Review of "Projections of oceanic  $N_2O$  emissions in the 21st century using the IPSL Earth System Model" by Martinez-Rey et al.

The Authors addressed most of my comments in the revised version of the manuscript, and I'm generally satisfied with the changes and additions. In particular, the discussion of the potential shortcomings of the IPSL model, and the implications for the analysis, have been substantially expanded in the revised manuscript. The description of the  $N_2O$  production parameterizations, their origin and rationale have been also clarified, and references have been provided.

Among the conclusions of the paper, the suggested increase in subsurface inventories because of increased stratification (and despite reduced production) is novel and thought provoking. The decrease in interior N<sub>2</sub>O production as a response to decreased export and remineralization is also interesting, but more in line with our expectations. The role and importance of oxygen minimum zones and suboxic waters, and of the low-O<sub>2</sub> production pathway, is still unclear, and perhaps leads to the least robust results of the paper. However, this reflects the uncertainty that still exists in our understanding of low-O<sub>2</sub> processes, and the shortcomings of the 3D GCM utilized. These limitations have been now thoroughly acknowledged by the Authors.

I think that the manuscript will be a useful first reference for anyone interested in the future evolution of  $N_2O$  emission under climate change. It also points to processes that should be robust and of first order importance in Earth System Models (and presumably in the real world), and it suggests several aspects of the  $N_2O$  cycle where more work is needed.

We thank the referee for the second review of the manuscript. We acknowledge indeed the improvements in the manuscript on the above mentioned topics thanks to the comments from the referees.

I still have few concerns regarding the box model formulation. I realize this is not an essential part of the paper, although it is used to gain some insight to interpret the GCM model behavior. Also, my concerns might not change the final message. However, equations 2-3, with the parameters explained in the revised text and Table S1, seem wrong as a box model of the surface and deep ocean. My problems are:

To be dimensionally consistent, and obtain rates of change (as in the left hand sides), both the mixing coefficient and the gas exchange parameter should have units of (1/time). For example, a simple interpretation would hold if the mixing coefficient was a volume transport (m3/s) divided by the box volume (m3), and the gas exchange coefficient a piston velocity (m/s) multiplied by the box surface area (m2) and divided by the box volume (m3). As they are now, expressed as dimensional fractions, these coefficients do not allow to calculate time rates of changes, and the box model does not have a clear physical meaning. Once one realizes that the mixing terms represent a volume transport divided by the volume of the box, the mixing coefficients (v) should not be the same for the surface and deep box - a given transport of water will have much smaller effects in the deep box because of the much larger volume - unless here the surface and deep volumes are equal, which I guess the Authors could have (somewhat oddly) assumed (in which case the surface gas exchange should be quite small, because of the thickness of the surface box).

Similarly the gas exchange term is puzzling - even after correcting for the units of k (or pi) (which dimensionally should not be a fraction), the atmospheric concentrations are missing - it looks like the surface box just outgasses  $N_2O$  to an atmosphere with zero mixing ratio (not a big effect but physically odd).

I suggest that the authors check their model formulation, or at least provide a consistent physical interpretation and derivation of the equations.

The comment from the referee made us aware of how misleading is the box model as it stands. The main purpose of the box model was to synthesize the GCM model behaviour and basically to simplify the description of the main mechanisms leading to changes in future  $N_2O$  emissions. However, it is clear that it is not the case and therefore we have decided to withdraw the box model analysis from the manuscript. We think that the main messages of the paper are still well explained and comprehensible without the box model.

Technical:

1. 353: initialize (typo)

1. 385: decrease increase: awkward wording

1. 487: smaller instead of less

These proposed changes have been included in the latest version of the manuscript, which is attached below with corrections marked in blue.

- 3 J. Martinez-Rey<sup>1</sup>, L. Bopp<sup>2</sup>, M. Gehlen<sup>3</sup>, A. Tagliabue<sup>4</sup> and N. Gruber<sup>5</sup>.
- 4
- 5 <sup>1</sup> Laboratoire des Sciences du Climat et de l'Environnement, IPSL, CEA/CNRS/UVSQ,
- 6 Bat. 712 Orme des Merisiers, F-91191 CE Saclay, Gif-sur-Yvette, France.
- 7 jorge.martinez-rey@lsce.ipsl.fr
- 8
- 9 <sup>2</sup> Laboratoire des Sciences du Climat et de l'Environnement, IPSL, CEA/CNRS/UVSQ,
- 10 Bat. 712 Orme des Merisiers, F-91191 CE Saclay, Gif-sur-Yvette, France.
- 11 laurent.bopp@lsce.ipsl.fr
- 12
- 13 <sup>3</sup> Laboratoire des Sciences du Climat et de l'Environnement, IPSL, CEA/CNRS/UVSQ,
- 14 Bat. 712 Orme des Merisiers, F-91191 CE Saclay, Gif-sur-Yvette, France.
- 15 marion.gehlen@lsce.ipsl.fr
- 16
- 17 <sup>4</sup> School of Environmental Sciences, University of Liverpool, 4 Brownlow Street,
- 18 Liverpool L69 3GP, UK.
- 19 a.tagliabue@liverpool.ac.uk
- 20
- 21 <sup>5</sup> Environmental Physics, Institute of Biogeochemistry and Pollutant Dynamics, ETH,

- 22 CHN E31.2, Universitaetstrasse 16, 8092 Zürich, Switzerland.
- 23 nicolas.gruber@env.ethz.ch
- 24

27 The ocean is a substantial source of nitrous oxide (N2O) to the atmosphere, but little is 28 known on how this flux might change in the future. Here, we investigate the potential 29 evolution of marine N<sub>2</sub>O emissions in the 21st century in response to anthropogenic 30 climate change using the global ocean biogeochemical model NEMO-PISCES. Assuming 31 nitrification as the dominant N2O formation pathway, we implemented two different 32 parameterizations of N<sub>2</sub>O production which differ primarily at low oxygen (O<sub>2</sub>) 33 conditions. When forced with output from a climate model simulation run under the 34 business-as-usual high CO<sub>2</sub> concentration scenario (RCP8.5), our simulations suggest a 35 decrease of 4 to 12 % in N<sub>2</sub>O emissions from 2005 to 2100, i.e., a reduction from 4.03 / 36 3.71 to 3.54 / 3.56 TgN yr<sup>-1</sup> depending on the parameterization. The emissions decrease 37 strongly in the western basins of the Pacific and Atlantic oceans, while they tend to 38 increase above the Oxygen Minimum Zones (OMZs), i.e., in the Eastern Tropical Pacific 39 and in the northern Indian Ocean. The reduction in N2O emissions is caused on the one 40 hand by weakened nitrification as a consequence of reduced primary and export 41 production, and on the other hand by stronger vertical stratification, which reduces the 42 transport of N2O from the ocean interior to the ocean surface. The higher emissions over 43 the OMZ are linked to an expansion of these zones under global warming, which leads to 44 increased N<sub>2</sub>O production associated primarily with denitrification. While there are many uncertainties in the relative contribution and changes in the N<sub>2</sub>O production 45 46 pathways, the increasing storage seems unequivocal and determines largely the decrease in 47 N<sub>2</sub>O emissions in the future. From the perspective of a global climate system, the 48 averaged feedback strength associated with the projected decrease in oceanic N2O 49 emissions amounts to around -0.009 W m<sup>-2</sup>K<sup>-1</sup>, which is comparable to the potential 50 increase from terrestrial N2O sources. However, the assessment for a compensation 51 between the terrestrial and marine feedbacks calls for an improved representation of N<sub>2</sub>O 52 production terms in fully coupled next generation of Earth System Models.

53

- 54 1 Introduction
- 55

56 Nitrous oxide ( $N_2O$ ) is a gaseous compound responsible for two key feedback 57 mechanisms within the Earth's climate. First, it acts as a long-lived and powerful 58 greenhouse gas (Prather et al., 2012) ranking third in anthropogenic radiative forcing 59 after carbon dioxide ( $CO_2$ ) and methane ( $CH_4$ ) (Myrhe et al., 2013). Secondly, the 60 ozone ( $O_3$ ) layer depletion in the future might be driven mostly by  $N_2O$  after the drastic 61 reductions in CFCs emissions start to show their effect on stratospheric chlorine levels

reductions in CFCs emissions start to show their effect on stratospheric chlorine levels
(Ravishankara et al., 2009). The atmospheric concentration of N<sub>2</sub>O is determined by the

63 natural balance between sources from land and ocean and the destruction of  $N_2O$  in the

64 atmosphere largely by photolysis (Crutzen, 1970; Johnston, 1971). The natural sources

65 from land and ocean amount to ~6.6 and 3.8 TgN yr<sup>-1</sup>, respectively (Ciais et al., 2013).

66 Anthropogenic activities currently add an additional 6.7 TgN yr<sup>-1</sup> to the atmosphere,

 $\,67$   $\,$  which has caused atmospheric  $N_2O$  to increase by 18% since pre-industrial times (Ciais

68 et al., 2013), reaching 325 ppb in the year 2012 (NOAA ESRL Global Monitoring

69 Division, Boulder, Colorado, USA, http://esrl.noaa.gov/gmd/).

 $70 \qquad \text{Using a compilation of } 60,000 \text{ surface ocean observations of the partial pressure of } N_2O$ 

71 (pN<sub>2</sub>O), Nevison et al. (2004) computed a global ocean source of 4 TgN yr<sup>-1</sup>, with a

72 large range of uncertainty from 1.2 to 6.8 TgN yr<sup>-1</sup>. Model derived estimates also differ

73 widely, i.e., between 1.7 and 8 TgN yr<sup>-1</sup> (Nevison et al., 2003; Suntharalingam et al.,

74 2000). These large uncertainties are a consequence of too few observations and of poorly

75 known N<sub>2</sub>O formation mechanisms, reflecting a general lack of understanding of key

76 elements of the oceanic nitrogen cycle (Gruber and Galloway, 2008; Zehr and Ward,

2002), and of N<sub>2</sub>O in particular (e.g., Zamora et al., 2012, Bange et al., 2009 or Freing

78~ et al., 2012, among others). A limited number of interior ocean  $N_2O$  observations were

79 made available only recently (Bange et al., 2009), but they contain large temporal and

80 spatial gaps. Information on the rates of many important processes remains insufficient,

81 particularly in natural settings. There are only few studies from a limited number of

82 specific regions such as the Arabian Sea, Central and North Pacific, Black Sea, the

83 Bedford Basin and the Scheldt estuary, which can be used to derive and test model

84 parameterisations (Mantoura et al., 1993; Bange et al., 2000; Elkins et al., 1978; Farias et

al., 2007; Frame and Casciotti, 2010; Westley et al., 2006; Yoshida et al., 1989; Punshon

and Moore, 2004; De Wilde and De Bie, 2000).

