



36 hydrothermal sources, which may well be of importance in a region known for its strong precipitation rates and volcanic
37 activity.

38

39 1. Introduction

40

41 Nitrogen is known to be the most common limiting nutrient for phytoplankton growth in the modern world ocean
42 (Moore et al., 2013), especially in the Low Nutrient, Low Chlorophyll (LNLC) ecosystems (Arrigo, 2005; Gruber,
43 2005). Characterizing the processes governing nitrogen sources and sinks to and from the ocean is therefore central to
44 understanding oceanic production, organic matter export and food web structure. Atmospheric dinitrogen (N_2) dissolved
45 in seawater is by far the dominant form of N present in the ocean, i.e. the $N_2:NO_3^-$ ratio typically exceeds 100 in surface
46 waters. However, most phytoplankton species cannot assimilate N_2 , and grow using reactive forms of nitrogen such as
47 nitrate, ammonium and dissolved organic compounds. Some planktonic prokaryotic microorganisms, called
48 “diazotrophs” use an enzyme, the nitrogenase, to fix N_2 and convert it into ammonia (NH_3) and ultimately ammonium
49 (NH_4^+). At the global scale, they provide the major external source of reactive nitrogen to the ocean (Gruber, 2008), and
50 support up to 50% of new production in tropical and subtropical (LNLC) regions (Bonnet et al., 2009; Capone, 1997;
51 Deutsch et al., 2007; Karl et al., 1997; Moutin et al., 2008; Raimbault and Garcia, 2008). These organisms are
52 physiologically and taxonomically diverse including cyanobacteria, bacteria, archaea (Zehr and Bombar, 2015; Delmont
53 et al., 2017)(Delmont et al., 2017; Zehr and Bombar, 2015).

54 Autotrophic diazotrophs have been far more intensively studied than heterotrophic diazotrophs, whose contribution to
55 global N_2 fixation remains unclear (Turk-Kubo et al., 2014; Bombar et al., 2016; Moisaner et al., 2017)(Bombar et al.,
56 2016; Moisaner et al., 2017; Turk-Kubo et al., 2014). Autotrophic diazotrophs have been characterized both in the field
57 and through laboratory experiments and their physiology is consequently better (Bergman et al., 2013; Küpper et al.,
58 2008; Mulholland et al., 2001; Mulholland and Capone, 2000; Ohki et al., 1992; Ramamurthy and Krishnamurthy,
59 1967; Rubin et al., 2011). Cyanobacterial (autotrophic) diazotrophs are composed of 3 main groups: 1) the filamentous
60 diazotrophs including the colonial, non-heterocyst-forming *Trichodesmium*, 2) the heterocyst-forming symbionts
61 associated with diatoms (DDAs), and 3) the unicellular cyanobacterial diazotrophs (UCYN, phylogenetically divided
62 into three groups: UCYN-A, B, and C). It has been established that autotrophic diazotrophs growth rates are typically
63 one order of magnitude lower than those of non-diazotrophs (Breitbarth et al., 2008; Falcón et al., 2005; Goebel et al.,
64 2008; LaRoche and Breitbarth, 2005). This can be related to the high energetic demand (Postgate, 1982) required to
65 convert N_2 to NH_3 as compared to that necessary to assimilate nitrate or ammonia. This low growth rate (compared to
66 other phytoplankton species) mainly constrains their ecological niches to nitrate-poor regions, where they can be
67 competitive. Moreover, their geographical distribution is constrained by nutrient availability in the photic layer (mainly
68 iron and phosphate) (Berman-Frank, 2001; Bonnet et al., 2009; Mills et al., 2004; Moutin et al., 2005, 2008; Rubin et
69 al., 2011; Rueter, 1988) and temperature (Staal et al., 2003). *Trichodesmium* sp. are present only in water where
70 temperature is above 20°C (Capone, 1997; LaRoche and Breitbarth, 2005; Montoya et al., 2004), while some UCYN
71 can be found in colder and deeper waters (Bonnet et al., 2015a; Church et al., 2005; Moisaner et al., 2010)

72 The spatial distribution and rates of N_2 fixation have been inferred at the global scale using several tools. (Deutsch et
73 al., 2007) have introduced the tracer P^* which represents the excess of P relative to the standard N quota. A decrease in
74 this tracer is then interpreted as N_2 fixation, since N_2 fixation extracts PO_4 alone. More recently, (Luo et al., 2014)



75 developed a multiple linear regression that relates N_2 fixation from the MAREDAT database (Luo et al., 2012) to
76 environmental conditions (nutrients, SST, irradiance, MLD,...) in order to build a statistical model for global N_2 fixation
77 distribution. Finally, numerical models have also been used and they allow to overcome the scarcity of observations that
78 may limit the two previous approaches. Indeed, models can be used to investigate the spatial and temporal variability of
79 dinitrogen fixation and to study the controlling environmental factors.

80

81 Among those studies focusing on the spatial distribution of dinitrogen fixation, (Berthelot et al., 2017; Bonnet et al.,
82 2009, 2015a; Garcia et al., 2007; Shiozaki et al., 2014) based on oceanographic campaigns have reported high N_2
83 fixation rates in the Western Tropical South Pacific (WTSP), that has been recently identified as a globally important
84 hot spot of N_2 fixation with rates $> 600 \mu\text{mol N m}^{-2}\cdot\text{d}^{-1}$ (Bonnet et al., 2017). Very high abundances of *Trichodesmium*
85 have been historically reported in this region (Dupouy et al., 2000, 2011; Moisander et al., 2008; Neveux et al., 2006;
86 Shiozaki et al., 2014); Stenegren et al., This issue) and have recently been identified as the major contributor to N_2
87 fixation in this region (Berthelot et al., 2017); Bonnet et al., This issue). However, the reasons for such an ecological
88 success of diazotrophs in this region are still poorly understood.

89 In this study, we aim at bringing new insights on this known, but poorly understood, “nitrogen fixation hotspot”. This
90 study ambitions to understand the spatial and temporal distribution (i.e. seasonal variability) of *Trichodesmium* and to
91 evaluate the potential impact of *Trichodesmium* fixers on the biogeochemical conditions of the WTSP. We will
92 specifically address the following overarching questions: (i) What are the mechanisms that structure the *Trichodesmium*
93 distribution in the WTSP, particularly around the South West Pacific islands, and (ii) what is the biogeochemical impact
94 of N_2 fixation in this region? Noteworthy, this study is also taking advantage of the sampling done during the
95 OUTPACE cruise, which nicely complement the data coverage in the south west Pacific, and allow a better
96 characterization of the processes responsible for the spatial and seasonal variability of the N_2 fixation.

97 To fulfill our objectives, we have implemented an explicit representation of the nitrogen fixers in a biogeochemical
98 model, based on the *Trichodesmium* physiology. The first section of this study describes the experimental design and the
99 observation used in our study, while the second part of the paper provides a validation of our reference simulation with
100 an analysis of the *Trichodesmium* compartment and its impacts on the biogeochemical conditions of the Tropical
101 Pacific. In the discussion, the impact of iron from islands sediment on dinitrogen fixation is considered as well as the
102 added value of an explicit dinitrogen fixer compartment rather than a simpler implicit representation of dinitrogen
103 fixation. Finally, implications and limits of our modeling exercise are detailed in the conclusion.

104

105 2. Methods

106 2.1 Coupled dynamical (ROMS)-primary production (PISCES) model

107

108 In this study, we used a coupled dynamical-biogeochemical framework based on the regional ocean dynamical model
109 ROMS (Regional Oceanic Modeling System, (Shchepetkin and McWilliams, 2005) and the state of the art
110 biogeochemical model PISCES (Pelagic Interactions Scheme for Carbon and Ecosystem Studies). The ocean model
111 configuration is based on the nested version of ROMS (Penven et al., 2006) and covers the tropical Pacific region
112 [33°S-33°N;110°E-90°W]. It has 41 terrain-following vertical levels with 2-5 m vertical resolution in the top 50 meters
113 of the water column, then 10-20 m resolution in the thermocline and 200-1000 m resolution in the deep ocean. The



114 horizontal resolution is 1° . The turbulent vertical mixing parameterization is based on the non-local K profile
115 parameterization (KPP) of (Large et al., 1994). Open boundary conditions are treated using a mixed active/passive
116 scheme (Marchesiello et al., 2001). This scheme allows to force our regional configuration with large-scale boundary
117 conditions from a $\frac{1}{2}^\circ$ global model simulation (details available in (Couvelard et al., 2008), while allowing anomalies to
118 radiate out of the domain. The use of similar ROMS configurations in the WTSP is largely validated through studies
119 demonstrating skills in simulating both the surface (Jullien et al., 2012, 2014; Marchesiello et al., 2010) and subsurface
120 ocean circulation (Couvelard et al., 2008). To compute the momentum and fresh water/heat fluxes, we use a
121 climatological forcing strategy. The momentum forcing is computed from a 1993-1996 7-day climatology of the ERS1-
122 2 scatterometer stress (<http://cersat.ifremer.fr/oceanography-from-space/our-domains-of-research/air-sea-interaction/ers-ami-wind>). Indeed, ERS derived forcing has been shown to produce adequate simulations of the Pacific Ocean
123 dynamics (e.g. (Cravatte et al., 2007)). Heat and fresh water forcing is computed from the COADS climatology 1-day
124 averaged outputs are stored for analysis. ROMS forces on line a biogeochemical model with the noticeable use of a
125 WENO5 advection scheme (i.e. five order weighted essentially non-oscillatory scheme; Shchepetkin and McWilliams,
126 1998).
127

128 The biogeochemical model PISCES simulates the marine biological productivity and the biogeochemical cycles of
129 carbon and the main nutrients (P, N, Si, Fe) (Aumont and Bopp, 2006a). In PISCES, there are five modeled limiting
130 nutrients for phytoplankton growth: nitrate, ammonium, phosphate, silicate and iron. Those nutrients are delivered to
131 the ocean through dust deposition, river runoff and mobilization from the sediment. In addition to the nutrients,
132 Dissolved Inorganic Carbon (DIC), total alkalinity and dissolved oxygen are also simulated. Three non-living organic
133 compartments are represented: semi-labile dissolved organic matter, small sinking particles, and large sinking particles.
134 In this study, we used a modified version of PISCES which differs in the use of a full quota formulation (with only
135 variable Redfield ratios; (Droop, 1983) rather than the mixed Monod-quota approach (with fixed ratios for nitrogen,
136 phosphorus and silica; Monod, 1942) used in the standard PISCES version (Aumont and Bopp, 2006). In this “quota”
137 version of PISCES (Kwiatkowski et al, 2017, submitted), the phytoplankton growth is limited by the internal
138 availability in nutrients. For the purpose of this study, we also implemented in this quota version an explicit
139 representation of *Trichodesmium*. Therefore, five living compartments are modeled with three phytoplankton groups
140 (nanophytoplankton, diatoms, and *Trichodesmium*) and two zooplankton groups (microzooplankton, and
141 mesozooplankton).

