capturing the episodic upwelling 470 in the SLD region which is further investigated by looking at its biological production. 471

The SLD biological production is highly exaggerated by the model for both constZc and varZc simulations (Figure 11a, b). The seasonal mean biological export for the monsoon period is 51.54 g C m⁻² yr⁻¹ as per satellite-derived estimates. However, the constZc and varZc simulations overestimate it at 167.71 \pm 59.04 g C m⁻² yr⁻¹ and 151.51 \pm 46.4 g C m⁻² yr⁻¹, respectively. This exaggerated export is visible in climatological annual means where for constZc and varZc simulations they are 144.43 \pm 49.8 g C m⁻² yr⁻¹ and 156.08 \pm 43.8 g C m⁻² yr⁻¹, respectively.

For constZc simulation, new production is overestimated from March to October when compared to observations and the second peak is observed in November (Figure 11a). But the overestimate in new production with varZc is observed only during JJAS period by a value of 26.23 g C m⁻² yr⁻¹. For the SLD region, the varZc parameterization overestimates the export production but minimizes the excess new production, especially in the monsoon period by 64.15 \pm 36.4 g C m⁻² yr⁻¹. This indicates that the varZc parameterization is somewhat successful in capturing the upwelling episode during the monsoon over SLD. All values are summarized inTable 1 to 4.

The solubility and biological pumps are modified by the varZc parameterization significantly when compared with the constZc simulation (Figure 9b). Over the monsoon period, the strength of the solubility and biological pumps are improved by 2.81 ± 1.1 g C m⁻² yr⁻¹ and 66.68 ± 9.7 g C m⁻² yr⁻¹, respectively. Similarly, the annual mean strength of solubility and biological pump are increased by 0.99 ± 1.2 g C m⁻² yr⁻¹ and 52.5 ± 5.1 g C m⁻² yr⁻¹ respectively. All values are provided in Table 5 and 6.

493

494 **3.4 Sumatra Coast (SC)**

The seasonal variation in Zc over the SC region lies between 40 m and 46 m (Figure 5). The seasonal maximum occurs during January to March, especially in March with a depth of 45.5 m. During the monsoon period, Zc shoals slightly with a minimum of 41.1 m in July. The variation in Zc is relatively small as compared to the other regions which is consistent with its relatively low production throughout the year.

The seasonality of CO₂ flux and pCO₂ captured by constZc and varZc simulations are shown in Figure 12a, b. The varZc simulations overestimate both CO₂ flux and pCO₂, especially during the monsoon. It is found that the constZc simulation is better compared to varZc simulation. The varZc simulation overestimates the seasonal mean CO₂ flux and pCO₂ by 1.19 mol m⁻² yr⁻¹ and 29.61 μ atm, respectively, compared to observations (Table 1). However, constZc produces a better estimate compared with observations for CO₂ flux and pCO₂. The constZc simulation delivers a better annual mean than varZc (Table 1, 2). The annual mean bias in constZc and 507 varZc simulations for CO₂ flux is -0.0033 mol m^{-2} yr⁻¹ and 0.31 mol m^{-2} yr⁻¹, respectively. 508 Similarly, pCO₂ bias is 1.95 µatm and 9.07 µatm for constZc and varZc simulations.

Biological production simulated by the model along SC explains the overestimation of CO_2 509 flux and pCO_2 (Figure 13). Both constZc and varZc simulations greatly overestimate export 510 production in the model. However, a small enhancement in the new production during JJAS in 511 constZc case is an indicator of upwelling episodes. The seasonal mean new production during 512 the monsoon from constZc and varZc are 63.64 ± 30.9 g C m⁻² yr⁻¹ and 78.11 ± 29.1 g C m⁻² yr⁻¹. 513 respectively (Table 4). The seasonal mean export production during the monsoon from 514 observations is 58.87 g C m⁻² yr⁻¹ (Table 3). The constZc simulation represents a better new 515 production, which is seen as a relatively small exaggeration of CO_2 flux and pCO_2 . The 516 biological response of SC is found to be better with constZc which is in contradiction to a 517 general improvement found with varZc in the other regions examined here. Such discrepancies 518 519 over the SC could be due to the effect of Indonesian Throughflow (Bates et al., 2006) which is not completely resolved in the model due to coarse spatial resolution (also see Valsala et al., 520 2010). 521

The overestimation of export production by varZc simulation is also evident by the increase in strength of the biological and solubility pumps, respectively (Figure 9c). The annual mean and JJAS mean DIC increases in the production zone due to the biological pump is 67.21 ± 1.3 g C m⁻² yr⁻¹ and 83.62 ± 0.5 g C m⁻² yr⁻¹, respectively. Similarly the increase in DIC due to the effect of solubility pump during the JJAS period and annual mean are 10.95 ± 5.2 g C m⁻² yr⁻¹ and 3.87 ± 2.2 g C m⁻² yr⁻¹ respectively (see table 5 and 6).

528

529 **3.5 Seychelles-Chagos Thermocline Ridge (SCTR)**

The SCTR is a unique open-ocean upwelling region with a prominent variability in air-sea interactions (Xie et al., 2002). Wind-driven mixing and upwelling of subsurface nutrient-rich water play a major role in the biological production of this region (Dilmahamod et al., 2016). The seasonal cycle in Zc is shown in Figure 5. The maximum Zc occurs in November at about 44.94 m and the minimum at 33.2 m in July. The shoaling of Zc during the monsoon period shows that the biological parameterization captures the upwelling response over this region.

The seasonality of CO_2 flux and p CO_2 are shown in Figure 14. The Takahashi observations 536 of CO₂ flux shows a peak in June with outgassing of CO₂ during the upwelling episodes. 537 However, both constZc and varZc simulations underestimate this variability. The seasonality of 538 539 CO_2 flux in varZc shows a significant improvement when compared to constZc simulation, but underestimated when compared to observations. The seasonal mean CO₂ flux during the 540 monsoon from observations, constZc and varZc simulations are 0.82 mol m⁻² yr⁻¹, -0.32 ± 0.3 541 mol m⁻² yr⁻¹ and -0.05 ± 0.4 mol m⁻² yr⁻¹, respectively. This represents a reduction in the seasonal 542 mean sink of CO₂ flux in the SCTR region during the monsoon by 0.27 \pm 0.1 mol m⁻² yr⁻¹ 543 bringing it closer to a source region (see Table 1 for details). 544

The improved CO₂ flux is also supported by the seasonal cycle in pCO₂. Based on observations, the seasonal mean of pCO₂ with constZc during JJAS is underestimated by 11.47 μ atm, varZc simulation underestimates it by 6.45 μ atm. So it is evident that varZc simulation capture the upwelling episodes better, marked by a larger pCO₂ during JJAS period. However, the magnitude of pCO₂ is still underestimated compared to observations (Table 2). 550 Figure 15 shows the biological production of constZc and varZc simulations for SCTR. It is clear that both simulations overestimate the export production and underestimate the new 551 production. The JJAS mean export production from observations, constZc and varZc are 51.08 g 552 C m⁻² yr⁻¹, 57.39 \pm 14.2 g C m⁻² yr⁻¹ and 99.23 \pm 29.8 g C m⁻² yr⁻¹, respectively. The varZc 553 simulations exaggerate the model export production by 48.14 g C m⁻² yr⁻¹. The varZc simulation 554 improves the JJAS mean new production by 1.14 ± 2.2 g C m⁻² yr⁻¹ (Table 4). The DIC variations 555 due to the biological pump over the monsoon period and the annual mean also support the 556 exaggerated export production. During the monsoon period, the varZc simulation strengthens the 557 biological and solubility pump by 72.64 \pm 6.2 g C m⁻² yr⁻¹ and -4.56 \pm 1.6 g C m⁻² yr⁻¹, 558 respectively when compared to the constZc simulation (Figure 9d). This is also reflected in the 559 annual mean DIC variations due to the biological and solubility pump effects (see table 5 and 6). 560 561 This slight improvement in the model new production, especially during the monsoon period signals that the varZc better captures the upwelling over SCTR. Considering the annual mean 562 values of model export and new production, constZc simulation is reasonably faithful to 563 observations. 564

The underestimation of CO_2 and pCO_2 , as well as the exaggeration of model export 565 production and a slight, overestimate in model new production may be due to two reasons; (1) 566 SCTR is a strongly coupled region with remote forcing of the mixed layer - thermocline 567 interactions (Zhou et al., 2008) which can affect the seasonality in biological production that the 568 model may not be resolving reasonably, (2) the bias associated with physical drivers, especially 569 wind stress may underestimate the CO₂ flux as well biological production. A similar 570 overestimation of biological production was also reported in a coupled biophysical model 571 (Dilmahamod et al., 2016). 572

Table 1 - 4 shows the entire summary of seasonal and annual mean CO₂ flux, pCO₂ and biological production reported in Section 3.