87 N<sub>2</sub>O is formed in the ocean interior through two major pathways and consumed only in 88 oxygen minimum zones through denitrification (Zamora et al., 2012). The first 89 production pathway is associated with nitrification (conversion of ammonia, NH4\*, into 90 nitrate, NO3<sup>-</sup>), and occurs when dissolved O2 concentrations are above 20 µmol L<sup>-1</sup>. We 91 subsequently refer to this pathway as the high-O2 pathway. The second production 92 pathway is associated with a series of processes when  $O_2$  concentrations fall below ~5 93 µmol L-1 and involve a combination of nitrification and denitrification (hereinafter 94 referred to as low-O2 pathway) (Cohen and Gordon, 1978; Goreau et al., 1980; Elkins et 95 al., 1978). As nitrification is one of the processes involved in the aerobic remineralization 96 of organic matter, it occurs nearly everywhere in the global ocean with a global rate at 97 least one order of magnitude larger than the global rate of water column denitrification 98 (Gruber, 2008). A main reason is that denitrification in the water column is limited to 99 the OMZs, which occupy only a few percent of the total ocean volume (Bianchi et al., 100 2012). This is also the only place in the water column where N<sub>2</sub>O is being consumed. 101 The two production pathways have very different N2O yields, i.e., fractions of nitrogen-102 bearing products that are transformed to  $N_2O$ . For the high- $O_2$  pathway, the yield is 103 typically rather low, i.e., only about 1 in several hundred molecules of ammonium 104 escapes as N2O (Cohen and Gordon, 1979). In contrast, in the low-O2 pathway, and 105 particularly during denitrification, this fraction may go up to as high as 1:1, i.e., that all 106 nitrate is turned into  $N_2O$  (Tiedje, 1988). The relative contribution of the two pathways 107 to global N2O production is not well established. Sarmiento and Gruber (2006) 108 suggested that the two may be of equal importance, but more recent estimates suggest

111 Two strategies have been pursued in the development of parameterizations for  $N_2O$ 112 production in global biogeochemical models. The first approach builds on the 113 importance of the nitrification pathway and its close association with the aerobic 114 remineralization of organic matter. As a result the production of  $N_2O$  and the 115 consumption of  $O_2$  are closely tied to each other, leading to a strong correlation between 116 the concentration of  $N_2O$  and the apparent oxygen utilization (AOU). This has led to the 117 development of two sets of parameterizations, one based on concentrations, i.e., directly

118 as a function of AOU (Butler et al., 1989) and the other based on the rate of oxygen 119 utilization, i.e. OUR (Freing et al., 2009). Additional variables have been introduced to 120 allow for differences in the yield, i.e., the ratio of N<sub>2</sub>O produced over oxygen consumed, 121 such as temperature (Butler et al., 1989) or depth (Freing et al., 2009). In the second 122 approach, the formation of N<sub>2</sub>O is modeled more mechanistically, and tied to both 123 nitrification and denitrification by an O2 dependent yield (Suntharalingam and 124 Sarmiento, 2000; Nevison et al., 2003; Jin and Gruber, 2003). Since most models do not 125 include nitrification explicitly, the formation rate is actually coupled directly to the 126 remineralization of organic matter. Regardless of the employed strategy, all 127 parameterizations depend to first order on the amount of organic matter that is being 128 remineralized in the ocean interior, which is governed by the export of organic carbon to 129 depth. The dependence of N<sub>2</sub>O production on oxygen levels and on other parameters 130 such as temperature only acts at second order. This has important implications not only 131 for the modeling of the present-day distribution of N2O in the ocean, but also for the 132 sensitivity of marine N<sub>2</sub>O to future climate change.

133 Over this century, climate change will perturb marine N<sub>2</sub>O formation in multiple ways. 134 Changes in productivity will drive changes in the export of organic matter to the ocean 135 interior (Steinacher et al., 2010; Bopp et al., 2013) and hence affect the level of marine 136 nitrification. Ocean warming might change the rate of N2O production during 137 nitrification (Freing et al., 2012). Changes in carbonate chemistry (Bindoff et al., 2007) 138 might cause changes in the C:N ratio of the exported organic matter (Riebesell et al., 139 2007), altering not only the rates of nitrification, but also the ocean interior oxygen levels 140 (Gehlen et al., 2011). Finally, the expected general loss of oxygen (Keeling et al., 2010; 141 Cocco et al., 2012; Bopp et al., 2013) could substantially affect N<sub>2</sub>O production via both 142 nitrifier denitrification and classic denitrification.

143 Ocean biogeochemical models used for IPCC's 4<sup>th</sup> assessment report estimated a decrease

144 between 2% and 13% in primary production (PP) under the business-as-usual high CO<sub>2</sub>

145 concentration scenario A2 (Steinacher et al., 2010). A more recent multi-model analysis

146 based on the models used in IPCC's 5<sup>th</sup> assessment report also suggest a large reduction of

147 PP down to 18% by 2100 for the RCP8.5 scenario (Bopp et al., 2013). In these

simulations, the export of organic matter is projected to decrease between 6% and 18%

149 in 2100 (Bopp et al., 2013), with a spatially distinct pattern: in general, productivity and

150 export are projected to decrease at mid- to low-latitudes in all basins, while productivity 151 and export are projected to increase in the high-latitudes and in the South Pacific 152 subtropical gyre (Bopp et al., 2013). A wider spectrum of responses was reported 153 regarding changes in the ocean oxygen content. While all models simulate decreased 154 oxygen concentrations in response to anthropogenic climate change (by about 2 to 4% in 155 2100), and particularly in the mid-latitude thermocline regions, no agreement exists with 156 regard to the hypoxic regions, i.e., those having oxygen levels below 60 µmol L<sup>-1</sup> (Cocco 157 et al., 2012; Bopp et al., 2013). Some models project these regions to expand, while 158 others project a contraction. Even more divergence in the results exists for the suboxic 159 regions, i.e., those having  $O_2$  concentrations below 5 µmol L<sup>-1</sup> (Keeling et al., 2010; 160 Deutsch et al., 2011; Cocco et al., 2012; Bopp et al., 2013), although the trend for most 161 models is pointing towards an expansion. At the same time, practically none of the 162 models is able to correctly simulate the current distribution of oxygen in the OMZ (Bopp 163 et al., 2013). In summary, while it is clear that major changes in ocean biogeochemistry 164 are looming ahead (Gruber, 2011), with substantial impacts on the production and 165 emission of N2O, our ability to project these changes with confidence is limited. 166 In this study, we explore the implications of these future changes in ocean physics and 167 biogeochemistry on the marine N2O cycle, and make projections of the oceanic N2O

168 emissions from year 2005 to 2100 under the high CO<sub>2</sub> concentration scenario RCP8.5.
169 We analyze how changes in biogeochemical and physical processes such as net primary

170 production (NPP), export production and vertical stratification in this century translate

171  $\,$  into changes in oceanic  $N_2O$  emissions to the atmosphere. To this end, we use the

172 NEMO-PISCES ocean biogeochemical model, which we have augmented with two

 $173 \quad \ \ different \ N_2O \ parameterizations, \ permitting \ us \ to \ evaluate \ changes \ in \ the \ marine \ N_2O$ 

174 cycle at the process level, especially with regard to production pathways in high and low

175 oxygen regimes. We demonstrate that while future changes in the marine  $N_2O$  cycle will 176 be substantial, the net emissions of  $N_2O$  appear to change relatively little, i.e., they are

177 projected to decrease by about 10% in 2100.

178

- 179 2. Methodology
- 180
- 181 2.1 NEMO-PISCES Model

- 183 Future projections of the changes in the oceanic N<sub>2</sub>O cycle were performed using the 184 PISCES ocean biogeochemical model (Aumont and Bopp, 2006) in offline mode with 185 physical forcings derived from the IPSL-CM5A-LR coupled model (Dufresne et al., 186 2013). The horizontal resolution of NEMO ocean general circulation model is 2° x 2° cos 187 Ø (Ø being the latitude) with enhanced latitudinal resolution at the equator of  $0.5^{\circ}$ . 188 PISCES is a biogeochemical model with five nutrients (NO3, NH4, PO4, Si and Fe), two 189 phytoplankton groups (diatoms and nanophytoplankton), two zooplankton groups 190 (micro and mesozooplankton), and two non-living compartments (particulate and 191 dissolved organic matter). Phytoplankton growth is limited by nutrient availability and 192 light. Constant Redfield C:N:P ratios of 122:16:1 are assumed (Takahashi et al., 1985), 193 while all other ratios, i.e., those associated with chlorophyll, iron, and silicon (Chl:C, 194 Fe:C and Si:C) vary dynamically.
- 195

### 196 2.2 N<sub>2</sub>O parameterizations in PISCES

197

210

198 We implemented two different parameterizations of  $N_2O$  production in NEMO-PISCES. 199 The first one, adapted from Butler et al. (1989) follows the oxygen consumption 200 approach, with a temperature dependent modification of the  $N_2O$  yield (P.TEMP). The 201 second one is based on Jin and Gruber (2003) (P.OMZ), following the more mechanistic 202 approach, i.e., it considers the different processes occurring at differing oxygen

203 concentrations in a more explicit manner.