142 In our configuration, the growth rate of *Trichodesmium* is limited by light, temperature, phosphorus and iron
143 availability. *Trichodesmium* growth rate is computed as follows:

$$144 \text{ Growth rate} = (\mu_{\text{FixN}_2} + \mu_{\text{TriNO}_3} + \mu_{\text{TriNH}_4}),$$

145 where μ_{FixN_2} denotes growth due to dinitrogen fixation, μ_{TriNO_3} and μ_{TriNH_4} represent growth sustained by NO_3^- and
146 NH_4^+ uptake, respectively. Dinitrogen fixation is activated when reactive nitrogen species are limiting. Otherwise,
147 *Trichodesmium* grows on NO_3^- and NH_4^+ just like standard nanophytoplankton do. Moreover, a fraction of fixed
148 nitrogen is released by the simulated *Trichodesmium*. (Berthelot et al., 2015) estimated this fraction at less than 10%
149 while considering all diazotrophs. Because we are only representing *Trichodesmium*, we set up this fraction at 5% of
150 total amount of fixed nitrogen.

151 Dinitrogen fixation is controlled by the availability of phosphate, iron and light and is modulated by temperature. The
152 complete set of equations of *Trichodesmium* is detailed in Appendix 1. This setup reproduces the dinitrogen fixation
153 through an explicit representation of the *Trichodesmium* biomass (to be compared with often used implicit
154 parameterization that links directly environmental parameters to nitrogen fixation without requiring the *Trichodesmium*



155 biomass to be simulated).

156

157 **2.2 Setup of experiments**

158

159 Below are summarized the set of experiments that have been performed in this study (Table 1). A first simulation over
160 20-years (1993-2013) has been performed as a reference experiment, hereafter referred to as "TRI". This reference
161 simulation uses the explicit dinitrogen fixation module described above. In a second experiment called
162 "TRI_NoFeSed", the model setup is identical to the reference experiment, except that iron input from the sediments is
163 turned off between 156°E and 240°E. In a third experiment "N2_imp", the explicit dinitrogen fixation module is
164 replaced by the implicit parameterization described in (Aumont et al., 2015). Finally, a fourth experiment "N2_Wo"
165 corresponds to a model setup in which no explicit nor implicit description of dinitrogen fixation is activated.
166 Comparison between TRI and TRI_NoFeSed experiments allow to estimate the impact of iron input from island
167 sediments on the dinitrogen fixation, while the impact of dinitrogen fixation on the biogeochemical conditions in the
168 Pacific Ocean can be investigated by comparing TRI and N2_Wo. Finally, the TRI and N2_imp experiments are used to
169 evaluate the added value of an explicit description of dinitrogen fixation relative to an implicit inexpensive
170 parameterization.

171

172 **2.3 Observational datasets.**

173

174 Several different databases have been used to evaluate the model skills. For nitrate and phosphate, the CSIRO ½°
175 global Atlas of Regional Seas (CARS, <http://www.marine.csiro.au/~dunn/cars2009/>) has been used. Iron has been
176 evaluated with the global database from (Tagliabue et al., 2012), to which the dissolved iron data from the OUTPACE
177 cruise (Guieu et al., under review) have been added. This database is a compilation of 13125 dissolved iron
178 observations covering the global ocean and encompassing the period 1978–2008. The global MARine Ecosystem DATA
179 (MAREDAT, <https://doi.pangaea.de/10.1594/PANGAEA.793246>) database of N₂ fixation has been expanded with data
180 from recent cruises performed in the WTSP (MOORSPICE, (Berthelot et al., 2017), DIAPALIS ((Garcia et al., 2007),
181 NECTALIS (<http://www.spc.int/oceanfish/en/ofpsection/ema/biological-research/nectalis>), PANDORA (Bonnet et al.,
182 2015a), OUTPACE (Bonnet et al., This issue), Mirai (Shiozaki et al., 2014) has been used for dinitrogen fixation rates.
183 This database contains 3079 data points at the global ocean scale, of which ~1300 are located in our simulation region
184 (Luo et al., 2012). Finally, we have used surface chlorophyll concentrations from the GLOBCOLOUR project
185 (<http://ftp.acri.fr>) which spans the 1998-2013 time period.

186

187 **3.Results**

188 **3.1 Model Validation**

189

190 In this subsection, we aim at validating our reference simulation "TRI" with the data previously mentioned. In the
191 Pacific, phosphate and nitrate concentrations show maxima in the upwelling regions, i.e. along the western American
192 coast, and in the equatorial upwelling (Fig. 1a,c), and minima in the subtropical gyres. First, phosphate patterns show



193 modeled values and structures in qualitatively good agreement with observations. In contrast, the nitrate structure shows
194 some biases. We observe concentrations higher than $1 \mu\text{mol.L}^{-1}$ all along the equator in CARS, while in the model
195 nitrate concentrations are lower than this value west of 170°W . More generally the model tends to underestimate nitrate
196 concentrations.

197 The regions most favorable for *Trichodesmium* can be defined by temperature within $26\text{--}29^{\circ}\text{C}$. The model reproduces
198 relatively well the spatial distribution of this temperature preferendum. This distribution exhibits a significant seasonal
199 variability, mainly as a result of the variability of the 26°C isotherm. The latter moves by $\sim 5^{\circ}$ latitudinally between
200 summer and winter in the WTSP, and by $\sim 15^{\circ}$ in the Western Tropical North Pacific (WTNP) (Fig. 1a). Along the
201 equator, this isotherm migrates by 15° eastward during summer (Fig. 1a). This temporal variability is well reproduced
202 by the model (Fig. 1b). In contrast, nitrate and phosphate seasonal variability remains low (not shown).

203 Another important feature that needs to be properly reproduced by the model is the iron distribution in the upper ocean.
204 The median value as well as the dispersion of the iron surface concentrations over the tropical Pacific, are displayed for
205 both the data and the model in Figure 2a. No statistical differences can be distinguished, the model being sampled at the
206 same time and same location as the data. This latter result shows a good agreement between the data and the model at
207 the tropical Pacific scale (Fig 2b compared to Fig. 2c). The best sampled area is the central Pacific ocean where
208 simulated iron concentrations are low (0.1 to $0.3 \text{ nmol Fe.L}^{-1}$), which is consistent with the observations. The southwest
209 Pacific is characterized by relatively high surface iron concentrations, between 0.4 and $0.8 \text{ nmol Fe.L}^{-1}$, both in the data
210 and in the model. Large scale patterns are thus well represented by the model. Nevertheless, the model tends to
211 overestimate iron levels in the south Pacific gyre, between 180° and 140°W at about 20°S .

212 Figure 3 displays a comparison between surface Chlorophyll concentrations from GLOBCOLOUR data (a), and from
213 TRI (b) and TRI_imp (c) simulations. Strong chlorophyll concentrations are found in the eastern equatorial Pacific
214 upwelling and along Peru in both the observations and our 2 simulations, with mean values of $0.3 \text{ mg Chl.m}^{-3}$. The
215 equatorial rich tongue simulated by the model (Fig. 3b,c) is however too narrow compared to the observations,
216 especially in the northern hemisphere. Similarly, the model is unable to simulate the elevated chlorophyll levels around
217 the Costa Rica dome and the localized enhanced chlorophyll off Papua New Guinea. In TRI (Fig. 3b), chlorophyll
218 values in the South West Pacific region vary between 0.1 and $0.2 \text{ mg Chl.m}^{-3}$, with maxima located in the vicinity of the
219 Fiji and Vanuatu islands. These values are within the range of the data, even if the data tend to be slightly higher (up to
220 $0.3 \text{ mg Chl.m}^{-3}$ near the coasts). The spatial structure is well represented, with maxima simulated around the islands.
221 Those localized chlorophyll enhancement suggest a specific island effect. In the subtropical gyres, the simulation
222 predicts chlorophyll concentrations of $\sim 0.05 \text{ mg Chl.m}^{-3}$ which are higher than the observations ($< 0.025 \text{ mg Chl.m}^{-3}$).
223 In contrast in TRI_imp (Fig. 3c), chlorophyll values in the South West Pacific and in the North hemisphere are too low
224 in comparison with the ocean colour data (Fig. 3a). TRI simulation thus appears in better agreement with the
225 observations than TRI_imp.