575

576 **4. Sensitivity Simulations**

577 From the analysis of four major upwelling regions over the Indian Ocean, it is evident that 578 the biological parameterization of varZc better captures upwelling episodes and thus it enhances the model export production. This is most clearly visible over the WAS region. In order to 579 580 quantify how much the varZc parameterization contributes to the seasonality of carbon cycle, two additional sensitivity simulations are carried out; viz. (1) with annual mean offline currents 581 and (2) annual mean offline temperatures with the goal of suppressing the dynamical and 582 thermodynamical effects of seasonal upwelling over WAS (see Section 2 for details). The focus 583 584 on this region is motivated by its prominence as the most productive zone of the Indian Ocean. Moreover, the improvement in the biological processes in the model by the varZc 585 586 parameterization is best captured in this region. The results are discussed below.

587

588 4.1 Impact of varZc parameterization on seasonality of carbon cycle with annual 589 mean currents.

To quantify the impact of varZc parameterization, the model is forced with annual mean currents only over WAS region with unaltered currents in the rest of the ocean. The hypothesis is that the muting of the seasonal variability of Ekman divergence removes the upwelling and the biological pump contribution to production and carbon cycle. The comparison of constZc and

27

varZc then allows us to decipher the impact of varZc in capturing the impacts of upwelling on biological production and the carbon cycle. The smooth blending of currents at the boundary of the WAS domain is achieved by a linear smoothing function as given in Section 2.

The model biological responses (inferred by comparing with the control run) in terms of the 597 CO₂ flux shows a flat pattern over the monsoon period for constZc simulation (Figure 16a). 598 While the varZc simulation forced with the annual mean currents shows an enhanced CO₂ flux 599 indicating the outgassing of CO₂ flux over WAS due to wind-driven upwelling (Figure 16b). 600 This qualitatively shows that the varZc itself has improved the seasonality in the biological 601 processes (export and new production) and captured the upwelling episodes during the monsoon. 602 The varZc parameterization is responsible for an improvement of 0.48 ± 0.04 mol m⁻² yr⁻¹ and 603 0.13 ± 0.02 mol m⁻² yr⁻¹ in the JJAS seasonal and annual mean CO₂ fluxes, respectively. This 604 improves the overall model CO₂ flux in the control run especially in July (Figure 16b). 605

Similar improvements are also noticed in pCO₂ (Figure 17). In the constZc simulation with annual mean currents, the pCO₂ dips down during JJAS monsoon period which indicates the inadequacy of constZc in capturing the upwelling enriched pCO₂ difference (Figure 17a, b). The varZc simulation slightly modifies the pCO₂ in the 'right' direction during JJAS despite the annual mean currents.

The export and new productions in the model explain the modification of CO_2 flux and p CO_2 by varZc parameterization. The biological export production is highly underestimated in the constZc simulation forced with annual mean currents while the varZc simulation captures the seasonal upswing in production (Figure 18). The improved JJAS mean and annual mean export production by 43.51 ± 8.6 g C m⁻² yr⁻¹ and 30.28 ± 13.7 g C m⁻² yr⁻¹, respectively is a clear indication of the positive impacts of a varZc. Similarly, the improvement in JJAS mean and annual mean new production (Figure 19) from varZc simulated with annual mean currents are $17.39 \pm 0.8 \text{ g C m}^{-2} \text{ yr}^{-1}$ and $14.81 \pm 0.1 \text{ g C m}^{-2} \text{ yr}^{-1}$, respectively. In short the varZc biological parameterization improves the export and new productions in the model. This helps the model to capture the upwelling episodes over the study regions. Table 7 summarizes all the results of biological sensitivity runs.

622

4.2 Impact of varZc parameterization on seasonality of carbon cycle with annual mean temperatures.

By imposing the annual mean temperature over WAS region, we are suppressing the cooling 625 effect of temperature due to upwelling and quantifying how much the model seasonality is 626 improved due to varZc parameterization. (see Section 2 for details). The varZc simulations 627 forced with annual mean SST has larger JJAS mean and annual mean CO₂ fluxes by 0.88 ± 0.1 628 mol m⁻² yr⁻¹ and 0.28 \pm 0.07 mol m⁻² yr⁻¹, respectively (Figure 20 and Table 8). For a given 629 630 annual mean SST the solubility pump largely controls the CO₂ emission during JJAS if a varZc is prescribed, likely by the enrichment of DIC (inferred from Figure 8b). Similarly, the 631 improvement in pCO₂ (Figure 21) with varZc simulation is also remarkable. The JJAS mean and 632 annual mean improvements from the implementation of varZc are 11.05 ± 1.9 µatm and $1.91 \pm$ 633 1.4 μ atm, respectively. The detailed quantification of CO₂ and pCO₂ responses for this 634 experimental setup is given in Table 8. The above analysis adds supporting evidence that the 635 varZc simulation strengthens the seasonality of the model compared to the constZc case. This is 636

presumably accomplished by the more accurate Zc and production zone implied with a variableZc.

639

640 5. Summary and Conclusions

641 A spatially and temporally varying Zc parameterization as a function of solar radiation and Chl-a is implemented in the biological pump model of OCMIP-II for a detailed analysis of 642 biological fluxes in the upwelling zones of the Indian Ocean. The varZc parameterization 643 644 improves the seasonality of model CO₂ flux and pCO₂ variability, especially during the monsoon period. A significant improvement is observed in WAS where the monsoon wind-driven 645 upwelling dominates biological production. The magnitude of CO₂ flux matches with 646 observations, especially in July when monsoon winds are at their peak. Monsoon triggers 647 648 upwelling in SLD as well which acts as a source of CO₂ to the atmosphere. The seasonal and annual mean are underestimated with constZc and the SLD is reduced to a sink of CO₂ flux. The 649 varZc simulation modifies the seasonal and annual means of the CO₂ flux of SLD and depict it as 650 651 a source of CO₂ especially during the monsoon, but the magnitude is still underestimated compared to Takahashi et al. (2009) observations. The SCTR variability is underestimated by 652 both constZc and varZc simulations, portraying it as a CO_2 sink region whereas observations 653 over the monsoon period indicate that the thermocline ridge driven by the open ocean wind-654 stress curl is, in fact, an oceanic source of CO₂. However, the varZc simulation reduces the 655 656 magnitude of the sink in this region bringing it relatively close to observations.

657 VarZc biological parameterization strengthens the export and new productions in the model,
658 which allows it to represent a better seasonal cycle of CO₂ flux and pCO₂ over the study regions.

The WAS export production is remarkably improved by 62.37 ± 7.8 g C m⁻² yr⁻¹ compared to 659 660 constZc. This supports our conclusion that the varZc parameterization increases the strength of biological export in the model. Over the SLD, the JJAS seasonal mean export and new 661 production are underestimated in varZc compared to constZc simulations, but the annual mean 662 export production is improved. Export production at SC and SCTR are highly exaggerated and 663 there is hardly any improvement in new production with a variable Zc especially over the 664 monsoon period. The inability of varZc parameterization to improve the seasonality of SC and 665 SCTR may be due to the interannual variability of biological production associated with the 666 667 Indonesian throughflow and remote forcing of the mixed layer-thermocline interactions and the effect of biases in the wind stress data used as a physical driver in the model. 668

Sensitivity experiments carried out by prescribing annual mean currents or temperatures over 669 selected subdomains reveal that the varZc retains the seasonality of carbon fluxes, pCO₂, and 670 671 export and new productions closer to observations. This strongly supports our contention that varZc parameterization improves export and new productions and it is also efficient in capturing 672 upwelling episodes of the study regions. This points out the significant role of having a close 673 balance in seasonal biological export and new production in models to capture the seasonality in 674 the carbon cycle. This also confirms the role of biological and solubility pumps in producing the 675 seasonality of carbon cycle in the upwelling zones. 676