The P.TEMP parameterization assumes that the N<sub>2</sub>O production is tied to nitrification only with a yield that is at first order constant. This is implemented in the model by tying the N<sub>2</sub>O formation in a linear manner to O<sub>2</sub> consumption. A small temperature dependence is added to the yield to reflect the potential impact of temperature on metabolic rates. The production term of N<sub>2</sub>O, i.e.,  $J^{P.TEMP}(N_2O)$ , is then mathematically formulated as:

$$J^{P.TEMP}(N_2 O) = (\gamma + \theta T) J(O_2)_{consumption}$$
(1)

where 
$$\gamma$$
 is a background yield (0.53 x 10<sup>-4</sup> mol N<sub>2</sub>O/mol O<sub>2</sub> consumed),  $\theta$  is the

212 temperature dependency of  $\gamma$  (4.6 x 10<sup>-6</sup> mol N<sub>2</sub>O (mol O<sub>2</sub>)<sup>-1</sup> K<sup>-1</sup>), T is temperature (K),

- and  $J(O_2)_{consumption}$  is the sum of all biological O<sub>2</sub> consumption terms within the model. The same ratio between constants  $\gamma$  and  $\theta$  is used in the model as in the original formulation from Butler et al. (1989). Although this parameterization is very simple, a recent analysis of N<sub>2</sub>O observations supports such an essentially constant yield, even in the OMZ of the Eastern Tropical Pacific (Zamora et al., 2012).
- 218 The P.OMZ parameterization, formulated after Jin and Gruber (2003), assumes that the 219 overall yield consists of a constant background yield and an oxygen dependent yield. The 220 former is presumed to represent the  $N_2O$  production by nitrification, while the latter is 221 presumed to reflect the enhanced production of  $N_2O$  at low oxygen concentrations, in 222 part driven by denitrification, but possibly including nitrification as well. This 223 parameterization includes the consumption of  $N_2O$  in suboxic conditions. This gives:

$$J^{P.OMZ}(N_2O) = (\alpha + \beta f(O_2))J(O_2)_{consumption} - k N_2O$$
<sup>(2)</sup>

225 where  $\alpha$  is, as in Eq.(1), a background yield (0.9  $\cdot$  10<sup>-4</sup> mol N<sub>2</sub>O/mol O<sub>2</sub> consumed),  $\beta$  is 226 a yield parameter that scales the oxygen dependent function (6.2  $\cdot$  10<sup>-4</sup>),  $f(O_2)$  is a unitless 227 oxygen-dependent step-like modulating function, as suggested by laboratory experiments 228 (Goreau et al., 1980) (Fig. S1, Supplementary Material), and k is the 1<sup>st</sup> order rate 229 constant of  $N_2O$  consumption close to anoxia (zero otherwise). For k, we have adopted a 230 value of 0.138 yr<sup>-1</sup> following Bianchi et al. (2012) while we set the consumption regime 231 for  $O_2$  concentrations below 5 µmol L<sup>-1</sup>. The constant  $\alpha$  is in the same order of 232 magnitude as the one proposed by Jin and Gruber (2003), while  $\beta$  is two orders of 233 magnitude smaller. The use of the original value would result in a significant increase of 234 N<sub>2</sub>O production associated with OMZs and, hence, in a departure from the assumption 235 of dominant nitrification.

 $236 \qquad \text{The P.OMZ parameterization permits us the independent quantification of the $N_2O$}$ 

237 formation pathways associated with nitrification and those associated with low-oxygen

238 concentrations (nitrification/denitrification) and their evolution in time over the next

239 century. Specifically, we consider the source term  $\alpha J(O_2)_{consumption}$  as that associated with

240 the nitrification pathway, while we associated the source term  $\beta f(O_2) J(O_2)_{consumption}$  with

241 the low-oxygen processes (Fig. S2, Supplementary Material).

 $242 \qquad N_2O \ \text{production is inhibited by light in the model, and therefore $N_2O$ production in $$$ 

243 P.TEMP and P.OMZ parameterizations only occurs below a fixed depth of 100m.

- 244 We employ a standard bulk approach for simulating the loss of N<sub>2</sub>O to the atmosphere
- via gas exchange. We use the formulation of Wanninkhof et al. (1992) for estimating the
- 246 gas transfer velocity, adjusting the Schmidt number for N<sub>2</sub>O and using the solubility
- 247 constants of N<sub>2</sub>O given by Weiss and Price (1980). We assume a constant atmospheric
- 248 N<sub>2</sub>O concentration of 284 ppb in all simulations to explore future changes inherent to
- 249 ocean processes without feedbacks due to changes in the atmosphere.
- 250

251 2.3 Experimental design

252

253 NEMO-PISCES was first spun up during 3000 years using constant pre-industrial 254 dynamical forcings fields from IPSL-CM5A-LR (Dufresne et al., 2013) without 255 activating the N2O parameterizations. This spin-up phase was followed by a 150-yr long 256 simulation, forced by the same dynamical fields now with N2O production and N2O sea-257 to-air flux embedded. The N2O concentration at all grid points was prescribed initially to 258 20 nmol L<sup>-1</sup>, which is consistent with the MEMENTO database average value of 18 259 nmol L-1 below 1500m (Bange et al., 2009). During the 150-yr spin-up, we diagnosed 260 the total N<sub>2</sub>O production and N<sub>2</sub>O sea-to-air flux and adjusted the  $\alpha$ ,  $\beta$ ,  $\gamma$  and  $\theta$ 261 parameters in order to achieve a total N2O sea-to-air flux in the two parameterizations at 262 equilibrium close to 3.85 TgN yr<sup>-1</sup> (Ciais et al., 2013). In addition, the relative 263 contribution of the high-O2 pathway in the P.OMZ parameterization was set to 75% of 264 the total  $N_2O$  production based on Suntharalingam et al. (2000), where a sensitivity 265 model analysis on the relative contribution of high- and low-O2 production pathways 266 showed that a higher contribution of nitrification (75%) than denitrification (25%) 267 achieved the best model performance compared to the data product from Nevison et al. 268 (1995). P.TEMP can be considered as 100% nitrification, testing in this way the 269 hypothesis of nitrification as the dominant pathway of N<sub>2</sub>O production on a global scale. 270 Nitrification could contribute with up to 93% of the total production based on 271 estimations considering N<sub>2</sub>O production along with water mass transport (Freing et al., 272 2012).

- 273 Projections in NEMO-PISCES of historical (from 1851 to 2005) and future (from 2005
- 274 to 2100) simulated periods were done using dynamical forcing fields from IPSL-CM5A-
- 275 LR. These dynamical forcings were applied in an offline mode, i.e. monthly means of

276 temperature, velocity, wind speed or radiative flux were used to force NEMO-PISCES. 277 Future simulations used the business-as-usual high CO<sub>2</sub> concentration scenario (RCP8.5) 278 until year 2100. Century scale model drifts for all the biogeochemical variables presented, 279 including N2O sea-to-air flux, production and inventory, were removed using an 280 additional control simulation with IPSL-CM5A-LR pre-industrial dynamical forcing 281 fields from year 1851 to 2100. Despite the fact that primary production and the export 282 of organic matter to depth were stable in the control simulation, the air-sea N2O 283 emissions drifted (an increase of 5 to 12% in 200 yr depending on the parameterization) 284 due to the short spin-up phase (150 yrs) and to the choice of the initial conditions for 285 N<sub>2</sub>O concentrations.

286

287 3. Present-day oceanic N<sub>2</sub>O

288

289 3.1 Contemporary N<sub>2</sub>O fluxes

290

291 The model simulated air-sea N2O emissions show large spatial contrasts, with flux 292 densities varying by one order of magnitude, but with relatively small differences between 293 the two parameterizations (Fig. 1a and 1b). This is largely caused by our assumption that 294 the dominant contribution (75%) to the total N<sub>2</sub>O production in the P.OMZ 295 parameterization is the nitrification pathway, which is then not so different from the 296 P.TEMP parameterization, where it is 100%. As a result, the major part of  $N_2O$  is 297 produced in the subsurface via nitrification, contributing directly to imprint changes into 298 the sea-to-air N2O flux without a significant meridional transport (Suntharalingam and 299 Sarmiento, 2000).

300 Elevated N<sub>2</sub>O emission regions (> 50 mgN m<sup>-2</sup> yr<sup>-1</sup>) are found in the Equatorial and 301 Eastern Tropical Pacific, in the northern Indian ocean, in the northwestern Pacific, in the 302 North Atlantic and in the Agulhas Current. In contrast, low fluxes (< 10 mgN m<sup>-2</sup> yr<sup>-1</sup>) 303 are simulated in the Southern Ocean, Atlantic and Pacific subtropical gyres and southern 304 Indian Ocean. The large scale distribution of N<sub>2</sub>O fluxes is coherent with Nevison et al.

305 (2004) (Fig. 1c). This comes as a natural consequence of the relatively high contribution

306 of nitrification and hence hotspots of N<sub>2</sub>O emissions are associated with regions where

307 higher export of organic matter occurs in the model.

308 There are however several discrepancies between the model and the data product. At high 309 latitudes, the high N2O emissions observed in the North Pacific are not well represented 310 in our model, with a significant shift towards the western part of the Pacific basin, similar 311 to other modeling studies (e.g., Goldstein et al., 2003; Jin and Gruber, 2003). The OMZ 312 in the North Pacific, located at approximately 600m deep, is underestimated in the 313 model due to the deficient representation of the Meridional Overturning Circulation 314 (MOC) in the North Pacific in global ocean biogeochemical models, which in turn 315 might suppress low oxygenated areas and therefore one potential N<sub>2</sub>O source. 316 Discrepancies between model and observations also occur in the Southern Ocean, a 317 region whose role in global N<sub>2</sub>O fluxes remains debated due to the lack of observations 318 and the occurrence of potential artifacts due to interpolation techniques reflected in data 319 products such as that from Nevison et al., 2004. (e.g., Suntharalingam and Sarmiento, 320 2000, and Nevison et al, 2003). The model also overestimates N2O emissions in the 321 North Atlantic. The emphasis put on the nitrification pathway suggests that hotspots of 322 carbon export are at the origin of elevated concentrations of N2O in the subsurface. N2O 323 is quickly outgassed to the atmosphere, leading to such areas of high N2O emissions in 324 the model. 325 Model-data discrepancies can be seen as a function of latitude in Figure 1d. The modeled

 $N_2O$  flux maxima peak at around 40°S, i.e., around 10° north to that estimated by Nevison et al. (2004), although Southern Ocean data must be interpreted with caution. In the northern hemisphere the stripe in the North Pacific in not captured by the model,

329  $\,$  splitting the flux from the 45°N band into two peaks at 38°N and 55°N  $\,$ 

330

331 3.2 Contemporary N<sub>2</sub>O concentrations and the relationship to O<sub>2</sub>

332

The model results at present day were evaluated against the MEMENTO database (Bange et al., 2009), which contains about 25,000 measurements of co-located  $N_2O$  and dissolved  $O_2$  concentrations. Table 1 summarizes the standard deviation and correlation coefficients for P.TEMP and P.OMZ compared to MEMENTO. The standard deviation of the model output is very similar to MEMENTO, i.e., around 16 nmol L<sup>-1</sup> of  $N_2O$ . However, the correlation coefficients between the sampled data points from MEMENTO and P.TEMP / P.OMZ are 0.49 and 0.42 respectively. Largest