226 Part of the surface chlorophyll in Figure 3b is associated to *Trichodesmium*. The Figure 3d shows the annual mean
227 spatial distribution of surface *Trichodesmium* chlorophyll in the “TRI” simulation. This distribution displays two zonal
228 tongues in the tropics, one in each hemisphere. Maximum values are located in the South West Pacific (around Vanuatu
229 archipelago, New Caledonia, Fiji, and Papua New Guinea) and around Hawaii, where they reach $0.06 \text{ mg Chl.m}^{-3}$. In
230 the south Pacific, high chlorophyll biomass extends eastward until 130°W . Further east, concentrations drop to below
231 $0.02 \text{ mg Chl.m}^{-3}$. It's important to note that in the observations *Trichodesmium* has never observed beyond 170°W . This
232 bias in the model could be explain by the overestimation of iron concentrations in SPG. In the Northern Hemisphere,
233 between the coast of Philippines (120°E) and Hawaii (140°W), *Trichodesmium* chlorophyll concentrations are greater



234 than $0.03 \text{ mg Chl.m}^{-3}$. In the North East Pacific, *Trichodesmium* chlorophyll is lower, yet significant ($<0.03 \text{ mg Chl.m}^{-3}$).
235 Otherwise the equatorial Pacific and South-east Pacific oceans are overall poor in *Trichodesmium*.
236 In Figure 4, the dinitrogen fixation rates predicted by the model in “TRI” are compared to the observations from the
237 MAREDAT expanded database. Evaluation of the model behavior remains quite challenging because of the scarcity of
238 the observations. Some large areas are not properly sampled such as the north west tropical Pacific and the eastern
239 Pacific. Nevertheless, some regional patterns emerge from the observations. Maximum fixation rates (~ 600 to 1600
240 $\mu\text{mol N.m}^{-2}.\text{d}^{-1}$; Fig. 4a) are observed around the south west Pacific islands, in the Solomon Sea, around the Melanesian
241 archipelagoes and around Hawaii, four well known « hotspots » of N_2 fixation (Berthelot et al., 2015, 2017, Bonnet et
242 al., 2009, 2017; Böttjer et al., 2017). The modeled regional patterns of strong fixation are coherent with the observations
243 (Fig. 4b), showing values in the same range. In the south Pacific, the TRI simulation is able to reproduce the strong
244 east-west increasing gradient of N_2 fixation as reported by dinitrogen fixation (Shiozaki et al., 2014; Bonnet et al., This
245 issue; Fig 4c,d). In the equatorial Central Pacific, modeled values of mean fixation are negligible ($< 0.5 \mu\text{mol N.m}^{-2}.\text{d}^{-1}$)
246 in contrast to the observations which suggest low but non-negligible fixation rates (between 1 to $2 \mu\text{mol N.m}^{-2}.\text{d}^{-1}$;
247 (Bonnet et al., 2009; Halm et al., 2012). In general, dinitrogen fixation rates are overestimated by $\sim 70\%$ in TRI
248 compared to the data. Some recent studies have shown that the $^{15}\text{N}_2$ -tracer addition method (Montoya et al., 2004) used
249 in most studies reported in the MAREDAT database may underestimate N_2 fixation rates due to an incomplete
250 equilibration of the $^{15}\text{N}_2$ tracer in the incubation bottles, which may explain the differences observed between the
251 modeled and measured rates (Großkopf et al., 2012; Mohr et al., 2010). However, some other studies performed in the
252 South Pacific (Bonnet et al., 2016b; Shiozaki et al., 2015) compared the two methods, and did not found any significant
253 differences.

254

255 3.2 *Trichodesmium* Primary Production

256 We evaluated the direct relative contribution of *Trichodesmium* to PP (Fig. 6). The spatial distribution of this
257 contribution is very similar to the spatial distribution of *Trichodesmium* chlorophyll, with 2 distinct tongues located on
258 each side of the Equator in the tropical domain. In the Northern hemisphere, the tongue extends from the coast of
259 Philippines (120°E) to Hawaii (140°W) longitudinally and between 10°N and 25°N latitudinally. The maximum
260 contribution ($\sim 35\%$) is reached near Hawaii while in the rest of the tongue, values are close to 20%. In the Southern
261 Hemisphere, the region of elevated contribution extends from PNG (140°E) to about the center of the South Pacific
262 subtropical gyre at 130°W , and between 5°S and 25°S latitudinally. Maximum values are predicted in the vicinity of
263 Vanuatu and Fiji Islands, where they can reach 35%. Part of this elevated contribution is explained by the very low PP
264 rates simulated in this region for both nanophytoplankton and diatoms (less than $0.03 \text{ mol C.m}^{-3}.\text{yr}^{-1}$). Furthermore, the
265 island effect seems to represent an important factor for explaining the spatial distribution of *Trichodesmium* growth
266 rates. Indeed, maximum *Trichodesmium* chlorophyll concentrations and the largest contribution of *Trichodesmium* to PP
267 are achieved near the islands. Finally, in LNLC regions (red boxes; fig. 1c,d), we assess that *Trichodesmium* contribute
268 to 15% of total PP, which is in accordance with biogeochemical studies performed in these areas (Bonnet et al., 2015;
269 Berthelot et al., 2017; Caffin et al., This issue)

270

271 3.3 Seasonal variability of *Trichodesmium* biomass

272 Simulated dinitrogen fixation rates and *Trichodesmium* biomass (not shown) display a seasonal variability that is driven



273 by the seasonal variability of the environmental conditions (light, temperature, currents, nutrients). The regional
274 maxima of *Trichodesmium* biomass (exceeding 3 mmol C.m^{-2} ; integrated over the top 100m of the ocean) are found in
275 both hemispheres during the summer season (Fig. 7a,e) even if locally, maxima can be attained during other periods of
276 the year than summer. In the south Pacific, the area of elevated *Trichodesmium* biomass moves by 3° southward from
277 austral winter to austral summer. Along Australia and in the Coral Sea, *Trichodesmium* biomass exhibits a large seasonal
278 variability with very low winter biomass that contrast with elevated values in summer. A similar important variability,
279 which is shifted by six months, is simulated in the Northern hemisphere in the Micronesia region and in the Philippine
280 Sea.

281 Unfortunately, due to the scarcity of N_2 fixation data over the annual cycle, this seasonal cycle cannot be properly
282 assessed at the scale of the tropical Pacific Ocean. This is only feasible at the time series station ALOHA located in the
283 North Pacific gyre at $22^\circ 45', 158^\circ \text{W}$, where seasonal data of dinitrogen fixation are available from 2005 to 2012 (Böttjer
284 et al., 2017). They proved that vertically integrated dinitrogen fixation rates are statistically significantly lower from
285 November to March (less than $200 \mu\text{mol N m}^{-2}.\text{d}^{-1}$) than from April to October (about $263 \pm 147 \mu\text{mol N.m}^{-2}.\text{d}^{-1}$) as
286 highlighted in Figure 5a (blue dots). In the model (red dots; Fig. 5a), the maximum amplitude of the seasonal cycle
287 appears to be underestimated relative to the observations (i.e. respectively $\sim 170 \mu\text{mol N.m}^{-2}.\text{d}^{-1}$ and $\sim 250 \mu\text{mol N.m}^{-2}.\text{d}^{-1}$)
288 ¹). Dinitrogen fixation peaks one month earlier in the model than in the data (August for the model and September for
289 the data). The simulated dinitrogen fixation rates are minimum between December and May (averaging $\sim 241 \pm 27$
290 $\mu\text{mol N.m}^{-2}.\text{d}^{-1}$) and maximum the rest of the year (averaging $\sim 347 \pm 52 \mu\text{mol N.m}^{-2}.\text{d}^{-1}$). These values are comparable
291 to the data even if they are slightly higher.

292 In order to assess the seasonal cycle of N_2 fixation rates in the south Pacific (red box Fig 1c; $160^\circ\text{E}-230^\circ\text{E}$; $25^\circ\text{S}-14^\circ\text{S}$),
293 we have extracted the available data for each month from our database (blue dots; Fig. 5b), and the corresponding
294 model values in TRI (red dots; Fig 5b). In July no observations are available and in January, April and August, only one
295 data point is available for the entire region. The predicted seasonal cycle is broadly consistent with observations.
296 Minimum dinitrogen fixation rates ($239 \pm 205 \mu\text{mol N.m}^{-2}.\text{d}^{-1}$) occur during austral winter and autumn. Maximum rates
297 are reached in February and March, where they exceed $600 \mu\text{mol N.m}^{-2}.\text{d}^{-1}$ in the observations. The increase in
298 dinitrogen fixations rates occurs one month earlier than in the observations, in December instead of January, and
299 remains two to three fold higher from April to June. It is important to note here that the sampling spatial and temporal
300 distribution may distort the seasonal cycle. Using the model, it is possible to evaluate how well the seasonal cycle is
301 captured by the sampling (red dots compared to green dots; Figure 5b). The general structure of the seasonal cycle
302 remains relatively unaltered. However, the amplitude is significantly impacted since it reaches $1100 \mu\text{mol N.m}^{-2}.\text{d}^{-1}$ in
303 the sampling whereas it is about twice as low at $600 \mu\text{mol N.m}^{-2}.\text{d}^{-1}$ when all the model data points are considered. We
304 can conclude that the TRI simulation reproduces well the seasonal cycle of N_2 fixation rates at the Pacific scale, even
305 though more data are needed to improve the evaluation of the model skills.

306 To further investigate the mechanisms that drive the seasonal variability of *Trichodesmium* in the Pacific, we examined
307 the factors that control *Trichodesmium* abundance in the TRI simulation (not shown). This analysis indicates that this
308 seasonal variability is mainly controlled by primary production. Hence we further examine the limitation terms of
309 primary production (Fig. 8) in two representative regions characterized by elevated levels of N_2 fixation rates (red
310 boxes; Fig 1c). A detailed description of these limitation terms is given in Appendix 1. A limitation term reaching 1
311 means that growth is not limited whereas a limitation term equal to zero means that growth ceases.

312 *Trichodesmium* growth sustained by nitrate and ammonia is very small in LNLC regions due to their very low
313 availability and is therefore not considered further. Thus, our analysis is restricted to dinitrogen fixation. *Trichodesmium*



314 growth can be limited by iron and phosphate and is inhibited when reactive nitrogen (nitrate and ammonia) is available.
315 In the WTSP, the model suggests that iron is the sole nutrient that modulates *Trichodesmium* growth (red curve; Fig.
316 8a,b). The others limiting factors of *Trichodesmium* growth are light (green curve; Fig 8a,b) and temperature (purple
317 curve; Fig. 8a,b). The product of these 3 limiting factors gives the limiting coefficient of dinitrogen fixation (brown
318 curve; Fig 8a,b). The limiting factors vary according to the season and the hemisphere. In the south (north) Pacific,
319 temperature and light are less limiting during the austral summer (winter) than during the austral winter (summer). The
320 limiting factor associated to temperature varies from 0.8 to 1, and the light limiting factor varies from 0.15 to 0.3.
321 Unlike light and temperature, iron is less (more) limiting in the south (north) Pacific during winter (summer) than
322 during the austral summer (winter) with values varying between 0.4 and 0.7. Finally, *Trichodesmium* growth is more
323 limited during the austral winter (summer) in the south (north) Pacific. The seasonal variability is forced by light and
324 temperature, and iron mitigates its amplitude. Indeed, temperature and iron are anti-correlated because, in the boxes
325 over which this analysis is performed, the ocean dynamic is mainly 1D. Nutrients are thus brought by enhanced vertical
326 mixing which also cools temperature.