However, the underestimation of the seasonality of CO_2 flux over the SLD and overestimation over the SC as well as the SCTR are a cautionary flag for the study. This uncertainty poses an important scientific question as to whether the model biology over the SC and SCTR region is not resolving the seasonality in CO_2 flux and p CO_2 properly or whether the seasonality in the Zc is not able to fully capture the biological processes.

| 682 | To address these questions we have used an inverse modeling approach (Bayesian inversion) |
|-----|--|
| 683 | in order to optimize the spatially and temporally varying Zc using surface pCO_2 as the |
| 684 | observational constraint and computed the optimized biological production. The results will be |
| 685 | reported elsewhere. |
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700 Appendix – A

701 The time evolution equations of the model variables are given by

702
$$\frac{d[PO_4]}{dt} = L([PO_4]) + J_b PO_4$$
(A1)

703
$$\frac{d[DOP]}{dt} = L([DOP]) + J_b DOP$$
(A2)

704
$$\frac{dDIC}{dt} = L([DIC]) + J_b DIC + J_g DIC + J_v DIC$$
(A3)

705
$$\frac{d[ALK]}{dt} = L([ALK]) + J_bALK + J_vALK$$
(A4)

Where L is the 3D transport operator, which represents the effects of advection, diffusion, and convection. [] or square brackets indicate the concentrations in mol m⁻³. J_bPO_4 , J_bDOP , J_bDIC , J_bALK is the biological source/sink terms and J_vDIC , J_vALK are the virtual source-sink terms representing the changes in surface DIC and ALK, respectively, due to evaporation and precipitation. J_vDIC is the source-sink term due to air-sea exchange of CO₂.

711 The following equations represent for the biological processes in the model

For
$$Z < Zc$$
,

713
$$J_{prod} = \frac{1}{\tau} ([PO_4] - [PO_4^*]), \quad [PO_4] > [PO_4^*]$$
 (A5)

714
$$J_{DOP} = \sigma J_{prod} - \kappa [DOP]$$
 (A6)

715
$$J_{PO4} = -J_{prod} + \kappa[DOP]$$
(A7)

716
$$J_{ca} = Rr_{C:P}(1-\sigma)J_{prod}$$
(A8)

717
$$J_{DIC} = r_{C:P} J_{PO4} + J_{ca}$$
 (A9)

718
$$J_{ALK} = -r_{N:P}J_{PO4} + 2J_{ca}$$
 (A10)

For Z > Zc,

720
$$J_{prod} = 0,$$
 $[PO_4] \le [PO_4^*]$ (A11)

$$721 \quad J_{DOP} = -\kappa[DOP] \tag{A12}$$

722
$$J_{PO4} = -\frac{\partial F}{\partial Z} + \kappa [DOP]$$
(A13)

723
$$F_c = (1 - \sigma) \int_0^{Z_c} J_{prod} dZ$$
 (A14)

724
$$F(Z) = F_c(\frac{Z}{Zc})^{-a}$$
 (A15)

$$725 \quad J_{ca} = -\frac{\partial F_{Ca}}{\partial Z} \tag{A16}$$

726
$$F_{Ca} = Rr_{C:P}F_C e^{-(Z-Zc)/d}$$
 (A17)

727 Where Z is the depth and Zc is the compensation depth in the model. J_{DOP} and J_{PO4} are the biogeochemical sources and sinks and J_{prod} , J_{Ca} represents the biogeochemical flows with 728 respect to production and calcification. Within Zc, the production of organic phosphorous in the 729 model J_{prod} is calculated using equation A5. [PO₄] is the model phosphate concentration and 730 $[PO_4^*]$ is observational phosphate. τ is the restoration timescale assumed to be 30 days. Whenever 731 732 the model phosphate exceeds the observational phosphate, it allows production, below which the 733 production is zero. The observational phosphate data were taken from the World Ocean Atlas (WOA; Garcia et al., 2014). It is assumed that a fixed fraction (σJ_{prod}) of phosphate uptake is 734

converted into Dissolved Organic Phosphorus (DOP) which is a source for J_{DOP} (equation A6). 735 The phosphate not converted to DOP results in an instantaneous downward flux of particulate 736 organic phosphorus at Zc (equation A14). The decrease of flux with depth due to 737 remineralization is shown by a power law relationship as in equation A15. The values of the 738 constants a, κ , σ are 0.9, 0.2/year to 0.7/year, 0.67, respectively. The rate of production is used to 739 740 explain the formation of calcium carbonate cycle in surface waters (equation A8) and its export is given by equation A16, where R is the rain ratio, a constant molar ratio of exported particulate 741 742 organic carbon to the exported calcium carbonate flux at Zc. The exponential decrease of calcium carbonate flux with scale depth d is given by equation A17. The biological source or 743 sink of dissolved inorganic carbon (DIC) and alkalinity (ALK) is explained through equations 744 A9 and A10, respectively. Where the values of rain ratio (R) is taken as 0.07 and the Redfield 745 ratio, $r_{C:P} = 117$, and $r_{N:P} = 16$ and scale depth d is chosen as 3500m. 746

747

748 Biological and Solubility Pump calculations

The biological effect on DIC is calculated from Louanchi et al., (1996). The tendency of DICdue to biomass production and calcite formation in the production zone is expressed as below.

751
$$\left(\frac{\partial DIC}{\partial t}\right)_b = \left(\frac{\partial PO_4}{\partial t}\right)_b \times R_{C:P} - J_{Ca}$$
 (A18)

752 The total tendency of DIC in the production zone is:

753
$$\left(\frac{\partial DIC}{\partial t}\right)_{total} = \left(\frac{\partial DIC}{\partial t}\right)_b + \int_x \int_y F dx \, dy \tag{A19}$$

where $\left(\frac{\partial DIC}{\partial t}\right)_{b}$ is the evolution of DIC due to the impact of biology (i.e., biological pump). The first term in the R.H.S of Equation A18 is the rate of change of phosphate resulting from photosynthesis and respiration in the model (i.e., J_{po4} in this case) multiplied by the carbon to phosphorous Redfield ratio ($R_{C:P} = 117:1$) and J_{Ca} represents the calcite formation in the model (see Equation A8 & A16). The solubility pump is calculated as the surface integral of the flux F (Louanchi et al., 1996).

760

761 Appendix B

762 In order to compare our model production of organic phosphorous to the curve of Ryther et al., 763 (1956) we have merely scaled our total production to "relative photosynthesis", which is, according to Ryther et al., (1956) is an index between 0 and 1 indicating the strength of 764 765 production estimated as P₁/P_{max}. Here P₁ is the photosynthesis at each intensity (of light) of 766 different species and P_{max} is the maximum photosynthesis observed in the same control 767 experiment. The curve between relative photosynthesis and light intensity shows the relation between photosynthetic activity and light in marine phytoplankton. Since our method relates 768 biological production to a function of light (limitation) by Chl-a attenuation, it is the best curve 769 770 to cross-compare our results. In this case we scaled our total biological production within Zc into 771 relative values between 0-1 by P_1/P_{max} . in which P_1 is taken as the individual grid cell biological component of organic phosphorus production and P_{max} is the maximum production available in 772 the domain at any given instant. All the grid points are quite similar to the curve of Ryther et al., 773 774 (1956) as shown in Figure 2.