- 340 discrepancies are found mostly in the deep ocean and in the OMZs.
- 341 Figure 2 compares the global average vertical profile of the observed N<sub>2</sub>O against the 342 results from the two parameterisations. The in-situ observations show three characteristic 343 layers: the upper 100m layer with low (~10 nmol L-1) N<sub>2</sub>O concentration due to gas 344 exchange keeping N<sub>2</sub>O close to its saturation concentration, the mesopelagic layer, 345 between 100 and 1500m, where N<sub>2</sub>O is enriched via nitrification and denitrification in 346 the OMZs, and the deep ocean beyond 1500m, with a relatively constant concentration 347 of 18 nmol L-1 on average. Both parameterizations underestimate the N2O concentration 348 in the upper 100 meters, where most of the N2O is potentially outgassed to the 349 atmosphere. In the second layer, P.OMZ shows a fairly good agreement with the 350 observations in the 500 to 900m band, whereas P.TEMP is too low by ~10 nmol L<sup>-1</sup>. 351 Below 1500m, both parameterizations simulate too high N2O compared to the 352 observations. This may be caused by the lack or underestimation of a sink process in the 353 deep ocean, or by the too high concentrations used to initialize the model, which persist 354 due to the rather short spin-up time of only 150 yrs.
- 355 The analysis of the model simulated N<sub>2</sub>O concentrations as a function of model 356 simulated O<sub>2</sub> shows the differences between the two parameterizations more clearly (Fig. 357 3a and 3b). Such a plot allows us to assess the model performance with regard to N<sub>2</sub>O 358 (Jin and Gruber, 2003), without being subject to the strong potential biases introduced 359 by the model's deficiencies in simulating the distribution of O2. This is particularly 360 critical in the OMZs, where all models exhibit strong biases (Cocco et al., 2012; Bopp et 361 al., 2013) (see also Fig. 3c). P.TEMP (Fig. 3a) slightly overestimates N2O for dissolved 362 O2 concentrations above 100 µmol L-1, and does not fully reproduce either the high N2O 363 values in the OMZs or the N2O depletion when O2 is almost completely consumed. 364 P.OMZ (Figure 3b) overestimates the N<sub>2</sub>O concentration over the whole range of O<sub>2</sub>, 365 with particularly high values of N<sub>2</sub>O above 100 nmol L<sup>-1</sup> due to the exponential function 366 used in the OMZs. There, the observations suggest concentrations below 80 nmol L<sup>-1</sup> for 367 the same low O2 values, consistent with the linear trend observed for higher O2, which 368 seems to govern over most of the  $O_2$  spectrum, as suggested by Zamora et al. (2012). The 369 discrepancy at low O2 concentration may also stem from our choice of a too low N2O 370 consumption rate under essentially anoxic conditions. Finally, it should be considered 371 that most of the MEMENTO data points are from OMZs and therefore N2O
  - 12

372 measurements could be biased towards higher values than the actual open ocean average,

- 373 where our model performs better.
- 374
- 375 4. Future oceanic N<sub>2</sub>O
- 376
- 377 4.1 N<sub>2</sub>O sea-to-air flux
- 378

379 The global oceanic N<sub>2</sub>O emissions decrease relatively little over the next century (Fig. 4a) 380 between 4% and 12%. Namely, in P.TEMP, the emissions decrease by 0.15 TgN yr<sup>-1</sup> 381 from 3.71 TgN yr<sup>-1</sup> in 1985-2005 to 3.56 TgN yr<sup>-1</sup> in 2080-2100 and in P.OMZ, the 382 decrease is slightly larger at 12%, i.e., amounting to 0.49 Tg N yr<sup>-1</sup> from 4.03 to 3.54 383 TgN yr<sup>-1</sup>. Notable is also the presence of a negative trend in N<sub>2</sub>O emissions over the 20<sup>th</sup> 384 century, most pronounced in the P.OMZ parameterization. Considering the change over 385 the 20<sup>th</sup> and 21<sup>st</sup> centuries together, the model projects a decrease between 7 and 15%. 386 These relatively small global decreases mask more substantial changes at the regional scale,

387 with a mosaic of regions experiencing a substantial increase and regions experiencing a 388 substantial decrease (Fig. 4b and 4c). In both parameterizations, the oceanic N2O 389 emissions decrease in the northern and south western oceanic basins (e.g., the North 390 Atlantic and Arabian Sea), by up to 25 mgN m<sup>-2</sup>yr<sup>-1</sup>. In contrast, the fluxes are simulated 391 to increase in the Eastern Tropical Pacific and in the Bay of Bengal. For the Benguela 392 Upwelling System (BUS) and the North Atlantic a bi-modal pattern emerges in 2100. As 393 was the case for the present-day distribution of the N2O fluxes, the overall similarity 394 between the two parameterizations is a consequence of the dominance of the nitrification 395 (high-O<sub>2</sub>) pathway in both parameterizations. 396 Nevertheless there are two regions where more substantial differences between the two

parameterizations emerge: the region overlying the oceanic OMZ at the BUS and the
Southern Ocean. In particular, the P.TEMP parameterization projects a larger
enhancement of the flux than P.OMZ at the BUS, whereas the emissions in the Southern
Ocean are enhanced in the P.OMZ parameterization.

- 401
- 402 4.2 Drivers of changes in N<sub>2</sub>O emissions
- 403

Jorge 5/8/15 8:28 PM Deleted: the Jorge 5/8/15 8:29 PM Deleted: s Jorge 5/8/15 8:28 PM Deleted: increase Jorge 5/8/15 8:29 PM Deleted: to

408 The changes in N<sub>2</sub>O emissions may stem from a change in net N<sub>2</sub>O production, a change 409 in the transport of N<sub>2</sub>O from its location of production to the surface, or any 410 combination of the two, which includes also changes in N<sub>2</sub>O storage. Next we determine 411 the contribution of these mechanisms to the overall decrease in N<sub>2</sub>O emissions that our 412 model simulated for the  $21^{st}$  century.

413

414 4.2.1 Changes in N<sub>2</sub>O production

415

416 In both parameterizations, global N<sub>2</sub>O production is simulated to decrease over the 21st 417 century. The total N<sub>2</sub>O production in P.OMZ decreases by 0.41 TgN yr<sup>-1</sup> in 2080-2100 418 compared to the mean value over 1985-2005 (Fig. 5a). The parameterization P.OMZ allows to isolate the contributions of high- and low- $O_2$  and will be analysed in greater 419 420 detail in the following sections. N<sub>2</sub>O production via the high-O<sub>2</sub> pathway in P.OMZ 421 decreases in the same order than total production, by 0.35 TgN yr<sup>-1</sup> in 2080-2100 422 compared to present. The N2O production in the low-O2 regions remains almost 423 constant across the experiment. In P.TEMP parameterization, the reduction in N<sub>2</sub>O 424 production is much weaker than in P.OMZ due to the effect of the increasing 425 temperature. N<sub>2</sub>O production decreases by 0.07 TgN yr<sup>-1</sup> in 2080-2100 compared to 426 present (Fig. 5b).

427 The vast majority of the changes in the N<sub>2</sub>O production in the P.OMZ parameterization 428 is caused by the high-O<sub>2</sub> pathway with virtually no contribution from the low-O<sub>2</sub> 429 pathway (Fig. 5a). As the N<sub>2</sub>O production in P.OMZ parameterization is solely driven 430 by changes in the O<sub>2</sub> consumption (Eq. (2)), which in our model is directly linked to 431 export production, the dominance of this pathway implies that primary driver for the 432 future changes in N<sub>2</sub>O production in our model is the decrease in export of organic 433 matter (CEX). It was simulated to decrease by 0.97 PgC yr<sup>-1</sup> in 2100, and the high degree 434 of correspondence in the temporal evolution of export and N<sub>2</sub>O production in Fig. 5a 435 confirms this conclusion. 436 The close connection between N2O production associated with the high-O2 pathway and

- 437 changes in export production is also seen spatially (Fig. 5c), where the spatial pattern of 438 changes in export and changes in  $N_2O$  production are extremely highly correlated (shown
- 439 by stippling). Most of the small deviations are caused by lateral advection of organic

440 carbon, causing a spatial separation between changes in  $O_2$  consumption and changes in

441 organic matter export.

442 As there is an almost ubiquitous decrease of export in all of the major oceanic basins 443 except at high latitudes, N<sub>2</sub>O production decreases overall as well. Hotspots of reductions 444 exceeding -10 mgN m<sup>-2</sup>yr<sup>-1</sup> are found in the North Atlantic, the western Pacific and 445 Indian basins (Fig. 5c). The fewer places where export increases, are also the locations of 446 enhanced N2O production. For example, a moderate increase of 3 mgN m<sup>-2</sup> yr<sup>-1</sup> is 447 projected in the Southern Ocean, South Atlantic and Eastern Tropical Pacific. The 448 general pattern of export changes, i.e., decreases in lower latitudes, increase in higher 449 latitudes, is consistent generally with other model projection patterns (Bopp et al., 2013), 450 although there exist very strong model-to-model differences at the more regional scale.

451 Although the global contribution of the changes in the low-O2 N2O production is small, 452 this is the result of regionally compensating trends. In the model's OMZs, i.e., in the 453 Eastern Tropical Pacific and in the Bay of Bengal, a significant increase in N2O 454 production is simulated in these locations (Fig. 5d), with an increase of more than 15 455 mgN m<sup>-2</sup> yr<sup>-1</sup>. This increase is primarily driven by the expansion of the OMZs in our 456 model (shown by stippling), while changes in export contribute less. In effect, NEMO-457 PISCES projects a 20% increase in the hypoxic volume globally, from 10.2 to 12.3 x 10<sup>6</sup> 458 km<sup>3</sup>, and an increase in the suboxic volume from 1.1 to 1.6 x 10<sup>6</sup> km<sup>3</sup> in 2100 (Fig. 5e). 459 Elsewhere, the changes in the N2O production through the low-O2 pathway are 460 dominated by the changes in export, thus following the pattern of the changes seen in the 461 high-O<sub>2</sub> pathway. Overall these changes are negative, and happen to nearly completely 462 compensate the increase in production in the OMZs, resulting in the near constant 463 global N<sub>2</sub>O production by the low-O<sub>2</sub> production pathway up to year 2100.

464

465 4.2.2 Changes in storage of N<sub>2</sub>O

466

467 A steady increase in the  $N_2O$  inventory is observed from present to 2100. The pool of 468 oceanic  $N_2O$  down to 1500m, i.e., potentially outgassed to the atmosphere, increases by 469 8.9 TgN from 1985-2005 to year 2100 in P.OMZ, whereas P.TEMP is less sensitive to 470 changes with an increase of 4.0 TgN on the time period considered (Fig. 6a). The 471 inventory in the upper 1500m in P.OMZ is 237.0 TgN at present, while in P.TEMP in

the same depth band is 179.8 TgN. This means that the projected changes in the
inventory represent an increase of about 4% and 2% in P.OMZ and P.TEMP
respectively.