327

328 4. Discussion

329

330 The implementation of dinitrogen fixation in ocean models has become more and more complex over the past twenty
331 years. At first, the supply of new nitrogen by dinitrogen fixation has been embedded into models by developing implicit
332 parameterizations (Bissett et al., 1999). Subsequently explicit descriptions of diazotrophs have been developed, mainly
333 based on the knowledge derived from laboratory experiments focused on *Trichodesmium* sp. (Fennel et al., 2001; Hood
334 et al., 2001; Moore et al., 2001). Since the mid-2000s, studies focused on the role of iron in controlling the distribution
335 of diazotrophs and dinitrogen fixation (Keith Moore et al., 2006; Krishnamurthy et al., 2009; Moore et al., 2004;
336 Tagliabue et al., 2008).

337

338 4.1 Impact of iron from island sediments

339

340 Monteiro et al. (2011) and Dutkiewicz et al. (2012) have suggested that a realistic representation of marine iron
341 concentration is key to simulate the diazotrophs habitat. Monteiro et al. (2011) performed a sensitivity study and found
342 that a fivefold increase in the solubility of aeolian iron improves the biogeographical distribution of N₂ fixation in the
343 southwest Pacific. In the meantime, a recent study has challenged this view by showing no increase in dinitrogen
344 fixation in response to increased dust deposition (Luo et al., 2014). In any cases, the sedimentary and hydrothermal
345 sources were not taken into account in those studies, although they are likely significant sources (Bennett et al., 2008;
346 Johnson et al., 1999; Moore et al., 2004; Tagliabue et al., 2010; Toner et al., 2009). In parallel, Dutkiewicz et al. (2012)
347 evaluated the sensitivity of the biogeographical distribution of N₂ fixation to the aeolian source of iron in a model which
348 takes into account the iron sediment supplies, and conclude for minor changes in south west Pacific, while in the north
349 Pacific the change was larger. Indeed, there are many high islands that could deliver significant amounts of iron to the
350 ocean (Radic et al., 2011) in the southwest Pacific.

351 To assess the impact of the sediment source of iron on the *Trichodesmium* production, we used the “TRI_NoFeSed”
352 experiment in which this specific source of iron has been turned off for the islands between 160°E and 120°W (Table 1).



353 In this simulation, the iron and the *Trichodesmium* chlorophyll concentrations decrease by 58% and 51% respectively
354 (Fig. 9a,b), in the WTSP (red box Fig 1c). Figure 9c displays the iron distribution simulated in TRI_NoFeSed, and
355 shows that the maximums around the islands disappear. Furthermore, in the south Pacific, iron decreases due to the
356 reduction of the zonal advection of iron downstream of the islands. The iron flux from the sediments around the islands
357 also affects the spatial structure of *Trichodesmium* chlorophyll (Fig. 9d,e), most noticeably in the south Pacific, with
358 maxima shifted from the south Pacific islands region (e.g. Fiji, New Caledonia, Vanuatu) in the TRI simulation to the
359 coastal regions near Australia and Papua New Guinea in the TRI_NoFeSed simulation. In the northern hemisphere, the
360 effects of the sediment flux of iron are less important with a shift of the *Trichodesmium* chlorophyll maxima towards the
361 Philippine Sea and a localized effect near Hawaii. This sensitivity test demonstrates that *Trichodesmium* are highly
362 sensitive to the iron distribution in our model and hence that the spatial patterns of *Trichodesmium* chlorophyll in the
363 south west Pacific are tightly controlled by the release of iron from the coastal sediments of the Pacific islands.

364

365 4.2 *Trichodesmium* impacts on biogeochemistry

366

367 One of the questions we want to address is the quantification of the *Trichodesmium* impact on primary production (PP),
368 at the Pacific scale with a focus on the WTSP region. In the oligotrophic waters of the south Pacific, dinitrogen fixation
369 can be an important source of bio-available nitrogen in the water column through *Trichodesmium* recycling which can
370 feed other phytoplankton. To evaluate that impact, we calculated the relative increase of PP between the TRI simulation
371 and the N2_Wo simulation in which no dinitrogen fixation is considered (Fig 10a). As expected, the spatial structure of
372 the primary production differences between both simulations matches the dinitrogen fixation spatial distribution in the
373 TRI simulation (two tongues, one in each hemisphere). In the north Pacific the maximum increase of the primary
374 production due to the dinitrogen fixation is located around Hawaii, where it exceeds 120%. In the remaining part of the
375 northern hemisphere tongue, primary production increases by 50% to 100%. In the southern hemisphere, values are
376 more homogeneous in the tongue (from 80 to 100%), even though there is a local maximum around Fiji and Vanuatu
377 (up to 120%). Out of these northern and southern tongues, the increase of PP is less than 20%. In average on our
378 domain, the increase of PP is 19% and in LNLC regions, it reaches approximately 50%.

379 From total primary production only, it is not possible to disentangle the increase of primary production directly due to
380 *Trichodesmium* themselves and the indirect increase due to the impact of dinitrogen fixation on the other phytoplankton
381 groups (nanophytoplankton and diatoms). Indeed, as mentioned in the method section, *Trichodesmium* also releases a
382 fraction of the recently fixed N₂ as bio-available nitrogen (mostly ammonia and dissolved organic nitrogen). Figure 10b
383 displays the difference of PP due to diatoms and nanophytoplankton only. The main large scale patterns constituted of
384 the northern and southern tongues persist, but the intensity of the differences contrasts with those found when
385 considering total primary production (Fig. 10a). Indeed, the increase of total primary production (Fig 10a) in those two
386 tongues is twice as high as when the direct effect of *Trichodesmium* is excluded. This analysis stresses the importance of
387 the bio-available nitrogen released by diazotrophs as we attribute about half of the total production increase to this
388 release. Indeed, recent isotopic studies tracing the passage of diazotroph-derived nitrogen into the planktonic food web
389 reveal that part of the recently fixed nitrogen is released to the dissolved pool and quickly taken up (24-48%) by
390 surrounding planktonic communities (Berthelot et al., 2016; Bonnet et al., 2016a, 2016b)).

391 With the simulation N2_imp, we aim at comparing an implicit dinitrogen fixation formulation to the explicit
392 formulation used in TRI. Figure 10c displays the relative change of total PP between the TRI and the N2_imp



393 simulations (see Table 2). The implicit formulation displays a similar spatial distribution than the explicit distribution
394 but is predicting a lower total primary production, especially in the the southern Pacific where explicit formulation
395 leads to an increase of about 45% in total PP compared to the one related to the implicit formulation. On average across
396 our domain, total primary production is about ~ 9% higher when nitrogen fixation is explicitly modeled relative to an
397 implicit formulation.

398 This difference becomes even weaker (2%) if only primary production by nanophytoplankton and diatoms is
399 considered, with noticeable differences restricted to the areas of maximum dinitrogen fixation in the southern
400 hemisphere (around the islands). PP sustained by the release of bio-available nitrogen is thus similar in the TRI and
401 N2_imp simulations, but an explicit representation of dinitrogen fixation allows a better description of dinitrogen
402 fixation patterns. Indeed, the areas of intense dinitrogen fixation rates cannot be properly simulated in the vicinity of the
403 islands, especially in the southern hemisphere, by the tested implicit parameterization.

404

405 4.3 Limitations

406

407 In this study, we simulate dinitrogen fixation through the explicit representation of only one type of diazotrophs, the
408 *Trichodesmium* sp. This choice has been motivated by evidences that it represents one of the main nitrogen fixers in the
409 western tropical Pacific (Bonnet et al., 2015a; Dupouy et al., 2011; Shiozaki et al., 2014) and by the relatively good
410 knowledge (compared to other dinitrogen fixers) we have about its physiology (Ramamurthy and Krishnamurthy, 1967;
411 Ohki et al., 1992; Mulholland and Capone, 2000; Mulholland et al., 2001; Küpper et al., 2008; Rubin et al., 2011;
412 Bergman et al., 2013). However, our model remains simple and some of the mechanisms that drive the behavior of
413 *Trichodesmium* have not been implemented in our model. As an example, the ability of *Trichodesmium* to group in
414 colonies and to vertically migrate (Kromkamp and Walsby, 1992; Villareal and Carpenter, 2003; Bergman et al., 2013)
415 (Bergman et al., 2013; Kromkamp and Walsby, 1992; Villareal and Carpenter, 2003) is well documented. The reason of
416 these mechanisms remains unclear, but several hypotheses have been put forward such as avoiding nitrogenase
417 exposition to di-oxygen (Carpenter, 1972; Gallon, 1992; Paerl et al., 1989), or maximizing light (on the surface) and
418 nutrients (at depth) acquisition (Letelier and Karl, 1998; Villareal and Carpenter, 1990; White et al., 2006), or even
419 increasing the efficiency of the uptake of atmospheric iron (Rubin et al., 2011). Our model does not represent those
420 processes, nor does it model the resulting vertical migration of *Trichodesmium*. Moreover, the release of fixed
421 dinitrogen as reactive nitrogen bioavailable to other phytoplanktonic organisms has been set to a constant value of 5%.
422 This percentage is known to be highly variable and therefore this value is in the lowest range of the observations. An
423 increase in this value would increase the PP due to nanophytoplankton and diatoms in the TRI simulation, and thus
424 decrease the relative contribution of *Trichodesmium* to total PP, which would be closer to the last observations
425 (Berthelot et al. 2017; Bonnet et al. 2017).