775

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Table: 1 WAS = Western Arabian Sea, SLD = Sri Lanka Dome, SC = Sumatra Coast, SCTR = Seychelles-Chagos Thermocline Ridge. JJAS mean and the climatological annual mean of CO₂ flux from Takahashi observations, constZc, and varZc simulations. Units are mol $m^{-2} yr^{-1}$.

| | | | CO ₂ flux (me | ol $\mathbf{m}^{-2} \mathbf{y} \mathbf{r}^{-1}$) | | | | |
|---------|------|---------------------------|--------------------------|---|--------------------------|--------------------------|--|--|
| Regions | | JJAS Mea | n | Annual Mean | | | | |
| | OBS | constZc | varZc | OBS | constZc | varZc | | |
| WAS | 1.99 | 1.44 ± 0.2 | 2.31 ± 0.4 | 0.94 | 0.80 ± 0.1 | 1.07 ± 0.2 | | |
| SLD | 1.79 | $\textbf{-0.008} \pm 0.2$ | 0.24 ± 0.09 | 0.8 | $\textbf{-0.02}\pm0.1$ | 0.10 ± 0.2 | | |
| SC | 0.31 | 0.60 ± 0.5 | 1.51 ± 1.01 | 0.21 | 0.21 ± 0.3 | 0.53 ± 0.5 | | |
| SCTR | 0.82 | $\textbf{-0.32}\pm0.3$ | -0.05 ± 0.4 | 0.55 | $\textbf{-0.02} \pm 0.1$ | $\textbf{-0.07} \pm 0.2$ | | |

Table: 2 Same as Table 1, but for pCO₂. Units are µatm.

| | | | pCO ₂ (µa | tm) | | |
|---------|--------|-------------------|----------------------|--------|------------------|-------------------|
| Regions | | JJAS Mea | n | | an | |
| | OBS | constZc | varZc | OBS | constZc | varZc |
| WAS | 397.58 | 389.18 ± 3.7 | 399.95 ± 5.01 | 394.69 | 389.62 ± 3.9 | 391.19 ± 4.7 |
| SLD | 382.44 | 371.67 ± 6.04 | 379.24 ± 8.9 | 380.21 | 370.76 ± 6.1 | 374.94 ± 9.6 |
| SC | 372.52 | 382.36 ± 12.7 | 402.14 ± 21.8 | 372.69 | 374.65 ± 9.3 | 381.76 ± 13.6 |
| SCTR | 377.18 | 365.71 ± 5.08 | 370.72 ± 7.4 | 379.89 | 372.69 ± 4.7 | 369.00 ± 5.4 |

| | | | Export Pro | duction (g | $C m^{-2} yr^{-1}$ | | | |
|---------|--------|--------------------|------------------|-------------|--------------------|-------------------|--|--|
| Regions | | JJAS Mean | l | Annual Mean | | | | |
| | OBS | constZc | varZc | OBS | constZc | varZc | | |
| WAS | 123.57 | 84.81 ± 16.04 | 147.19 ± 23.8 | 94.31 | 77.41 ± 15.1 | 122.54 ± 25.2 | | |
| SLD | 51.54 | 167.71 ± 59.04 | 151.51 ± 46.4 | 43.25 | 144.43 ± 49.8 | 156.08 ± 43.8 | | |
| SC | 58.87 | 260.11 ± 104.7 | 310.03 ± 99.5 | 54.53 | 172.52 ± 72.4 | 215.52 ± 70.8 | | |
| SCTR | 51.08 | 57.39 ± 14.2 | 99.23 ± 21.8 | 40.45 | 55.15 ± 17.9 | 80.35 ± 26.04 | | |

Table: 3 JJAS mean and the climatological annual mean of export production from satellite-derived Net Primary Production data, constZc, and varZc simulations. Units are g C m^{-2} yr⁻¹.

Table: 4 Model derived values for New production. Units are g C m⁻² yr⁻¹.

| | New Production (g C m ⁻² yr ⁻¹) | | | | | | | | |
|---------|--|------------------|------------------|-------------|-------------------|------------------|--|--|--|
| Regions | | JJAS Mea | n | Annual Mean | | | | | |
| | OBS | constZc | varZc | OBS | constZc | varZc | | | |
| WAS | | 150.84 ± 27.9 | 133.03 ± 19.5 | | 108.43 ± 23.4 | 81.47 ± 15.7 | | | |
| SLD | | 141.93 ± 64.1 | 77.78 ± 27.6 | | 111.05 ± 71.1 | 50.37 ± 26.3 | | | |
| SC | | 63.64 ± 30.9 | 78.11 ± 29.1 | | 56.69 ± 43.3 | 54.58 ± 23.3 | | | |
| SCTR | | 12.17 ± 16.3 | 13.32 ± 18.6 | | 13.74 ± 15.5 | 12.94 ± 13 | | | |

| | Biological Pump (g C m ⁻² yr ⁻¹) | | | | | | | |
|---------|---|-------------------|-----------------|-------------------|--|--|--|--|
| Regions | con | stZc | varZc | | | | | |
| | JJAS Mean | Annual Mean | JJAS Mean | Annual Mean | | | | |
| WAS | 45.18 ± 14.8 | 45.49 ± 14.38 | 151.7 ± 23.8 | 126.67 ± 24.3 | | | | |
| SLD | 89.39 ± 58.1 | 108.65 ± 48.6 | 156.07 ± 48.4 | 161.15 ± 43.5 | | | | |
| SC | 235.54 ± 95.4 | 155.21 ± 67.4 | 319.16 ± 94.9 | 222.92 ± 68.7 | | | | |
| SCTR | 30.49 ± 13.4 | 26.81 ± 16.8 | 103.13 ± 19.6 | 83.98 ± 23.6 | | | | |

Table 5: Biological pump impact over DIC in the model due to constZc and varZc simulations for JJAS and annual mean.

Table 6: Same as Table 5, But for Solubility pump.

| | | Solubility Pun | $np (g C m^{-2} yr^{-1})$ | _ | | |
|---------|-----------------|-----------------|---------------------------|-----------------|--|--|
| Regions | COI | nstZc | varZc | | | |
| | JJAS Mean | Annual Mean | JJAS Mean | Annual Mean | | |
| WAS | 17.29 ± 3.5 | 9.63 ± 2.1 | 27.72 ± 4.8 | 12.92 ± 2.7 | | |
| SLD | -0.09 ± 2.4 | -0.32 ± 2.3 | 2.9 ± 3.5 | 1.31 ± 3.5 | | |
| SC | 7.22 ± 6.9 | 2.56 ± 3.8 | 18.17 ± 12.1 | 6.43 ± 6.0 | | |
| SCTR | -3.95 ± 3.7 | -0.35 ± 2.3 | -0.61 ± 5.3 | -0.86 ± 2.8 | | |

| | WAS region forced with Annual Mean Currents | | | | | | | |
|---|---|------------------|---------------|-----------------|------------------|--------------------------|--|--|
| | | JJAS mean | | Clima | atological annua | al mean | | |
| | constZc | varZc | Improvement | constZc | varZc | Improvement | | |
| CO_2 flux (mol m ⁻² yr ⁻¹) | 0.80 ± 0.2 | 1.29 ± 0.2 | 0.48 ± 0.04 | 0.65 ± 0.1 | 0.79 ± 0.1 | 0.13 ± 0.02 | | |
| pCO ₂ (µatm) | 381.81 ± 3.4 | 387.24 ± 3.9 | 5.43 ± 0.5 | 388.68 ± 3.4 | 388.40 ± 3.6 | $\textbf{-0.28} \pm 0.1$ | | |
| Export production (g C m ⁻² yr ⁻¹) | 60.71 ± 4.7 | 104.22 ± 13.4 | 43.51 ± 8.6 | 74.30 ± 4.5 | 104.58 ± 18.3 | 30.28 ± 13.7 | | |
| New Production (g C m ⁻² yr ⁻¹) | 34.76 ± 2.3 | 52.16 ± 1.51 | 17.39 ± 0.8 | 29.91 ± 1.7 | 44.72 ± 1.6 | 14.81 ± 0.1 | | |

 Table 7: JJAS mean and climatological annual mean response from the model forced with annual mean currents.

Table 8 Same as Table 7 but from annual mean temperature simulation.

| | | WAS region forced with Annual mean temperature | | | | | | | | |
|--|-----------------|--|-----------------|------------------|-------------------|----------------|--|--|--|--|
| | | JJAS mean | | Clin | natological annua | l mean | | | | |
| | constZc | varZc | Improvement | constZc | varZc | Improvement | | | | |
| CO ₂ flux (mol m ⁻² yr ⁻¹) | 1.85 ± 0.2 | 2.74 ± 0.4 | 0.88 ± 0.1 | 0.81 ± 0.1 | 1.10 ± 0.2 | 0.28 ± 0.07 | | | | |
| pCO ₂ (µatm) | 393.20 ± 3.01 | 404.26 ± 4.9 | 11.05 ± 1.9 | 384.61 ± 3.3 | 386.52 ± 4.8 | 1.91 ± 1.4 | | | | |

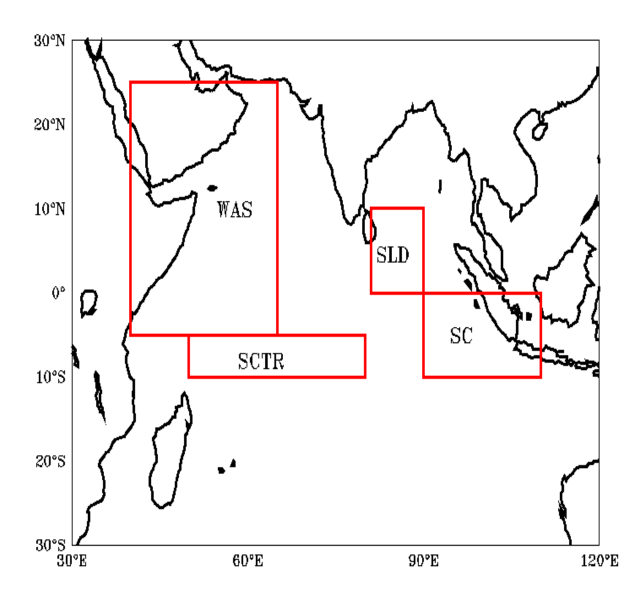


Figure 1: Red boxes shows the study regions (1) WAS (Western Arabian Sea, 40°E:65°E, 5°S:25°N) (2) SLD (Sri Lanka Dome, 81°E:90°E, 0°:10°N) (3) SCTR (Seychelles-Chagos Thermocline Ridge, 50°E:80°E, 5°S:10°S) and (4) SC (Sumatra Coast, 90°E:110°E, 0°:10°S).