 $\label{eq:2.1} \mbox{This increase in storage of $N_2$O in the ocean interior shows an homogeneous pattern for}$ 

476 P.TEMP, with particular hotspots in the North Pacific, North Atlantic and the eastern

477 boundary currents in the Pacific (Fig. 6b). The spatial variability is more pronounced in

478 P.OMZ (Fig. 6c), related in part to the enhanced production associated with OMZs.

479 Most of the projected changes in storage are associated with shoaling of the mixed layer

482 to increasing sea surface temperatures associated with global warming (Sarmiento et al.,

- 483 2004).
- 484

 $485 \qquad 4.2.3 \ \text{Effects of the combined mechanisms on $N_2O$ emissions}$ 

486

487 The drivers of the future evolution of oceanic  $N_2O$  emissions emerge from the preceding 488 analysis. Firstly, a decrease in the high-O<sub>2</sub> production pathway driven by a reduced 489 organic matter remineralization reduces  $N_2O$  concentrations below the euphotic zone. 490 Secondly, the increased  $N_2O$  inventory at depth is caused by increased stratification and 491 therefore to a less efficient transport to the sea-to-air interface, leading to a <u>smaller  $N_2O$ </u> 492 flux.

493 The global changes in N<sub>2</sub>O flux, N<sub>2</sub>O production and N<sub>2</sub>O storage for P.OMZ are 494 presented in Fig. 7. Changes in N<sub>2</sub>O flux and N<sub>2</sub>O production are mostly of the same 495 sign in almost all of the oceanic regions in line with the assumption of nitrification begin 496 the dominant contribution to N2O production. Changes in N2O production in the 497 subsurface are translated into corresponding changes in N2O flux. There is only one 498 oceanic region (Sub-Polar Pacific) where this correlation does not occur. N<sub>2</sub>O inventory 499 increases in all of the oceanic regions. The increase in inventory is particularly 500 pronounced at low latitudes along the eastern boundary currents in the Equatorial and 501 Tropical Pacific, Indian Ocean and also in smaller quantities in the Atlantic Ocean. 502 Figure 7 shows how the decrease in N<sub>2</sub>O production and increase in N<sub>2</sub>O storage occurs 503 in all oceanic basins.

Jorge 5/8/15 8:33 PM Deleted: less

#### Jorge 5/8/15 8:04 PM

Deleted: The synergy among the driving mechanisms can be explored with a box model pursuing two objectives. First, to separate the effect of physical (i.e., increased stratification) and the biogeochemical (i.e., reduction of  $N_2 O$ production in the high-O2 regions) mechanisms on N2O emissions. In this way we can reproduce future projections assuming that the only mechanisms ruling the N2O dynamics in the future were those that we have proposed in our hypothesis, i.e., increased stratification and reduction of N2O production in high-O2 regions. Secondly, to explore a wider range of values for both mixing (i.e., degree of stratification) and efficiency of N2O production in high-O2 conditions. In the particular NEMO-PISCES model projection we have studied, changes in mixing and export are unique and can not be explored individually.

... [1]

529

527		
530	The state variables upon which representation of $\mathrm{N}_2\mathrm{O}$ in models rely, i.e., oxygen and	
531	export of carbon, are compared to the CMIP5 model ensemble to put our analysis in	
532	context of the current state-of-the-art model capabilities. We focus here our analysis on	
533	suboxic waters (O <sub>2</sub> < 5 $\mu mol~L^{-1}$ ) and on export production. Whereas CMIP5 models	
534	tend to have large volumes of $O_2$ concentrations in the suboxic regime, it is not the case	
535	for our NEMO-PISCES simulation, which clearly underestimates the volume of low-	
536	oxygen waters as compared to the oxygen corrected World Ocean Atlas 2005	
537	(WOA2005*) (Bianchi et al., 2012). The fact that NEMO-PISCES forced by IPSL-	
538	CM5A-LR is highly oxygenated is confirmed by Figure 8, where the histogram of the full	
539	O2 spectrum of WOA2005* and NEMO-PISCES is shown. The O2 distribution in the	Del
540	model shows a deficient representation of the OMZs, with higher concentrations than	
541	those from observations. The rest of the O2 spectrum is well represented in our model.	
542	The $O_2$ distribution in the model (Fig. 9) shows a deficient representation of the OMZs,	lor
543	with higher concentrations than those from observations in WOA2005* and the other	Del
544	CMIP5 models. NEMO-PISCES is therefore biased towards the high $\mathrm{O}_2$ production	
545	pathway of N2O due to the modeled O2 fields.	
546	When turning to the export of organic matter, NEMO-PISCES is close to the CMIP5	
547	average value of 6.9 PgC yr <sup>-1</sup> . The overall distribution of export is also very similar to the	
548	CMIP5 model mean and both show smaller values than those from the data-based	
549	estimate of 9.84 PgC yr <sup>-1</sup> from Dunne et al., 2007 (Fig. 9).	
550	The uncertainties derived from present and future model projections can be estimated	Del
551	using the spread in the CMIP5 model projection of export of organic matter and	
552	assuming a linear response between nitrification (or export) and $\mathrm{N_2O}$ production in the	
553	subsurface, which is assumed to be quickly outgassed to the atmosphere. In NEMO-	
554	PISCES, a decrease in 13% in export leads to a maximum decrease in $\mathrm{N}_2\mathrm{O}$ emissions of	
555	12% in the P.OMZ scenario. Based on results by Bopp et al. (2013), changes in export of	
556	carbon span -7% to -18% in the CMIP5 model ensemble at the end of the 21st century	
557	and for RCP8.5. The spread would propagate to a similar range in projected $\mathrm{N}_2\mathrm{O}$	

558  $\,$  emissions across the CMIP5 model ensemble. Applying these values to present  $N_2O$ 

Jorge 5/8/15 8:04 PM Deleted: 9

Jorge 5/8/15 8:04 PM Deleted: 10

Jorge 5/8/15 8:03 PM Deleted: 10

562 emissions of 3.6 TgN yr<sup>-1</sup>, uncertainties are then bracketed between -0.25 and -0.65 TgN
563 yr<sup>-1</sup>.

564 Regarding the low-O<sub>2</sub> pathway, a similar approach is not that straight forward. Zamora et 565 al., (2012) found that a linear relationship between AOU and N<sub>2</sub>O production might 566 occur even at the OMZ of the ETP. Zamora et al. (2012) acknowledged the fact that the 567 MEMENTO database includes N<sub>2</sub>O advected from other regions and that mixing could 568 play a relevant role, smoothing the fit between N2O and AOU from exponential to linear. 569 However, Zamora et al. (2012) quoting Frame and Casciotti (2010), suggested that 570 regions were an exponential relationship in N2O production is present might be rare, that 571 other non-exponential N<sub>2</sub>O production processes might occur and therefore the plot they 572 presented could describe the actual linear relationship between N2O production and 573 oxygen consumption. Based on this hypothesis, we could refer again to the linear 574 relationship suggested in the high-O2 and export scenario. However, in this case the 575 CMIP5 model projections of changes in the hypoxic and suboxic volumes differ 576 substantially. Most models project an expansion of the OMZs in the +2% to +16% range 577 in the suboxic volume ( $O_2 < 5 \mu mol L^{-1}$ ). There are, however, models that project a slight 578 reduction of 2%. Spatial variability of projections add to the spread between CMIP5 579 models. These discrepancies suggest that uncertainties from this spread must be 580 interpreted with caution when estimating potential future N<sub>2</sub>O emissions.

581 The use of  $O_2$  consumption as a proxy for the actual  $N_2O$  production plays therefore a 582 pivotal role in the uncertainties in N2O model estimations. Future model development 583 should aim at the implementation of mechanistic parameterizations of N2O production 584 based on nitrification and denitrification rates. Further, in order to determine accurate 585 O2 boundaries for both N2O production and N2O consumption at the core of OMZs 586 additional measurements and microbial experiments are needed. The contribution of the 587 high-O2 pathway that was considered in this model analysis might be a conservative 588 estimate. Freing et al. (2012) suggested that the high-O2 pathway could be responsible of 589 93% of the total N2O production. Assuming that changes in the N2O flux are mostly 590 driven by N<sub>2</sub>O production via nitrification, that would suggest a larger reduction in the 591 marine N2O emissions in the future. However, the mismatch between NEMO-PISCES 592 and the Nevison et al. (2004) spatial distribution of  $N_2O$  emissions in the western part of

the basins suggests that changes in the future might not be as big as those projected in the

594 model in such regions. Changes would be then distributed more homogeneously.

595 The model assumption neglecting N2O production in the upper 100m avoids one 596 important source of uncertainty in estimating global oceanic N2O fluxes. In case 597 nitrification occurs in the euphotic layer, our results would be facing a significant 598 uncertainty of at least ±25% in N2O emissions according to Zamora and Oschlies (2014) 599 analysis using the UVic Earth System Climate Model. Finally, Zamora et al. (2012) 600 observed a higher than expected  $N_2O$  consumption at the core of the OMZ in the 601 Eastern Tropical Pacific, occurring at an upper threshold of 10 µmol L<sup>-1</sup>. The 602 contribution of OMZs to total N2O production remains an open question. N2O 603 formation associated with OMZs might be counterbalanced by its own local 604 consumption, leading to the attenuation of the only increasing source of N2O 605 attributable to the projected future expansion of OMZs (Steinacher et al., 2010; Bopp et 606 al., 2013).

607 The combined effect of climate change and ocean acidification has not been analyzed in 608 this study. N<sub>2</sub>O production processes might be altered by the response of nitrification to 609 increasing levels of seawater pCO<sub>2</sub> (Huesemann et al., 2002; Beman et al. 2011). Beman 610 et al. (2011) reported a reduction in nitrification in response to decreasing pH. This 611 result suggests that N2O production might decrease beyond what we have estimated only 612 due to climate change. Conversely, negative changes in the ballast effect could potentially 613 reinforce nitrification at shallow depth in response to less efficient POC export to depth 614 and shallow remineralization (Gehlen et al., 2011). Regarding N<sub>2</sub>O formation via 615 denitrification, changes in seawater pH as a consequence of higher levels of CO2 might 616 not be substantial enough to change the N2O production efficiency, assuming a similar 617 response of marine denitrifiers as reported for denitrifying bacteria have in terrestrial 618 systems (Liu et al., 2010). Finally, the C:N ratio in export production (Riebesell et al., 619 2007) might increase in response to ocean acidification, potentially leading to a greater 620 expansion of OMZs than simulated here (Oschlies et al., 2008; Tagliabue et al., 2011), 621 and therefore to enhanced N<sub>2</sub>O production associated with the low-O<sub>2</sub> pathway. 622 Changes in atmospheric nitrogen deposition have not been considered in this study. It

623 has been suggested that due to anthropogenic activities the additional amount of reactive

- 624
- nitrogen in the ocean could fuel primary productivity and N<sub>2</sub>O production. Estimates are
- 625 however low, around 3-4% of the total oceanic emissions (Suntharalingam et al., 2012).