426 Some studies, mostly based on extrapolated in-situ data, aimed at assessing the potential of dinitrogen fixation at global
427 or regional scale (Codispoti et al., 2001; Deutsch et al., 2001; Galloway et al., 2004). In the western south tropical
428 Pacific, Bonnet et al (2017) have estimated total nitrogen fixation at 15 to 19 Tg N.yr⁻¹. For the same region, nitrogen
429 fixation is predicted to amount to ~7 Tg N.yr⁻¹ in the TRI simulation. As already mentioned, this rather low predicted
430 estimates might be explained by the sole representation of *Trichodesmium* as nitrogen fixing organisms, which
431 dominate in the western tropical south Pacific (Dupouy et al., 2011, Stenegren et al., This issue). It has to be noted that
432 other diazotroph groups such UCYN-B and DDAs are abundant in the WTSP, representing 10-20% of the overall



433 diazotroph community (Stenegren et al., This issue). Moreover, the contribution of heterotrophic diazotrophic organism
434 is poorly studied and may account for a significant fraction of N_2 fixation (Moisander et al., 2017). Our model
435 estimation has also been computed from monthly averages and is thus not taking into account the high-frequency
436 variability that may explain at least some of the very high rates of dinitrogen fixation used in the study by Bonnet et al.
437 (2017). Our assessment based on a model could thus be seen as a lower limit for dinitrogen fixation in the western
438 tropical Pacific. Moreover, our model shows also a good qualitative agreement with the studies based on observations
439 that focus on the impact of dinitrogen fixation in tropical oligotrophic waters (Raimbault and Garcia, 2008; Shiozaki et
440 al., 2013). Indeed, in agreement with those studies, our reference simulation predicts that diazotrophs support a large
441 part of PP (~ 15%) in LNLC regions.

442 5. Conclusion

443 This study describes the spatial and temporal distribution of *Trichodesmium* at the scale of the tropical Pacific ocean,
444 and investigates the impact of a major diazotroph species (e.g *Trichodesmium* sp.) on the biogeochemistry of this
445 region. Toward this end, we performed a first 20-year simulation with the coupled 3D dynamical biogeochemical model
446 ROMS-PISCES in which we embedded an explicit representation of dinitrogen fixation based on *Trichodesmium*
447 physiology. This simulation was shown to be able to reproduce the main physical (SST) and biogeochemical (nutrients)
448 conditions of the tropical Pacific ocean. This includes the spatial distribution of surface chlorophyll and dinitrogen
449 fixation.

450 The validation of this simulation allows us to confidently assess the *Trichodesmium* distribution. The model predicts
451 that areas favorable to *Trichodesmium* growth extend from 150°E to 120°W in the south Pacific, and from 120°E to
452 140°W in the north Pacific, with local optimal conditions around the islands (i.e., Hawaii, Fiji, Samoa, New Caledonia,
453 Vanuatu). This broadly corresponds to the LNLC regions where *Trichodesmium* are predicted to be responsible for
454 15% of total primary production. The seasonal variability of the *Trichodesmium* habitat is dominantly controlled by
455 SST and light, while iron availability modulates the amplitude of the seasonal cycle.

456 In our study we also assess the role played by iron released from the island sediments, and show that this iron source
457 partly controls the spatial structure and the abundance of *Trichodesmium* in the western tropical south Pacific. However,
458 this region is in the center of the south Pacific convergence zone which is the largest convective area of the Southern
459 Hemisphere, with rainfall exceeding $6\text{mm}\cdot\text{day}^{-1}$, hence it would be interesting to assess the impact of river iron supply
460 on the diazotrophs activity. In addition, the Vanuatu archipelago and Tonga are located on the ring fire, hence
461 hydrothermal sources could have a strong impact on dinitrogen fixation. These two iron sources are not yet
462 implemented in our configuration but may improve simulations of dinitrogen fixation in the south western tropical
463 Pacific region. Finally, our explicit simulation of dinitrogen fixation has proven to be higher by 25% (while still in the
464 lower end of estimations from observations) than the more commonly used implicit parameterization.

465

466 6. Appendix

467

468 *Trichodesmium* preferentially fixes di-nitrogen at temperature between 20-34°C (Breitbarth et al., 2008). The
469 temperature effect on the growth rate is modeled using a 4th order polynomial function (Ye et al., 2012):

470



$$L_T^{\text{Tri}} = \frac{2,32 \cdot 10^{-5} \times T^4 - 2,52 \cdot 10^{-3} \times T^3 + 9,75 \cdot 10^{-2} \times T^2 - 1,58 \times T + 9,12}{0,25}$$

471

472

473 where $0.25d^{-1}$ is the maximum observed growth rate (Breitbarth et al., 2008). Hence, at 17°C the growth rate is zero
 474 and maximum growth rate is reached at 27.5°C .

475

476 We first evaluate if nitrate+ammonium limitation occurs. In that case, dinitrogen fixation can occur and it is limited by
 477 phosphorus and iron. Phosphorus and iron limitations (L^P , L^{Fe}) are calculated in a 2-step process as follows:

478 First, we determine the most limiting nutrient:

$$L_{\text{Tri}}^{\text{Fe}} = \frac{(\theta^{\text{Fe}} - \theta_0^{\text{Fe}}) \theta_{\text{opt}}^{\text{Fe}}}{(\theta_{\text{opt}}^{\text{Fe}} - \theta_0^{\text{Fe}}) \theta^{\text{Fe}}} \quad \text{and} \quad L_{\text{Tri}}^{\text{P}} = \frac{(\theta^{\text{P}} - \theta_{\text{min}}^{\text{P}}) \theta_{\text{opt}}^{\text{P}}}{(\theta_{\text{opt}}^{\text{P}} - \theta_{\text{min}}^{\text{P}}) \theta^{\text{P}}}$$

479

480 where $\theta^{\text{Nutrients}}$ represents the nutrient quota for Fe and PO_4 (i.e. the ratio between iron and carbon concentrations in
 481 *Trichodesmium*, for instance).

482

$$483 \quad \theta_0^{\text{Fe}} = \theta_{\text{min}}^{\text{Fe}} - m$$

484 where m represents the maintenance iron, the intracellular Fe : C present in the cell at zero net growth rate (Kustka et
 485 al., 2003).

486

487 $\theta_{\text{min}}^{\text{Nut}}$ and $\theta_{\text{opt}}^{\text{Nut}}$ are constant, whereas θ varies with time. The minimum of $L_{\text{Tri}}^{\text{Fe}}$ and $L_{\text{Tri}}^{\text{P}}$ defines the limiting nutrient.

488 We then evaluate dinitrogen fixation as follows:

489

490 If iron is limiting :

491

$$492 \quad \mu_{\text{FixN}_2} = \mu_{\text{max}}^{\text{Tri}} L_I \frac{(\theta^{\text{Fe}} - \theta_0^{\text{Fe}}) \theta_{\text{opt}}^{\text{Fe}} - L_{\text{Tri}}^{\text{N}} (\theta_{\text{opt}}^{\text{Fe}} - \theta_0^{\text{Fe}}) \theta^{\text{Fe}}}{\alpha \mu_{\text{max}}^{\text{Tri}} \theta_{\text{opt}}^{\text{Fe}} + (\theta_{\text{opt}}^{\text{Fe}} - \theta_0^{\text{Fe}}) \theta^{\text{Fe}}}$$

493 μ_{FixN_2} is the growth rate of *Trichodesmium* sustained by dinitrogen fixation.

494

495 else if phosphorus is limiting :

$$496 \quad \mu_{\text{fixN}_2} = \mu_{\text{max}}^{\text{Tri}} L_I L_P^{\text{Tri}} - (\mu_{\text{NO}_3}^{\text{Tri}} + \mu_{\text{NH}_4}^{\text{Tri}})$$

497

498 which allows to recalculate the actual iron limitation for *Trichodesmium* as follows

$$499 \quad L_{\text{Tri}}^{\text{Fe}} = \frac{(\theta^{\text{Fe}} - \theta_0^{\text{Fe}} - \alpha \mu_{\text{fixN}_2}) \theta_{\text{opt}}^{\text{Fe}}}{(\theta_{\text{opt}}^{\text{Fe}} - \theta_0^{\text{Fe}}) \theta^{\text{Fe}}}; \alpha = \frac{1}{\beta}$$

500

501 β is the marginal use efficiency and equals the moles of additional carbon fixed per additional mole of intracellular iron
 502 per day (Raven, 1988; Sunda and Huntsman, 1997).

503



504 Such recalculation takes into account the fact that *Trichodesmium* need much higher cell Fe : C ratios to achieve the
 505 same growth rate as if growing on ammonium. That limitation is similar to the previous iron michaelis-menten-type
 506 limitation except that the growth rate is zero as long as the Fe : C ratio does not reach 14 μmol Fe : mol C (Kustka et al.,
 507 2003).

508

509 To summarize *Trichodesmium* growth rate:

510

$$511 \text{ If } \mu_{FixN_2} = 0 : \mu_{NO_3}^{Tri} = \mu_{max}^{Tri} L_I L_P^{Tri} \frac{L_{NO_3}^{Tri}}{L_{NO_3}^{Tri} + L_{NH_4}^{Tri}} \text{ and } \mu_{NH_4}^{Tri} = \mu_{max}^{Tri} L_I L_P^{Tri} \frac{L_{NH_4}^{Tri}}{L_{NO_3}^{Tri} + L_{NH_4}^{Tri}}$$

512

513

$$514 \text{ If } \mu_{FixN_2} > 0 : \mu_{NO_3}^{Tri} = \mu_{max}^{Tri} L_I L_{NO_3}^{Tri} \min(L_P^{Tri}, L_{Fe}^{Tri}) \text{ and } \mu_{NH_4}^{Tri} = \mu_{max}^{Tri} L_I L_{NH_4}^{Tri} \min(L_P^{Tri}, L_{Fe}^{Tri}),$$

515

516

517 L_P^{Tri} , L_N^{Tri} , L_{Fe}^{Tri} are respectively the *Trichodesmium* limiting function by phosphate, nitrogen and iron. L_I is the
 518 *Trichodesmium* limiting function by temperature and light.