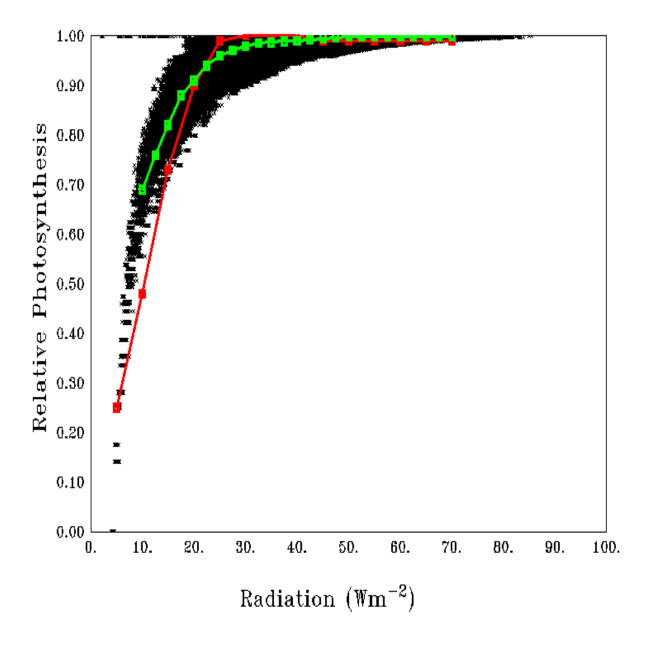


Figure 2: Scatter of average relative photosynthesis versus different light intensities in the model (black dots) and its mean (green curve). The red curve shows the theoretical P - I curve from Parsons et al., (1984).

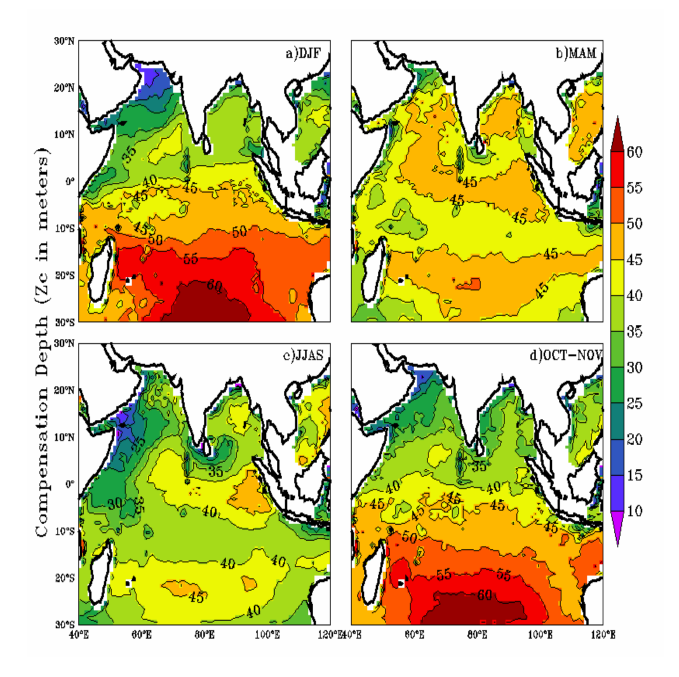


Figure 3: Seasonal-mean maps of varying compensation depth (varZc), (a) December to February (DJF), (b) March to May (MAM), (c) June to September (JJAS), (d) October to November (OCT-NOV). Units are meters.

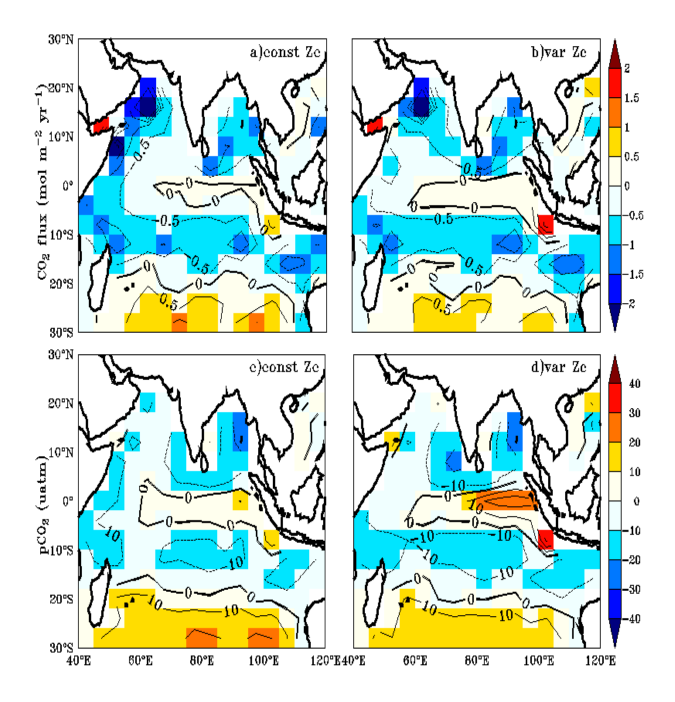


Figure 4: Annual mean biases in the model evaluated against Takahashi et al. (2009) observations for CO₂ flux (a, b) and pCO₂ (c, d) with constant Zc (constZc) and varying Zc (varZc). Units of CO₂ flux and pCO₂ are mol m^{-2} yr⁻¹ and µatm, respectively.

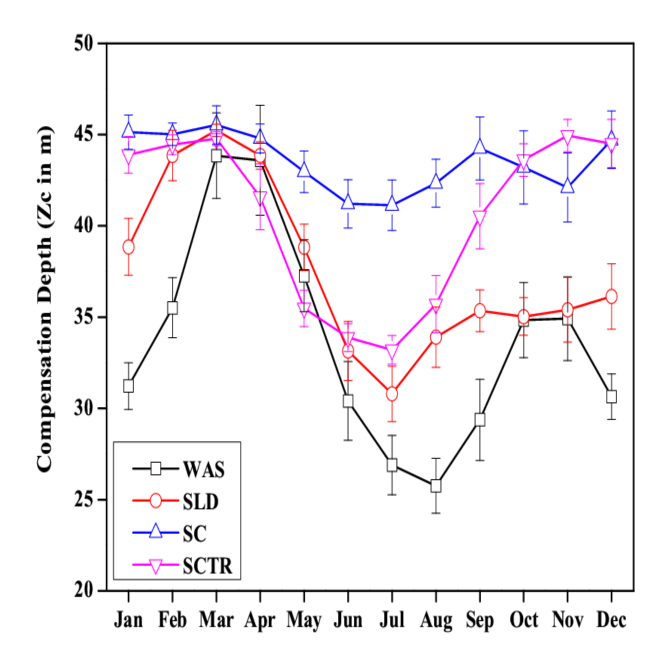


Figure 5: Seasonal variations in varZc over the study regions shown as climatology computed over 1990-2010. Error bar shows standard deviations of individual months over these years. Units are meters.