- 626 Longer simulation periods could reveal additional effects on N<sub>2</sub>O transport beyond 627 changes in upwelling or meridional transport of N2O in the subsurface (Suntharalingam 628 and Sarmiento, 2000) that have been observed in this transient simulation. Long-term 629 responses might include eventual ventilation of the N<sub>2</sub>O reservoir in the Southern Ocean, 630 highlighting the role of upwelling regions as an important source of N<sub>2</sub>O when longer 631 time periods are considered in model projections. Additional studies using other ocean 632 biogeochemical models might also yield alternative values using the same 633 parameterizations. N2O production is particularly sensitive to the distribution and 634 magnitude of export of organic matter and O2 fields defined in models.
- 635
- 636 6. Contribution of future N<sub>2</sub>O to climate feedbacks
- 637

638 Changes in the oceanic emissions of N2O to the atmosphere will have an impact on 639 atmospheric radiative forcing, with potential feedbacks on the climate system. Based on 640 the estimated 4 to 12% decrease in N2O sea-to-air flux over the 21st century under 641 RCP8.5, we estimated the feedback factor for these changes as defined by Xu-Ri et al. 642 (2012). Considering the reference value of the pre-industrial atmospheric N2O 643 concentration of 280 ppb in equilibrium, and its associated global N<sub>2</sub>O emissions of 11.8 644 TgN yr<sup>-1</sup>, we quantify the resulting changes in N<sub>2</sub>O concentration per degree for the two 645 projected emissions in 2100 using P.TEMP and P.OMZ. The model projects changes in 646 N<sub>2</sub>O emissions of -0.16 and -0.48 TgN yr<sup>-1</sup> respectively, whereas surface temperature is 647 assumed to increase globally by 3°C on average according to the physical forcing used in 648 our simulations. These results yield -0.05 and -0.16 TgN yr<sup>-1</sup> K<sup>-1</sup>, or alternatively -1.25 649 and -3.80 ppb K<sup>-1</sup> for P.TEMP and P.OMZ respectively. Using Joos et al. (2001) we 650 calculate the feedback factor in equilibrium for projected changes in emissions to be -651  $0.005 \text{ and } \text{-}0.014 \text{ W} \text{ m}^{\text{-}2}\text{K}^{\text{-}1} \text{ in P.TEMP and P.OMZ.}$ 652 Stocker et al. (2013) projected changes in terrestrial N<sub>2</sub>O emissions in 2100 using 653 transient model simulations leading to feedback strengths between +0.001 and +0.015 W

- 654 m<sup>-2</sup>K<sup>-1</sup>. Feedback strengths associated with the projected decrease of oceanic N<sub>2</sub>O
- 655 emissions are of the same order of magnitude as those attributable to changes in the
- 656 terrestrial sources of N<sub>2</sub>O, yet opposite in sign, suggesting a compensation of changes in
- 657 radiative forcing due to future increasing terrestrial N2O emissions. At this stage,

658 potential compensation between land and ocean emissions is to be taken with caution, as

659 it relies of a single model run with constant atmospheric N<sub>2</sub>O.

660

661 7. Conclusions

662

663 Our simulations suggest that anthropogenic climate change could lead to a global 664 decrease in oceanic  $N_2O$  emissions during the  $21^{st}$  century. This maximum projected 665 decrease of 12% in marine  $N_2O$  emissions for the business-as-usual high  $CO_2$  emissions 666 scenario would compensate for the estimated increase in  $N_2O$  fluxes from the terrestrial 667 biosphere in response to anthropogenic climate change (Stocker et al. 2013), so that the 668 climate- $N_2O$  feedback may be more or less neutral over the coming decades.

The main mechanisms contributing to the reduction of marine  $N_2O$  emissions are a decrease in  $N_2O$  production in high oxygenated waters as well as an increase in ocean vertical stratification that acts to decrease the transport of  $N_2O$  from the sub-surface to the surface ocean. Despite the decrease in both  $N_2O$  production and  $N_2O$  emissions, simulations suggest that the global marine  $N_2O$  inventory may increase from 2005 to 2100. This increase is explained by the reduced transport of  $N_2O$  from the production zones to the air-sea interface.

676 Differences between the two parameterizations used here are more related to 677 biogeochemistry rather than changes in ocean circulation. Despite sharing the high-O2 678 N<sub>2</sub>O production pathway, leading to a decrease in N<sub>2</sub>O emissions in both cases, the role 679 of warming in P.TEMP or higher N<sub>2</sub>O yields at low-O<sub>2</sub> concentrations in P.OMZ 680 translate into notable differences in the evolution of the two production pathways. 681 However, the dominant effect of changes in stratification in both parameterizations 682 drives ultimately the homogeneous response of the parameterizations considered in 683 model projections in the next century.

684 The  $N_2O$  production pathways demand however a better understanding in order to 685 enable an improved representation of processes in models. At a first order, the efficiencies 686 of the production processes in response to higher temperatures or increased seawater 687 pCO<sub>2</sub> are required. Second order effects such as changes in the O<sub>2</sub> boundaries at which 688 nitrification and denitrification occur must be also taken into account. In the absence of 689 process-based parameterizations, N<sub>2</sub>O production parameterizations will still rely on

- 690 export of organic carbon and oxygen levels. Both need to be improved in global
- 691 biogeochemical models.
- 692 The same combination of mechanisms (i.e., change in export production and ocean
- 693 stratification) have been identified as drivers of changes in oceanic N2O emissions during
- 694 the Younger Dryas by Goldstein et al. (2003). The N<sub>2</sub>O flux decreased, while the N<sub>2</sub>O
- 695 reservoir was fueled by longer residence times of  $N_2O$  caused by increased stratification.
- 696 Other studies point towards changes in the N<sub>2</sub>O production at the OMZs as the main
- 697 reason for variations in N<sub>2</sub>O observed in the past (Suthhof et al., 2001). Whether these
- 698 mechanisms are plausible drivers of changes beyond year 2100 remains an open question
- 699 that needs to be addressed with longer simulations.
- 700

- 701 8. Acknowledgements
- 702

703  $\,$  We thank Cynthia Nevison for providing us the  $N_2O$  sea-to-air flux dataset. We thank

704 Annette Kock and Herman Bange for the availability of the MEMENTO database

705 (https://memento.geomar.de). We thank Christian Ethé for help analyzing PISCES

706 model drift. Comments by Parvadha Suntharalingam and three anonymous reviewers

707 improved significantly this manuscript. Nicolas Gruber acknowledges the support of

708 ETH Zürich. This work has been supported by the European Union via the Greencycles

709 II FP7-PEOPLE-ITN-2008, number 238366.

- 710 7. References
- 711
- 712 Aumont, O., and Bopp, L.: Globalizing results from ocean in situ iron fertilization studies,
- 713 Global Biogeochemical Cycles, GB2017, 20, 10.1029/2005gb002591, 2006.
- 714 Bange, H. W., Rixen, T., Johansen, A. M., Siefert, R. L., Ramesh, R., Ittekkot, V., Hoffmann,
- 715 M. R., and Andreae, M. O.: A revised nitrogen budget for the Arabian Sea, Global
- 716 Biogeochemical Cycles, 14, 1283-1297, 10.1029/1999gb001228, 2000.
- 717 Bange, H. W., Bell, T. G., Cornejo, M., Freing, A., Uher, G., Upstill-Goddard, R. C., and
- 718 Zhang, G.: MEMENTO: a proposal to develop a database of marine nitrous oxide and methane
- 719 measurements, Environmental Chemistry, 6, 195-197, 10.1071/en09033, 2009.
- 720 Beman, J. M., Chow, C.-E., King, A. L., Feng, Y., Fuhrman, J. A., Andersson, A., Bates, N. R.,
- 721 Popp, B. N., and Hutchins, D. A.: Global declines in oceanic nitrification rates as a consequence
- 722 of ocean acidification, Proceedings of the National Academy of Sciences of the United States of
- 723 America, 108, 208-213, 10.1073/pnas.1011053108, 2011.
- 724 Bianchi, D., Dunne, J. P., Sarmiento, J. L., and Galbraith, E. D.: Data-based estimates of
- suboxia, denitrification, and N2O production in the ocean and their sensitivities to dissolved O-
- 726 2, Global Biogeochemical Cycles, 26, GB2009, 10.1029/2011gb004209, 2012.
- 727 Bindoff, N., Willebrand, J., Artale, V., Cazenave, A., Gregory, J., Gulev, S., Hanawa, K.,
- 728 Le Quere, C., Levitus, S., Norjiri, Y., Shum, C., Talley, L., and Unnikrishnan, A.:
- 729 Observations: Oceanic Climate Change and Sea Level, In Climate Change 2007: The
- 730 Physical Science Basis.Contribution of Working Group I to the Fourth Assessment
- 731 Report of the Intergovernmental Panel on Climate Change, 2007.
- 732 Bopp, L., Resplandy, L., Orr, J. C., Doney, S. C., Dunne, J. P., Gehlen, M., Halloran, P.,
- 733 Heinze, C., Ilyina, T., Seferian, R., Tjiputra, J., and Vichi, M.: Multiple stressors of ocean
- rade cosystems in the 21st century: projections with CMIP5 models, Biogeosciences, 10, 6225-6245,
- 735 10.5194/bg-10-6225-2013, 2013.
- 736 Butler, J. H., Elkins, J. W., Thompson, T. M., and Egan, K. B.: Tropospheric and dissolved
- 737 N<sub>2</sub>O of the west pacific and east-indian oceans during the el-nino southern oscillation event of
- 738 1987, Journal of Geophysical Research-Atmospheres, 94, 14865-14877,
- 739 10.1029/JD094iD12p14865, 1989.
- 740 Ciais, P., Sabine, C., Bala, G., Bopp, L., Brovkin, V., Canadell, J., Chhabra, A., DeFries,
- 741 R., Galloway, J., Heimann, M., Jones, C., Le Quéré, C., Myneni, RB., Piao, S. and