519 μ_{max}^{Tri} is the *Trichodesmium* maximum growth rate,

520

521 $\mu_{NO_3}^{Tri}$ and $\mu_{NH_4}^{Tri}$ are respectively the new and regenerated productions.

522

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528

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Table 1 : Models parameters for Trichodemium and nanophytoplakton.

Parameters	Symbol	Name in code	Unity	Value
Initial slope P-I tricho	αI	pislope1	$(W \cdot m^{-2})^{-1} d^{-1}$	0.072
Initial slope P-I nano	αI	pislope	$(W \cdot m^{-2})^{-1} d^{-1}$	2.0
Microzoo preference for tricho	pItri	xpref2t	-	0.5
Microzoo preference for nano	pIP	xpref2p	-	1.0
Mesozoo preference for tricho	pItri	xpref1	-	0.3
Mesozoo preference for nano	pIP	xprefp	-	0.3
Tricho feeding threshold for mesozoo	Tithresh	xthresh2tri	mol C L ⁻¹	1.10 ⁻⁸
Nanophyto feeding threshold for mesozoo	Pithresh	xthresh2phy	mol C L ⁻¹	1.10 ⁻⁸
Tricho feeding threshold for microzoo	Tithresh	xthreshtri	mol C L ⁻¹	1.10 ⁻⁸
Nanophyto feeding threshold for microzoo	Pithresh	xthreshphy	mol C L ⁻¹	1.10 ⁻⁸
NO ₃ half saturation of tricho	KTriNO ₃	conctno3	mol N L ⁻¹	1.10 ⁻⁶
NO ₃ half saturation of nanophyto	KPNO ₃	concnno3	mol N L ⁻¹	1.10 ⁻⁶
NH ₄ half saturation of tricho	KTriNH ₄	conctnh4	mol N L ⁻¹	5.10 ⁻⁷
NH ₄ half saturation of nanophyto	KPNH ₄	concnnh4	mol N L ⁻¹	5.10 ⁻⁷
PO ₄ half saturation of tricho	KTriPO ₄	conctpo4	mol P L ⁻¹	5.10 ⁻⁶
PO ₄ half saturation of nanophyto	KPPO ₄	concnpo4	mol P L ⁻¹	5.10 ⁻⁶
Iron half saturation for tricho	KtriFe	conctfer	mol Fe L ⁻¹	1.10 ⁻⁹
Iron half saturation for nanophyto	KPFe	concnfer	mol Fe L ⁻¹	1.10 ⁻⁹
Minimum size criteria for tricho	Imax	xsizetri	mol C L ⁻¹	1.10 ⁻⁶
Minimum size criteria for nanophyto	Imax	xsizephy	mol C L ⁻¹	1.10 ⁻⁶
Optimal Fe quota for tricho	$\theta Fe, \text{Triopt}$	qtfelim	mol Fe (mol C) ⁻¹	7.10 ⁻⁶
Optimal Fe quota for nanophyto	$\theta Fe, \text{Iopt}$	qnfelim	mol Fe (mol C) ⁻¹	7.10 ⁻⁶
Minimum Chl/C in tricho	$\theta Chl, \text{Trimin}$	chlctm	mg Chl (mg C) ⁻¹	0.033
Minimum Chl/C in nanophyto	$\theta Chl, \text{Imin}$	chlenm	mg Chl (mg C) ⁻¹	0.033
Maximum Fe/C in tricho	$\theta Fe, \text{Trimax}$	fectm	mol Fe (mol C) ⁻¹	1.10 ⁻⁴
Maximum Fe/C in nanophyto	$\theta Fe, \text{Imax}$	fecnm	mol Fe (mol C) ⁻¹	4.10 ⁻⁵
Maximum N/C in tricho	$\theta N, \text{Trimax}$	qntlm	mol N (mol C) ⁻¹	1.45
Maximum N/C in nanophyto	$\theta N, \text{Imax}$	qnnlm	mol N (mol C) ⁻¹	1.45
Maximum P/C in tricho	$\theta P, \text{Trimax}$	qptlm	mol P (mol C) ⁻¹	2.44
Maximum P/C in nanophyto	$\theta P, \text{Imax}$	qpnlm	mol P (mol C) ⁻¹	2.44
Excretion ratio of tricho	rTri	excret1	d ⁻¹	0.05
Excretion ratio of nanophyto	rI	excret	d ⁻¹	0.05
Maintenance iron	m	zfixbasal	mol Fe (mol C) ⁻¹	1.4.10 ⁻⁵
Marginal use efficiency	β	qfec	day ⁻¹	1.4.10 ⁻⁴



Table 2 : List and description of the different experiments.

Name configuration	N ₂ fixation	Iron from sediment
TRI	explicit	yes
TRI_NoFeSed	explicit	no
N2_imp	implicit	yes
Wo_N2	no	yes

790 **Figures caption**

Fig. 1 : Annual mean concentrations in $\mu\text{mol L}^{-1}$: a) PO₄ data from the CARS b) PO₄ simulated by ROMS-PISCES model c) NO₃ data from the CARS d) NO₃ simulated by ROMS-PISCES model. On panels a and b the black contours show the annual mean of patterns of temperature preferendum from observations (a) and model (b). The red contours displays the 26°C isotherm for austral winter (plain) and austral summer (dash). On panels (c) and (d) the red boxes represent the LNL regions.

Fig. 2 : Left : Boxplots of the 0-150 averaged Iron (nmol Fe.L^{-1}) data (blue) and the equivalent for the model (red) colocalised with the observations in space. The coloured box represents the 25-75% of the distribution, the whiskers the 10-90% distribution. The line into the coloured box is the median.

Right : Iron concentrations (nmol Fe.L^{-1}) as observed (b) and as simulated by the model (c). Iron concentrations has been averaged over the top 150m of the ocean. Model values have been sampled at the same location, the same month, and the same depth as the data.

Fig. 3 : Top : Annual mean Chlorophyll concentration (in mg Chl.m^{-3}) in surface from (a) GLOBCOLOUR data (b) TRI simulation (c) TRI_imp simulation.

Bottom (d) : Annual mean Trichodesmium chlorophyll (in mg Chl.m^{-3}) in surface from TRI simulation

Fig. 4 : Nitrogen fixation rates ($\mu\text{mol N.m}^{-2}\text{d}^{-1}$) as observed (left) and as simulated by TRI simulation (right).

Top : Nitrogen fixation rates has been integrated over the top 150m of the ocean.

Bottom : Nitrogen fixation rates has been averaged over the top 30m of the ocean.

Model values have been sampled at the same location, the same month, and the same depth as the data.

Fig. 5 : a) Depth-integrated (0 to 125m) rates of nitrogen fixation ($\mu\text{mol N.m}^{-2}\text{d}^{-1}$) at ALOHA for the data (blue) and TRI simulation (red).

b) Depth-integrated (0 to 150m) rates of nitrogen fixation ($\mu\text{mol N.m}^{-2}\text{d}^{-1}$) in the south pacific (red box, Fig. 1c) for the data (blue) and TRI simulation (red). The green curve is the averaged of the seasonal cycle from TRI simulation at the data positions. Model values have been sampled at the same location, the same month, and the same depth as the data.

Fig. 6 : Relative contribution (in percentage) of *Trichodesmium* to primary production.

Fig. 7 : *Trichodesmium* biomass (mmol C.m^{-2}) for a) austral summer and b) austral winter, integrated over the top 100m of the ocean.

Fig. 8 : Seasonal cycle of limitation terms for *Trichodesmium* production in a) South Pacific and b) North Pacific. The right scale represents the total limitation.

Fig. 9 : Top : Minimum, mean and maximum in the South box (Fig 1c) for (a) iron concentration, (b) chlorophyll concentration in *Trichodesmium*.

Bottom : Annual mean iron concentration (shading ; in nmol Fe.L^{-1}) and current (vectors ; in m.s^{-1}) for c) TRI_NoFeSed simulation and d) TRI simulation. Annual mean Chlorophyll concentration in *Trichodesmium* (mg Chl.m^{-3}) for e) TRI_NoFeSed simulation and f) TRI simulation. The concentrations have been averaged over the top 100m of the ocean.

Fig. 10 : Percentage increase of primary production between the TRI simulation and Wo_N2 simulation (top) and N2_imp simulation (bottom); for total primary production (left) and primary production from Nanophyto. + diatoms (right).