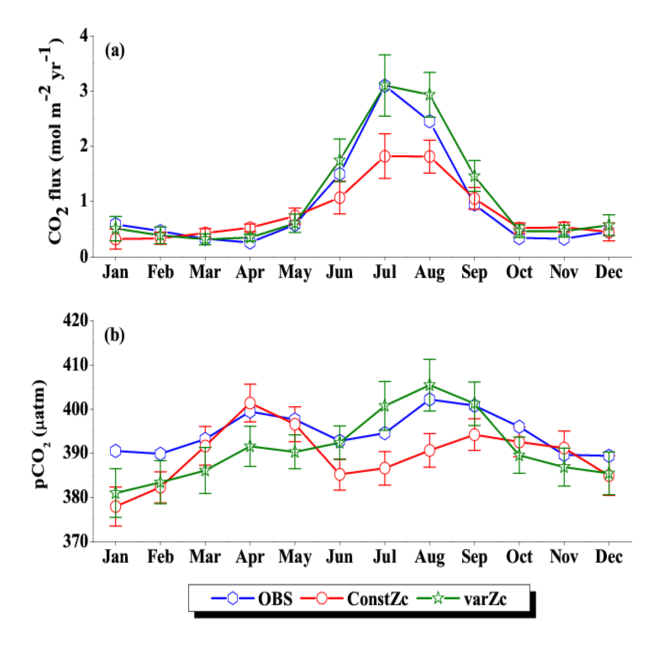


Figure 6: Comparison of model (a) CO_2 flux and (b) p CO_2 simulated with constZc and varZc with that of Takahashi et al. (2009) observations (OBS) over WAS as climatology computed over 1990-2010. Error bar shows standard deviations of individual months over these years. Units of CO_2 flux and p CO_2 are mol m⁻² yr⁻¹ and µatm, respectively. Legend is common for both graphs.

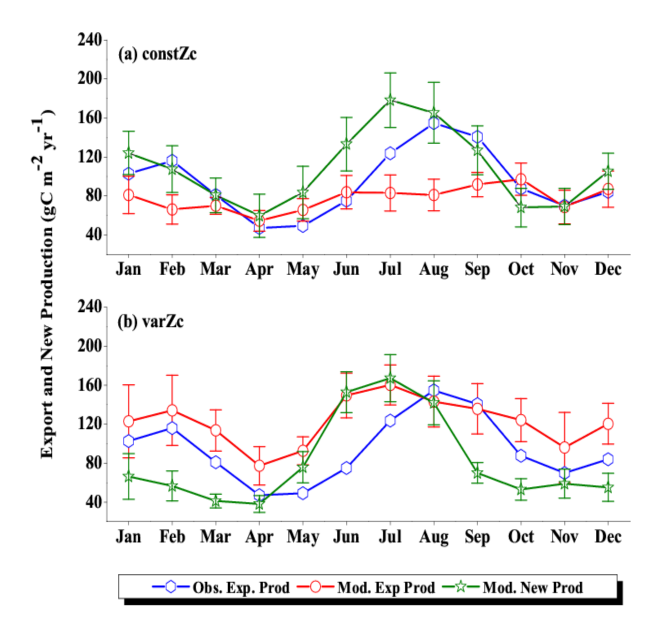


Figure 7: Comparison of model export production (Mod. Exp. Prod) and new production (Mod. New Prod) with satellite-derived export production (Obs. Exp. Prod) for (a) ConstZc and (b) varZc simulations for WAS. Units are g C m^{-2} yr⁻¹. Legends are common for both graphs.

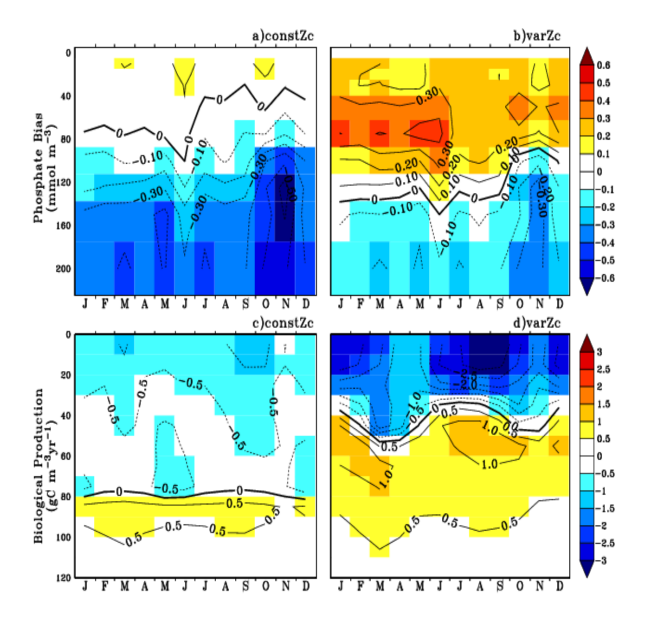


Figure 8: Annual mean bias of model phosphate when compared with climatological observational data (a) for constZc and (b) for varZc simulations. Corresponding annual mean biological source/sink profiles (c, d) in the model for WAS. Unit of phosphate is mmol m^{-3} and biological source/sink is g C m^{-3} yr⁻¹.

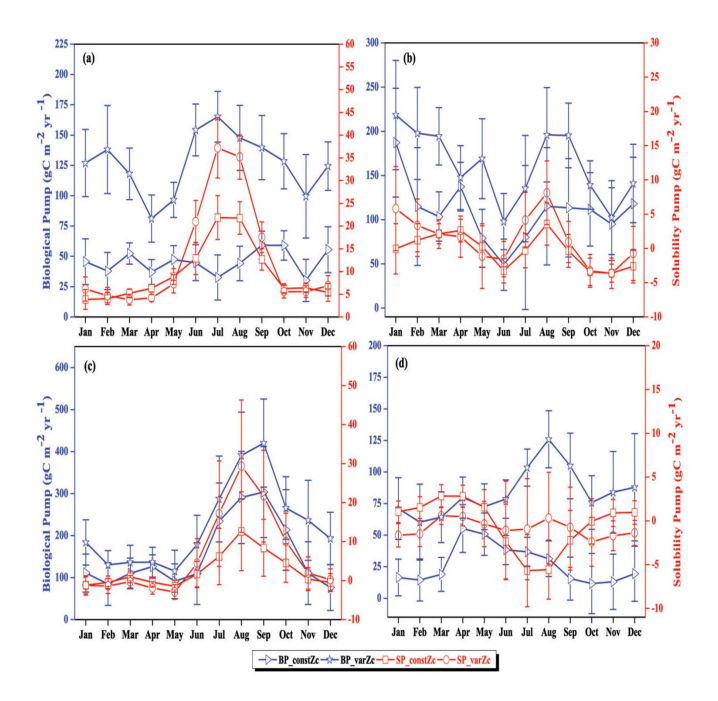


Figure 9: The strength of the biological pump (BP, black lines) and solubility pump (SP, red lines) from constZc and varZc simulations for (a) WAS (b) SLD (c) SC and (d) SCTR. The left axis shows the biological pump and the right axis shows the solubility pump. Error bar shows standard deviations of individual months over the years 1990 - 2010. Units are g C m⁻² yr⁻¹.

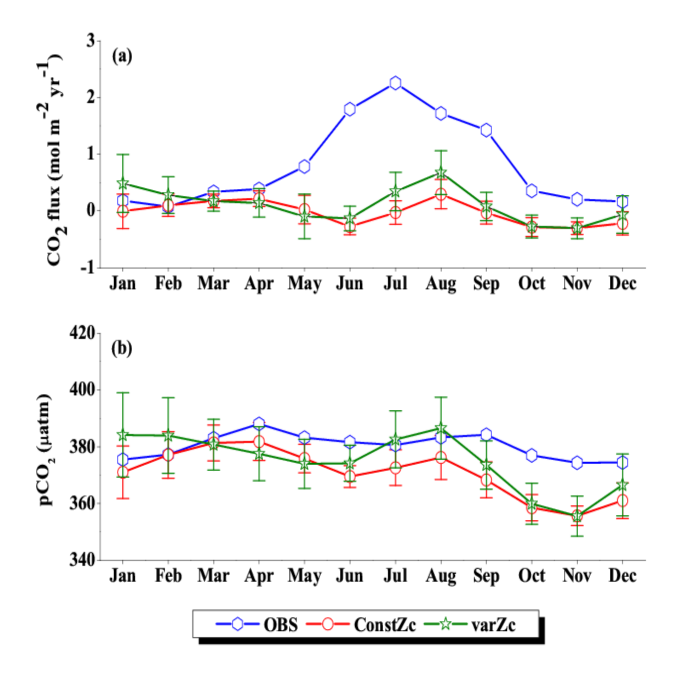


Figure 10: Same as Figure (6), but for SLD.