- 742 Thornton, P.: Carbon and Other Biogeochemical Cycles. In: Climate Change 2013: The
- 743 Physical Science Basis. Contribution of Working Group I to the Fifth Assessment Report
- of the Intergovernmental Panel on Climate Change, 2013.
- 745 Cocco, V., Joos, F., Steinacher, M., Froelicher, T. L., Bopp, L., Dunne, J., Gehlen, M., Heinze,
- 746 C., Orr, J., Oschlies, A., Schneider, B., Segschneider, J., and Tjiputra, J.: Oxygen and indicators
- of stress for marine life in multi-model global warming projections, Biogeosciences, 10, 1849-
- 748 1868, 10.5194/bg-10-1849-2013, 2013.
- 749 Cohen, Y., and Gordon, L. I.: Nitrous-oxide in oxygen minimum of eastern tropical north
- 750 pacific evidence for its consumption during denitrification and possible mechanisms for its
- 751 production, Deep-Sea Research, 25, 509-524, 10.1016/0146-6291(78)90640-9, 1978.
- 752 Cohen, Y., and Gordon, L. I.: Nitrous-oxide production in the ocean, Journal of Geophysical
- 753 Research-Oceans and Atmospheres, 84, 347-353, 10.1029/JC084iC01p00347, 1979.
- 754 Crutzen, P. J.: Influence of nitrogen oxides on atmospheric ozone content, Quarterly Journal of
- 755 the Royal Meteorological Society, 96, 320-326, 10.1002/qj.49709640815, 1970.
- 756 de Wilde, H. P. J., and de Bie, M. J. M.: Nitrous oxide in the Schelde estuary: production by
- nitrification and emission to the atmosphere, Marine Chemistry, 69, 203-216, 10.1016/s0304-
- 758 4203(99)00106-1, 2000.
- 759 Deutsch, C., Brix, H., Ito, T., Frenzel, H., and Thompson, L.: Climate-Forced Variability of
- 760 Ocean Hypoxia, Science, 333, 336-339, 10.1126/science.1202422, 2011.
- 761 Dufresne, J. L., Foujols, M. A., Denvil, S., Caubel, A., Marti, O., Aumont, O., Balkanski, Y.,
- 762 Bekki, S., Bellenger, H., Benshila, R., Bony, S., Bopp, L., Braconnot, P., Brockmann, P.,
- 763 Cadule, P., Cheruy, F., Codron, F., Cozic, A., Cugnet, D., de Noblet, N., Duvel, J. P., Ethe, C.,
- 764 Fairhead, L., Fichefet, T., Flavoni, S., Friedlingstein, P., Grandpeix, J. Y., Guez, L., Guilyardi,
- 765 E., Hauglustaine, D., Hourdin, F., Idelkadi, A., Ghattas, J., Joussaume, S., Kageyama, M.,
- 766 Krinner, G., Labetoulle, S., Lahellec, A., Lefebvre, M. P., Lefevre, F., Levy, C., Li, Z. X., Lloyd,
- 767 J., Lott, F., Madec, G., Mancip, M., Marchand, M., Masson, S., Meurdesoif, Y., Mignot, J.,
- 768 Musat, I., Parouty, S., Polcher, J., Rio, C., Schulz, M., Swingedouw, D., Szopa, S., Talandier,
- 769 C., Terray, P., Viovy, N., and Vuichard, N.: Climate change projections using the IPSL-CM5
- Earth System Model: from CMIP3 to CMIP5, Climate Dynamics, 40, 2123-2165,
- 771 10.1007/s00382-012-1636-1, 2013.
- 772 Dunne, J. P., Sarmiento, J. L., and Gnanadesikan, A.: A synthesis of global particle export
- from the surface ocean and cycling through the ocean interior and on the seafloor, Global

- 774 Biogeochemical Cycles, 21, GB4006,10.1029/2006gb002907, 2007.
- 775 Elkins, J. W., Wofsy, S. C., McElroy, M. B., Kolb, C. E., and Kaplan, W. A.: Aquatic sources
- 776 and sinks for nitrous-oxide, Nature, 275, 602-606, 10.1038/275602a0, 1978.
- 777 Farias, L., Paulmier, A., and Gallegos, M.: Nitrous oxide and N-nutrient cycling in the
- 778 oxygen minimum zone off northern Chile, Deep-Sea Research Part I-Oceanographic
- 779 Research Papers, 54, 164-180, 10.1016/j.dsr.2006.11.003, 2007.
- 780 Fletcher, S. E. M., Gruber, N., Jacobson, A. R., Gloor, M., Doney, S. C., Dutkiewicz, S.,
- 781 Gerber, M., Follows, M., Joos, F., Lindsay, K., Menemenlis, D., Mouchet, A., Muller, S. A.,
- and Sarmiento, J. L.: Inverse estimates of the oceanic sources and sinks of natural CO2 and
- the implied oceanic carbon transport, Global Biogeochemical Cycles, 21, GB1010,
- 784 10.1029/2006gb002751, 2007.
- 785 Frame, C. H., and Casciotti, K. L.: Biogeochemical controls and isotopic signatures of nitrous
- 786 oxide production by a marine ammonia-oxidizing bacterium, Biogeosciences, 7, 2695-2709,
- 787 10.5194/bg-7-2695-2010, 2010.
- 788 Freing, A., Wallace, D. W. R., Tanhua, T., Walter, S., and Bange, H. W.: North Atlantic
- 789 production of nitrous oxide in the context of changing atmospheric levels, Global
- 790 Biogeochemical Cycles, 23, GB4015, 10.1029/2009gb003472, 2009.
- 791 Freing, A., Wallace, D. W. R., and Bange, H. W.: Global oceanic production of nitrous oxide,
- Philosophical Transactions of the Royal Society B-Biological Sciences, 367, 1245-1255,
- 793 10.1098/rstb.2011.0360, 2012.
- 794 Gehlen, M., Gruber, N., Gangstø, R., Bopp, L., and Oschlies, A.: Biogeochemical consequences
- of ocean acidification and feedbacks to the earth system. Ocean acidification: 230-248, 2011.
- 796 Goldstein, B., Joos, F., and Stocker, T. F.: A modeling study of oceanic nitrous oxide during the
- 797 Younger Dryas cold period, Geophysical Research Letters, 30, 1092, 10.1029/2002gl016418,
- 798 2003.
- 799 Goreau, T. J., Kaplan, W. A., Wofsy, S. C., McElroy, M. B., Valois, F. W., and Watson, S. W.:
- 800 Production of NO<sub>2</sub><sup>-</sup> and N<sub>2</sub>O by nitrifying bacteria at reduced concentrations of oxygen, Applied
- and Environmental Microbiology, 40, 526-532, 1980.
- 802 Gruber, N., and Galloway, J. N.: An Earth-system perspective of the global nitrogen cycle,
- 803 Nature, 451, 293-296, 10.1038/nature06592, 2008.
- 804 Gruber, N.: Warming up, turning sour, losing breath: ocean biogeochemistry under global
- 805 change, Philosophical Transactions of the Royal Society a-Mathematical Physical and
- 806 Engineering Sciences, 369, 1980-1996, 10.1098/rsta.2011.0003, 2011.

- 807 Gruber, N.: The marine nitrogen cycle: Overview of distributions and processes. In:
- 808 Nitrogen in the marine environment, 2nd edition, 1-50, 2008.
- 809 Huesemann, M. H., Skillman, A. D., and Crecelius, E. A.: The inhibition of marine nitrification
- 810 by ocean disposal of carbon dioxide, Marine Pollution Bulletin, 44, 142-148, 10.1016/s0025-
- 811 326x(01)00194-1, 2002.
- 812 Jin, X., and Gruber, N.: Offsetting the radiative benefit of ocean iron fertilization by enhancing
- 813 N<sub>2</sub>O emissions, Geophysical Research Letters, 30, 2249, 10.1029/2003gl018458, 2003.
- 814 Johnston, H.: Reduction of stratospheric ozone by nitrogen oxide catalysts from supersonic
- 815 transport exhaust, Science, 173, 517-522, 10.1126/science.173.3996.517, 1971.
- 816 Joos, F., Prentice, I. C., Sitch, S., Meyer, R., Hooss, G., Plattner, G. K., Gerber, S., and
- 817 Hasselmann, K.: Global warming feedbacks on terrestrial carbon uptake under the
- 818 Intergovernmental Panel on Climate Change (IPCC) emission scenarios, Global Biogeochemical
- 819 Cycles, 15, 891-907, 10.1029/2000gb001375, 2001.
- 820 Keeling, R. F., Koertzinger, A., and Gruber, N.: Ocean Deoxygenation in a Warming World,
- 821 Annual Review of Marine Science, 2, 199-229, 10.1146/annurev.marine.010908.163855, 2010.
- 822 Liu, B., Morkved, P. T., Frostegard, A., and Bakken, L. R.: Denitrification gene pools,
- 823 transcription and kinetics of NO, N<sub>2</sub>O and N-2 production as affected by soil pH, Fems
- 824 Microbiology Ecology, 72, 407-417, 10.1111/j.1574-6941.2010.00856.x, 2010.
- 825 Mantoura, R. F. C., Law, C. S., Owens, N. J. P., Burkill, P. H., Woodward, E. M. S., Howland,
- 826 R. J. M., and Llewellyn, C. A.: Nitrogen biogeochemical cycling in the northwestern indian-
- 827 ocean, Deep-Sea Research Part II-Topical Studies in Oceanography, 40, 651-671, 1993.
- 828 Myhre, G., Shindell, D., Bréon, F.-M., Collins, W., Fuglestvedt, J., Huang, J., Koch, D.,
- 829 Lamarque, J.-F., Lee, D., Mendoza, B., Nakajima, T., Robock, A., Stephens, G.,
- 830 Takemura, T. and Zhang, H.: Anthropogenic and Natural Radiative Forcing. In:
- 831 Climate Change 2013: The Physical Science Basis. Contribution of Working Group I to
- the Fifth Assessment Report of the Intergovernmental Panel on Climate Change, 2013.
- 833 Nevison, C. D., Lueker, T. J., and Weiss, R. F.: Quantifying the nitrous oxide source
- from coastal upwelling, Global Biogeochemical Cycles, 18, GB1018,
- 835 10.1029/2003gb002110, 2004.
- 836 Nevison, C., Butler, J. H., and Elkins, J. W.: Global distribution of N<sub>2</sub>O and the Delta N<sub>2</sub>O-
- AOU yield in the subsurface ocean, Global Biogeochemical Cycles, 17, 1119,
- 838 10.1029/2003gb002068, 2003.