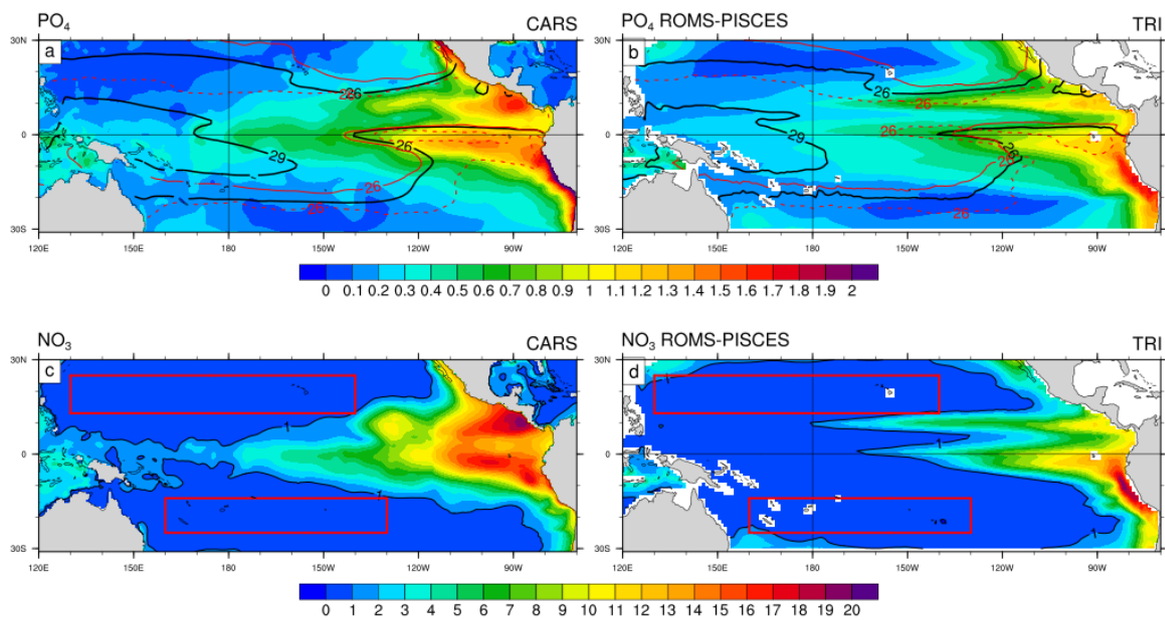


Figure 1

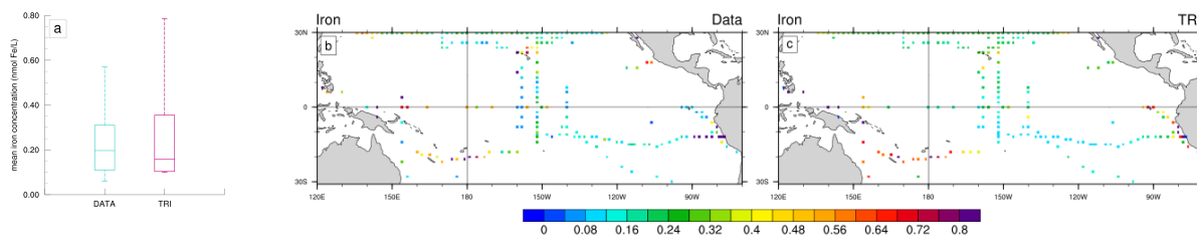


Figure 2

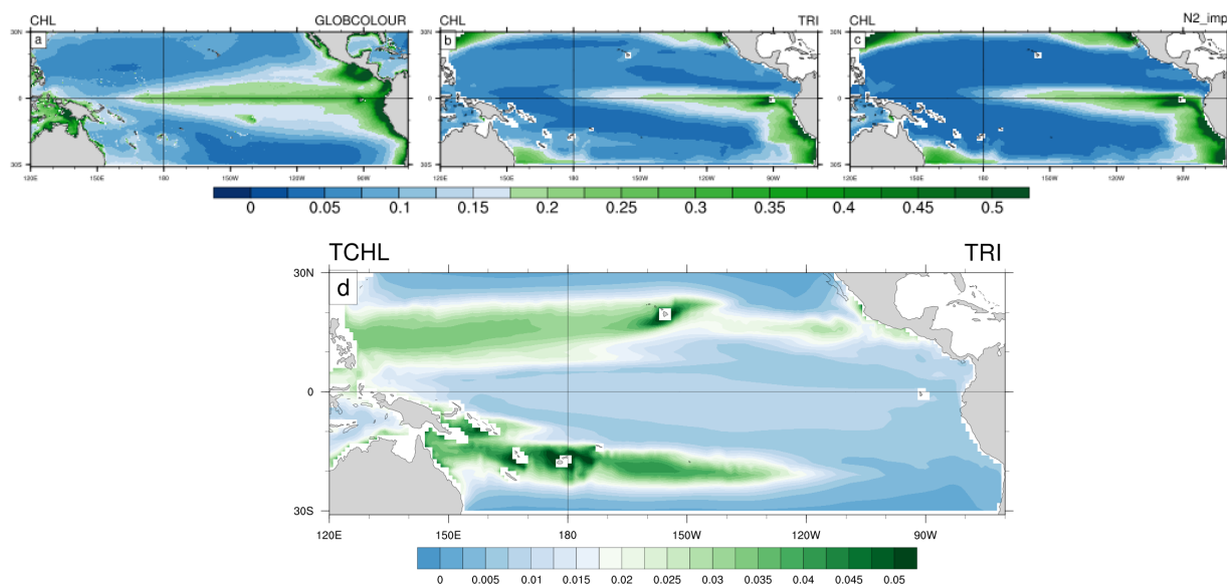


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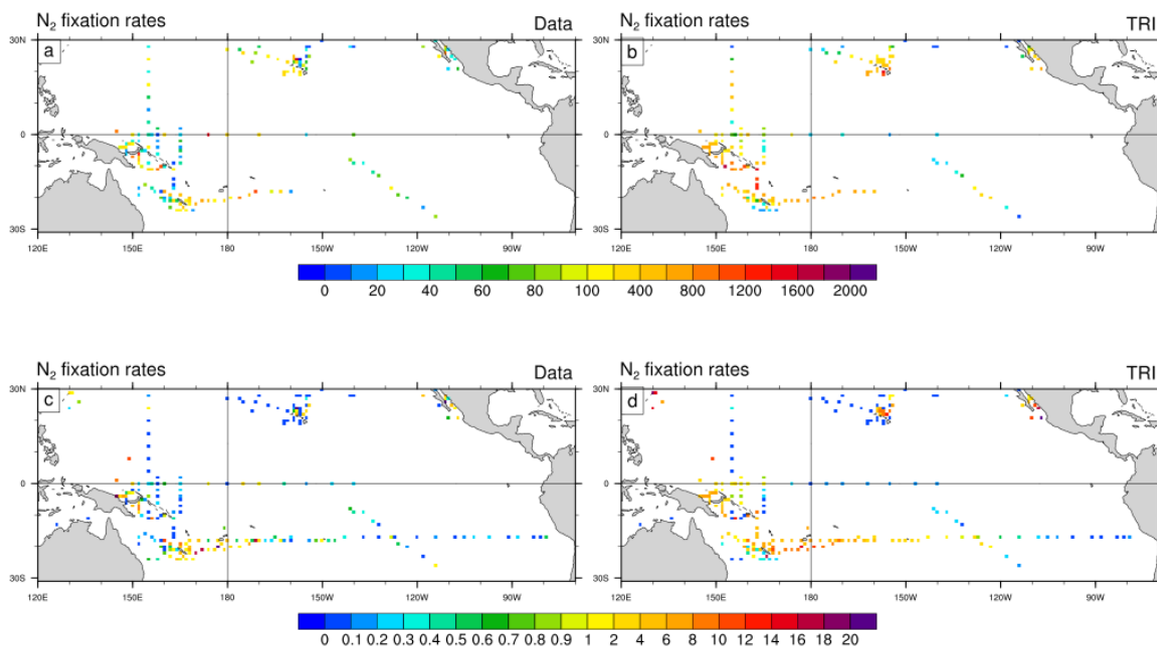


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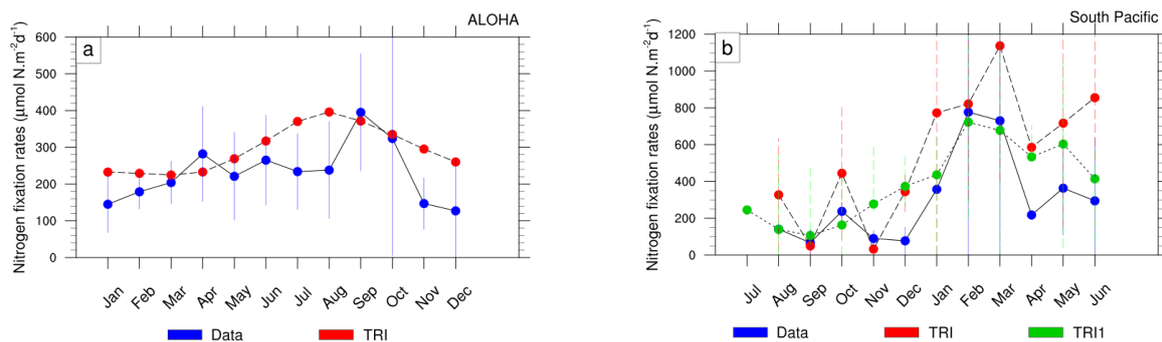