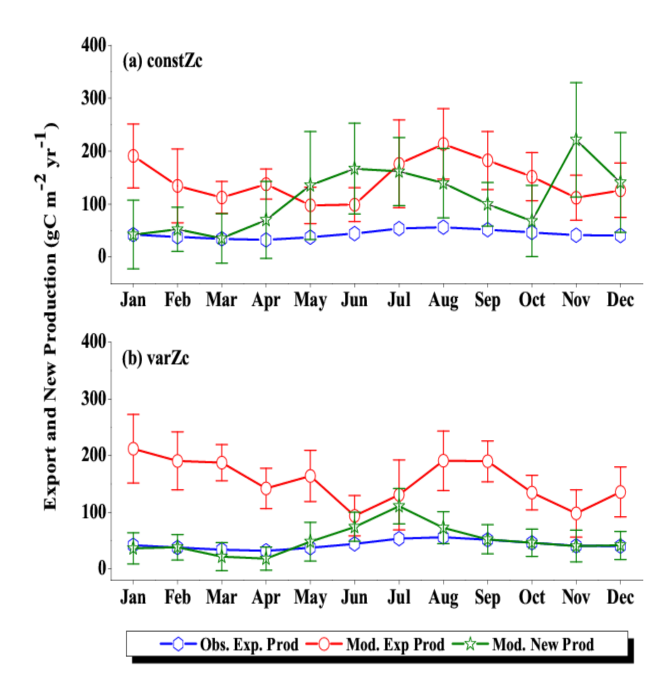


Figure 11: Same as Figure (7), but for SLD.

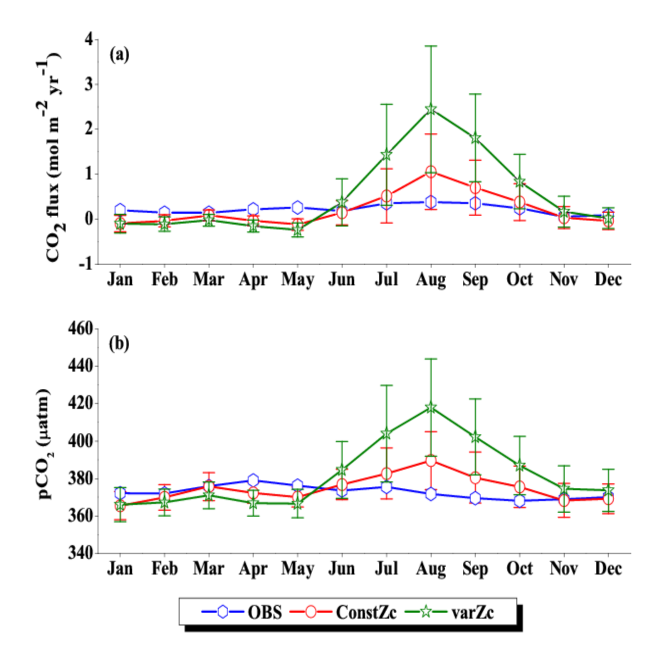


Figure 12: Same as Figure (6), but for SC.

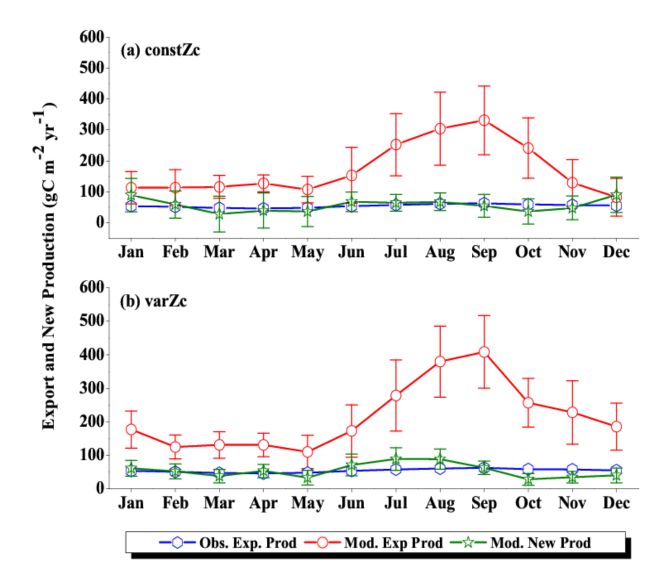


Figure 13: Same as Figure (7), but for SC.

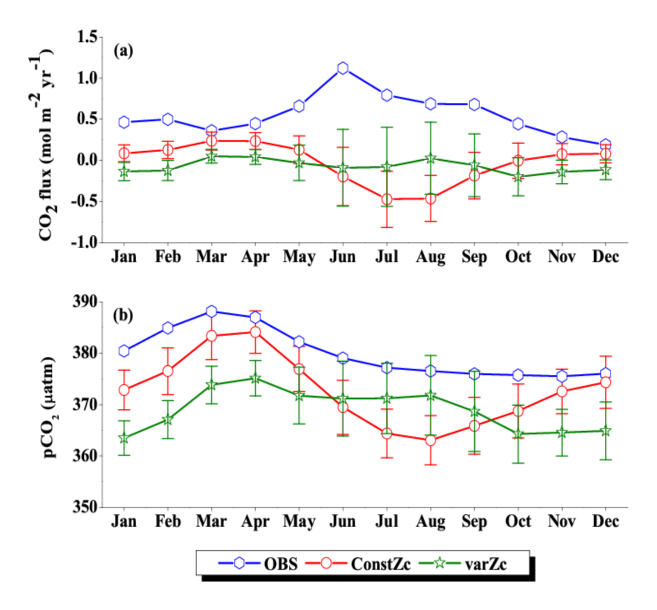


Figure 14: Same as Figure (6), but for SCTR.

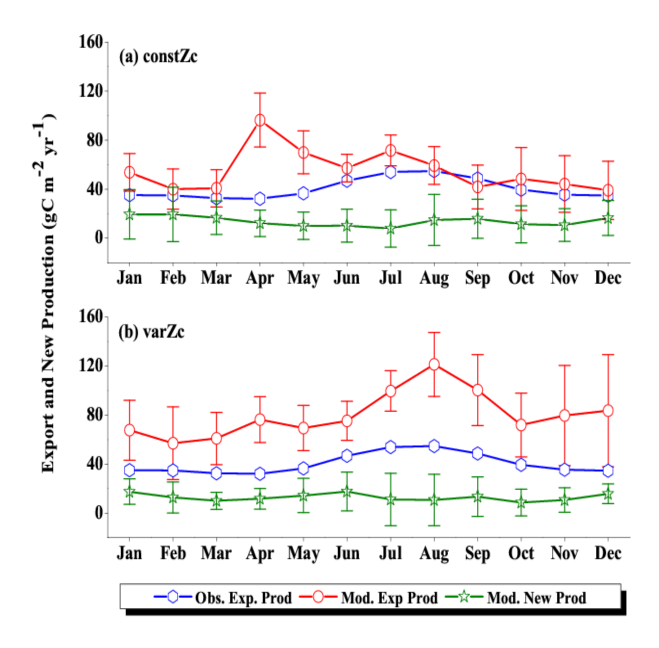


Figure 15: Same as Figure (7), but for SCTR.

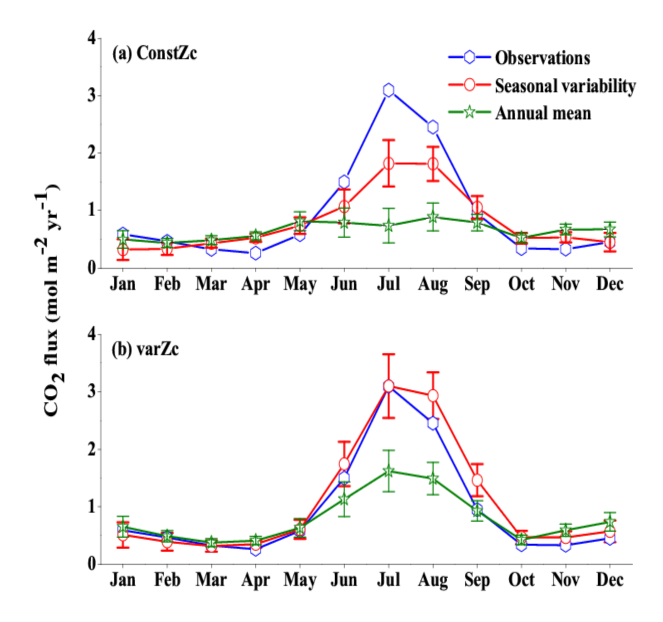


Figure 16: Response of CO_2 flux from the model forced with annual mean currents over the WAS as climatology computed over 1990-2010. Error bar shows standard deviations of individual months over these years. (a) constZc and (b) varZc. Units are mol m⁻² yr⁻¹. Legends are same for both graphs.

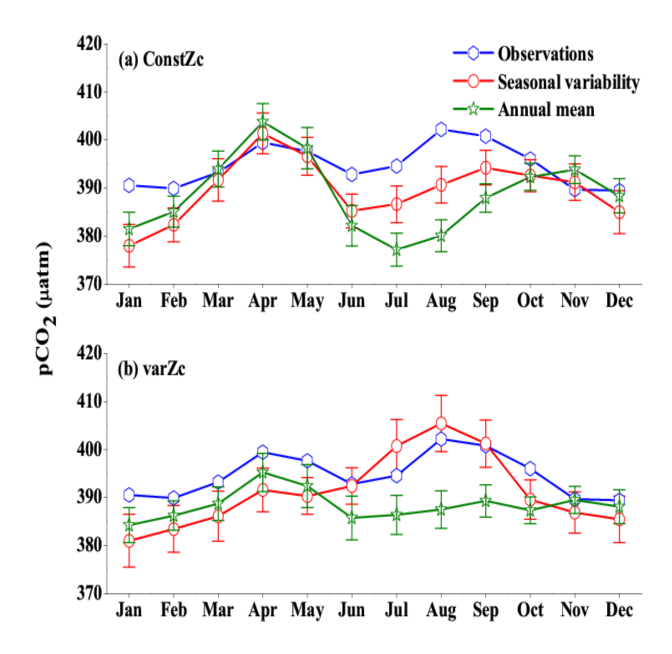


Figure 17: Same as Figure (16), but for pCO₂. Units are µatm.

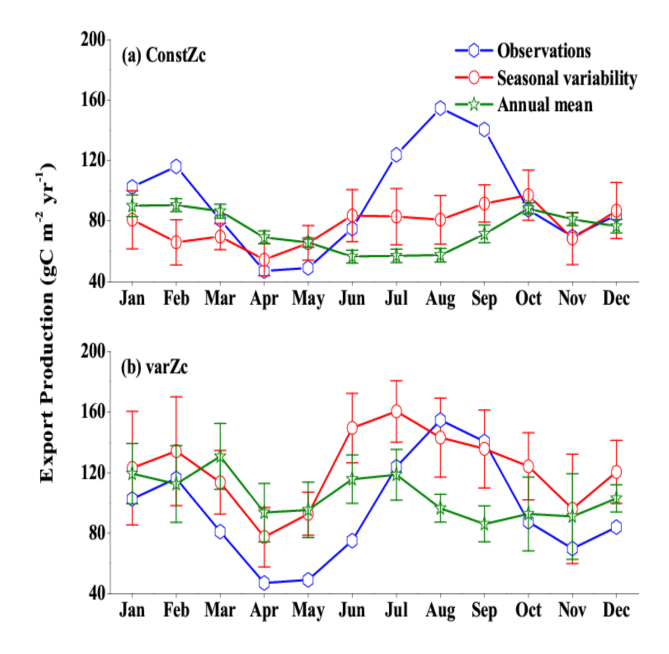


Figure 18: Response in the export production of the model forced with annual mean currents in the WAS as climatology computed over 1990-2010. Error bar shows standard deviations of individual months over these years. (a) constZc (b) varZc. Units are g C m^{-2} yr⁻¹. Legends are same for both graphs.

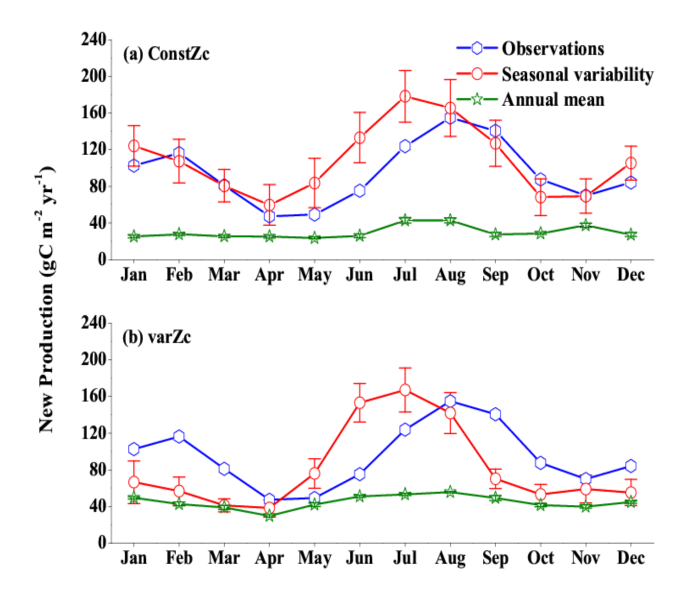


Figure 19: Same as Figure (18), but for New Production. Units are g C m⁻² yr⁻¹.

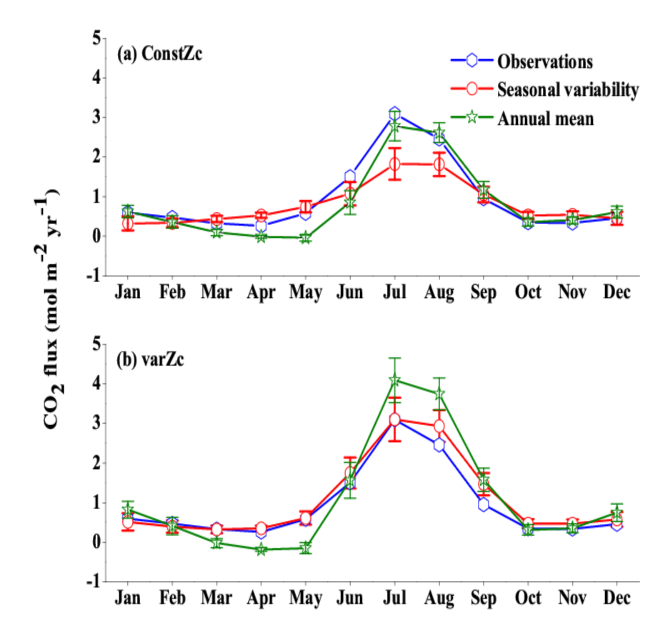


Figure 20: Response of CO_2 flux from the model forced with annual mean SST over the WAS as climatology computed over 1990-2010. Error bar shows standard deviations of individual months over these years. (a) constZc and (b) varZc. Units are mol m⁻² yr⁻¹. Legends are same for both graphs.

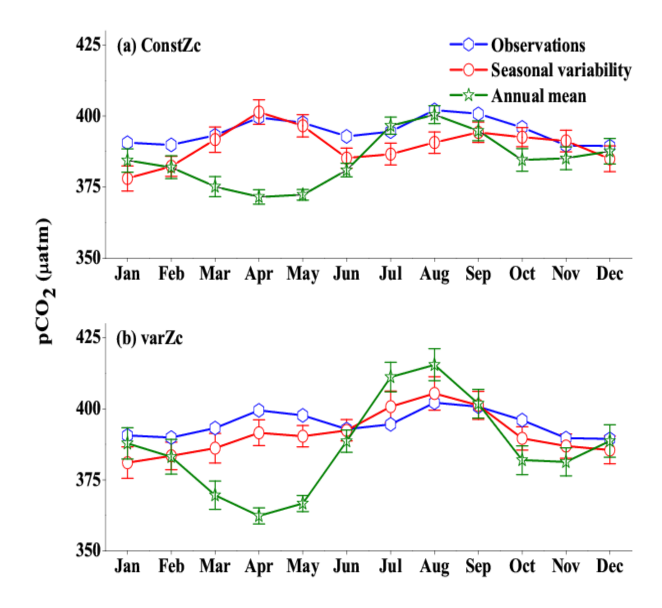


Figure 21: Same as Figure (20), But for pCO₂. Units are µatm.