Figure 5

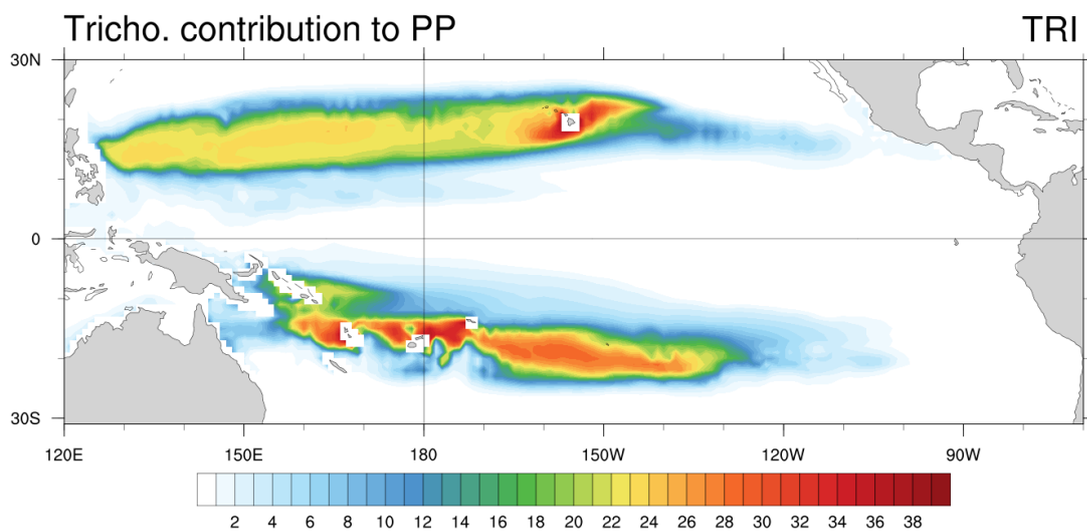


Figure 6

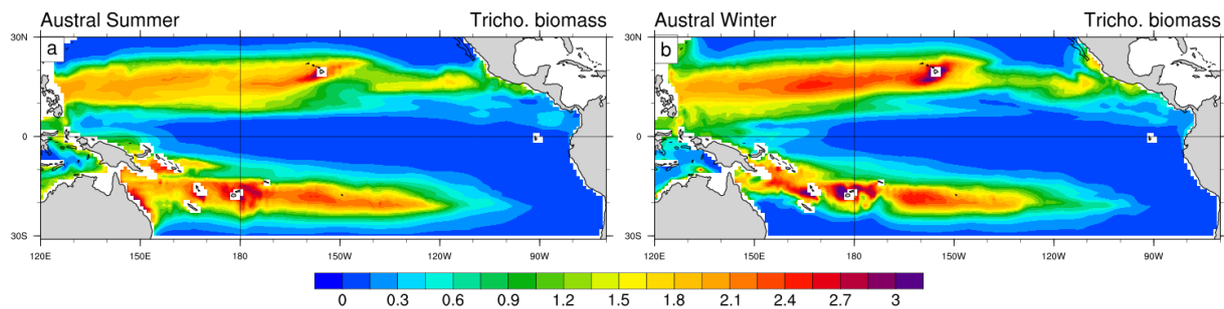


Figure 7

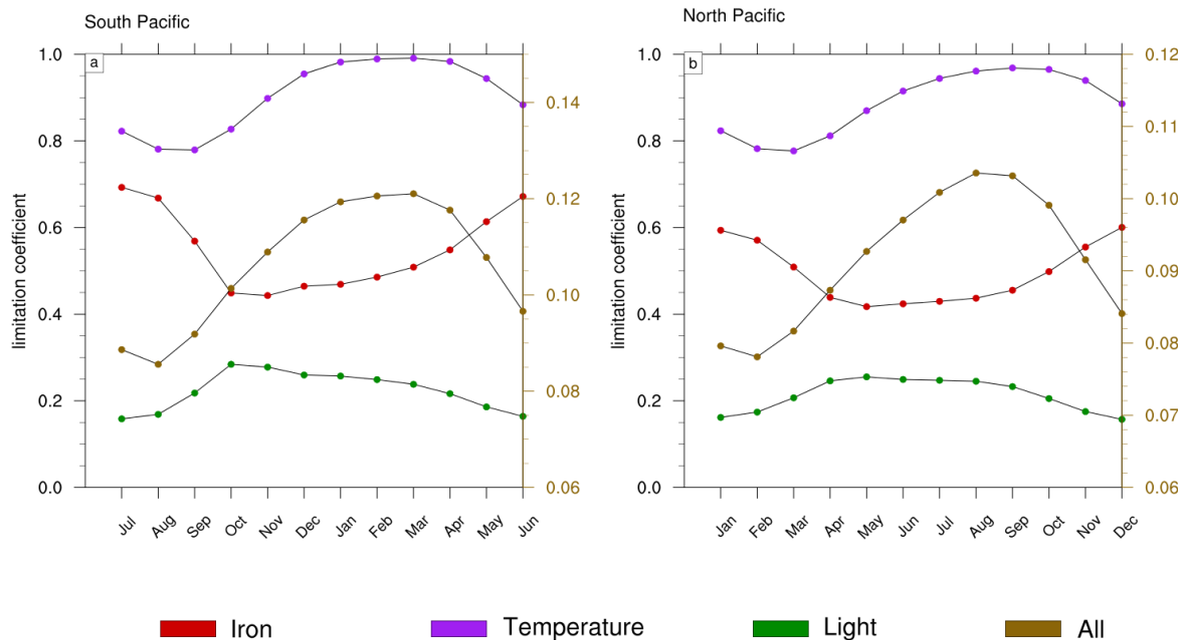


Figure 8

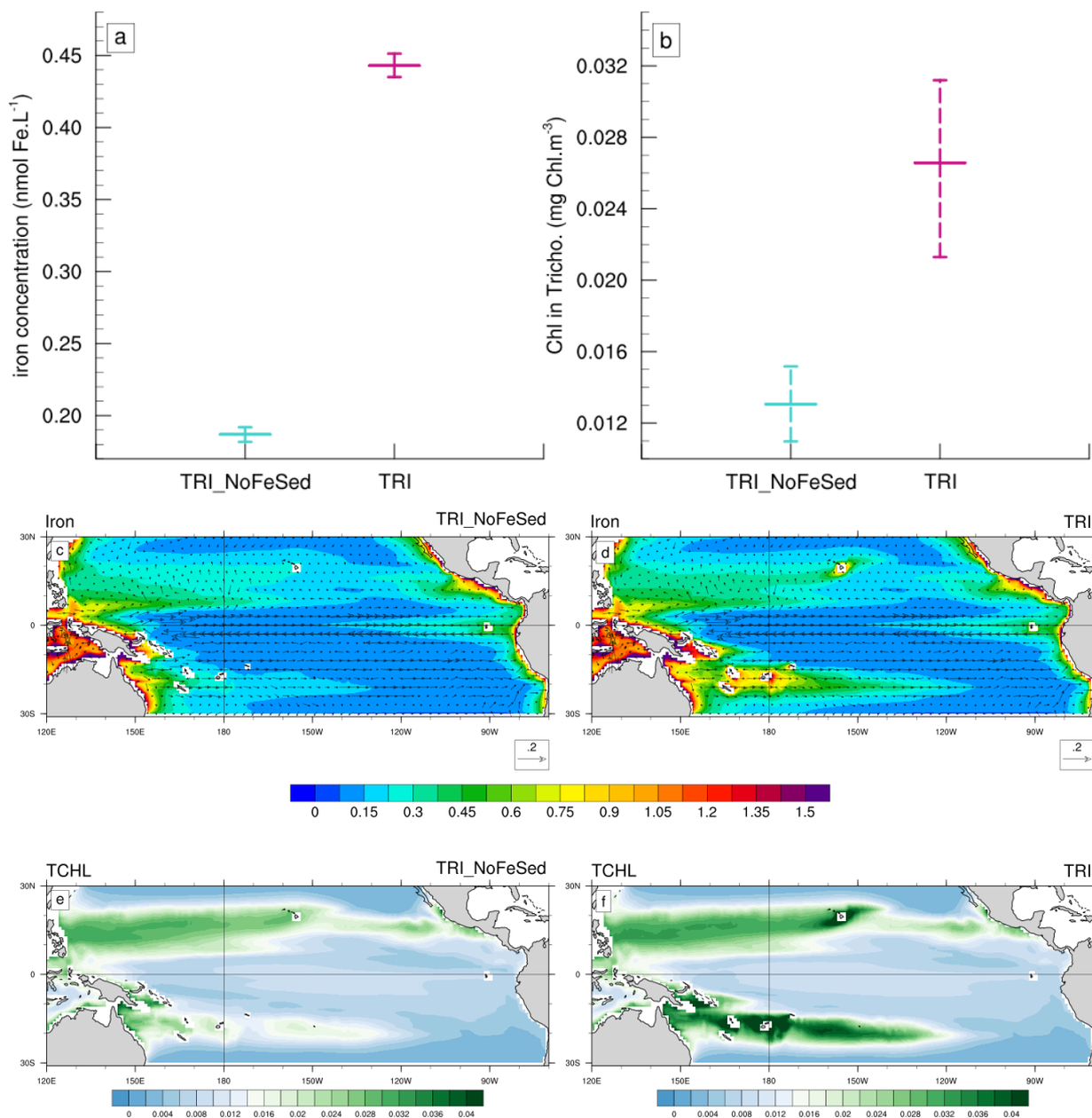


Figure 9

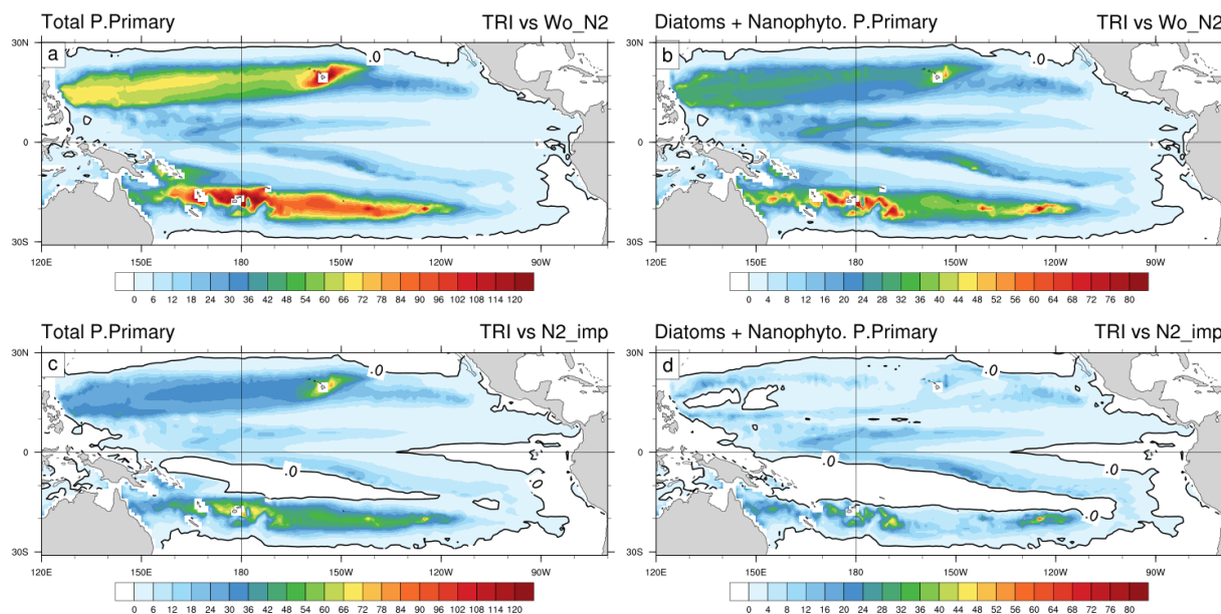


Figure 10