- 839 Nevison, C. D., Weiss, R. F., and Erickson, D. J.: Global oceanic emissions of nitrous-oxide,
- 840 Journal of Geophysical Research-Oceans, 100, 15809-15820, 10.1029/95jc00684, 1995.
- 841 Oschlies, A., Schulz, K. G., Riebesell, U., and Schmittner, A.: Simulated 21st century's increase
- 842 in oceanic suboxia by CO2-enhanced biotic carbon export, Global Biogeochemical Cycles, 22,
- 843 GB4008, 10.1029/2007gb003147, 2008.
- 844 Prather, M. J., Holmes, C. D., and Hsu, J.: Reactive greenhouse gas scenarios: Systematic
- 845 exploration of uncertainties and the role of atmospheric chemistry, Geophysical Research Letters,
- 846 39, L09803, 10.1029/2012gl051440, 2012.
- 847 Punshon, S., and Moore, R. M.: Nitrous oxide production and consumption in a eutrophic
- 848 coastal embayment, Marine Chemistry, 91, 37-51, 10.1016/j.marchem.2004.04.003, 2004.
- 849 Ravishankara, A. R., Daniel, J. S., and Portmann, R. W.: Nitrous Oxide (N2O): The Dominant
- 850 Ozone-Depleting Substance Emitted in the 21st Century, Science, 326, 123-125,
- 851 10.1126/science.1176985, 2009.
- 852 Resplandy, L., Levy, M., Bopp, L., Echevin, V., Pous, S., Sarma, V. V. S. S., and Kumar, D.:
- 853 Controlling factors of the oxygen balance in the Arabian Sea's OMZ, Biogeosciences, 9, 5095-
- 854 5109, 10.5194/bg-9-5095-2012, 2012.
- 855 Riebesell, U., Schulz, K. G., Bellerby, R. G. J., Botros, M., Fritsche, P., Meyerhoefer, M., Neill,
- 856 C., Nondal, G., Oschlies, A., Wohlers, J., and Zoellner, E.: Enhanced biological carbon
- 857 consumption in a high CO2 ocean, Nature, 450, 545-548, 10.1038/nature06267, 2007.
- 858 Sarmiento, J. L., Slater, R., Barber, R., Bopp, L., Doney, S. C., Hirst, A. C., Kleypas, J., Matear,
- 859 R., Mikolajewicz, U., Monfray, P., Soldatov, V., Spall, S. A., and Stouffer, R.: Response of ocean
- 860 ecosystems to climate warming, Global Biogeochemical Cycles, 18, GB3003,
- 861 10.1029/2003gb002134, 2004.
- 862 Steinacher, M., Joos, F., Frolicher, T. L., Bopp, L., Cadule, P., Cocco, V., Doney, S. C., Gehlen,
- 863 M., Lindsay, K., Moore, J. K., Schneider, B., and Segschneider, J.: Projected 21st century
- decrease in marine productivity: a multi-model analysis, Biogeosciences, 7, 979-1005, 2010.
- 865 Stocker, B. D., Roth, R., Joos, F., Spahni, R., Steinacher, M., Zaehle, S., Bouwman, L., Xu, R.,
- 866 and Prentice, I. C.: Multiple greenhouse-gas feedbacks from the land biosphere under future
- 867 climate change scenarios, Nature Climate Change, 3, 666-672, 10.1038/nclimate1864, 2013.
- 868 Suntharalingam, P., and Sarmiento, J. L.: Factors governing the oceanic nitrous oxide
- 869 distribution: Simulations with an ocean general circulation model, Global Biogeochemical
- 870 Cycles, 14, 429-454, 10.1029/1999gb900032, 2000.

- 871 Suntharalingam, P., Sarmiento, J. L., and Toggweiler, J. R.: Global significance of nitrous-oxide
- 872 production and transport from oceanic low-oxygen zones: A modeling study, Global
- 873 Biogeochemical Cycles, 14, 1353-1370, 10.1029/1999gb900100, 2000.
- 874 Suntharalingam, P., Buitenhuis, E., Le Quere, C., Dentener, F., Nevison, C., Butler, J. H.,
- 875 Bange, H. W., and Forster, G.: Quantifying the impact of anthropogenic nitrogen deposition on
- oceanic nitrous oxide, Geophysical Research Letters, 39, L07605, 10.1029/2011gl050778, 2012.
- 877 Suthhof, A., Ittekkot, V., and Gaye-Haake, B.: Millennial-scale oscillation of denitrification
- 878 intensity in the Arabian Sea during the late Quaternary and its potential influence on
- atmospheric N<sub>2</sub>O and global climate, Global Biogeochemical Cycles, 15, 637-649,
- 880 10.1029/2000gb001337, 2001.
- 881 Tagliabue, A., Bopp, L., and Gehlen, M.: The response of marine carbon and nutrient cycles to
- 882 ocean acidification: Large uncertainties related to phytoplankton physiological assumptions,
- 883 Global Biogeochemical Cycles, 25, GB3017, 10.1029/2010gb003929, 2011.
- 884 Takahashi, T., Broecker, W. S., and Langer, S.: Redfield ratio based on chemical-data from
- isopycnal surfaces, Journal of Geophysical Research-Oceans, 90, 6907-6924,
- 886 10.1029/JC090iC04p06907, 1985.
- 887 Tiedje, J.M.: Ecology of denitrification and dissimilatory nitrate reduction to
- ammonium. Biology of anaerobic microorganisms, 179–244, 1988.
- 889 Wanninkhof, R.: Relationship between wind-speed and gas-exchange over the ocean, Journal of
- 890 Geophysical Research-Oceans, 97, 7373-7382, 10.1029/92jc00188, 1992.
- 891 Weiss, R. F., and Price, B. A.: Nitrous-oxide solubility in water and seawater, Marine Chemistry,
- 892 8, 347-359, 10.1016/0304-4203(80)90024-9, 1980.
- 893 Westley, M. B., Yamagishi, H., Popp, B. N., and Yoshida, N.: Nitrous oxide cycling in the
- 894 Black Sea inferred from stable isotope and isotopomer distributions, Deep-Sea Research Part
- 895 Ii-Topical Studies in Oceanography, 53, 1802-1816, 10.1016/j.dsr2.2006.03.012, 2006.
- 896 Yoshida, N., Morimoto, H., Hirano, M., Koike, I., Matsuo, S., Wada, E., Saino, T., and
- 897 Hattori, A.: Nitrification rates and N-15 abundances of N<sub>2</sub>O and NO<sub>3</sub><sup>-</sup> in the western north
- 898 pacific, Nature, 342, 895-897, 10.1038/342895a0, 1989.
- 899 Zamora, L. M., and Oschlies, A.: Surface nitrification: A major uncertainty in marine N<sub>2</sub>O
- 900 emissions, Geophysical Research Letters, 41, 4247-4253, 10.1002/2014g1060556, 2014.
- 901 Zamora, L. M., Oschlies, A., Bange, H. W., Huebert, K. B., Craig, J. D., Kock, A., and
- 902 Loescher, C. R.: Nitrous oxide dynamics in low oxygen regions of the Pacific: insights from the
- 903 MEMENTO database, Biogeosciences, 9, 5007-5022, 10.5194/bg-9-5007-2012, 2012.

- 904 Zehr, J. P., and Ward, B. B.: Nitrogen cycling in the ocean: New perspectives on processes and
- 905 paradigms, Applied and Environmental Microbiology, 68, 1015-1024, 10.1128/aem.68.3.1015-

- 906 1024.2002, 2002.
- 907

- 908 Table 1: Standard deviation and correlation coefficients between P.TEMP and P.OMZ
- 909 parameterizations with respect to MEMENTO database observations (Bange et al., 2009).

	P.TEMP	P.OMZ	OBS
Standard deviation (in nmol $N_2 O \ L^{\text{-1}})$	12	18	16
Correlation coefficient with obs.	0.49	0.42	-

913	Y	
914		Jorge 5/8/15 8
915		Deleted:

## :03 PM

- 922 Fig.1:  $N_2O$  sea-to-air flux (in mgN  $m^{\text{-2}}\,\text{yr}^{\text{-1}})$  from (a) P.TEMP parameterzation averaged for the
- 923 1985 to 2005 time period in the historical simulation, (b) P.OMZ parameterization over the
- 924 same time period, (c) data product of Nevison et al. (2004) and (d) latitudinal  $N_2O$  sea-to-air
- 925

80°S

40°S

(d)

- 926

930





33

80°N

40°N

0°

Latitude

- $931 \qquad \mbox{Fig.2: Global average depth profile of $N_2O$ concentration (in nmol $L^{-1}$) from the MEMENTO}$
- 932 database (dots) (Bange et al., 2009), P.TEMP (blue) and P.OMZ (red). Model
- 933  $\hfill parameterizations are averaged over the 1985 to 2005 time period from the historical$
- 934 simulation.



- $937 \qquad \mbox{Fig. 3: Relationship between } O_2 \mbox{ concentration (in $\mu$mol $L^{-1}$) and $N_2O$ concentration (in $n$mol $L^{-1}$)}$
- 938 in the MEMENTO database (black) (Bange et al., 2009), compared to model (a) P.TEMP (blue)
- 939  $\ \ \,$  and (b) P.OMZ (red) parameterizations averaged over the 1985 to 2005 time period from the
- 940 historical simulation.



- $947 \qquad \mbox{Fig 4: (a) $N_2$O sea-to-air flux (in $TgN yr^{-1}$) from 1851 to $2100$ in $P.TEMP$ (blue) and $P.OMZ$}$
- 948 (red) using the historical and future RCP8.5 simulations. Dashed lines indicate the mean value
- 949 over the 1985 to 2005 time period. Change in  $N_2O$  sea-to-air flux (mgN m  $^{-2}yr \cdot ^{1})$  from the
- $950 \qquad {\rm averaged} \ 2080\text{-}2100 \ {\rm to} \ 1985\text{-}2005 \ {\rm time} \ {\rm periods} \ {\rm in} \ {\rm future} \ {\rm RCP8.5} \ {\rm and} \ {\rm historical} \ {\rm simulations} \ {\rm in}$
- 951 (b) P.TEMP and (c) P.OMZ parameterizations.







958 (c)

- 959 Fig 5: (a) Anomalies in export of organic matter at 100m (green), low-O2 production pathway 960 (short dashed red), high-O2 production pathway (long dashed red) and total P.OMZ production 961 (red) from 1851 to 2100 using the historical and future RCP8.5 simulations. (b) Anomalies in 962 export of organic matter at 100m (green) and P.TEMP production (blue) over the same time 963 period. (c) Change in high-O2 production pathway of N2O (in mgN m<sup>-2</sup> yr<sup>-1</sup>) in the upper 964 1500m between 2080-2100 to 1985-2005 averaged time periods. Hatched areas indicate 965 regions where change in export of organic matter at 100m deep have the same sign as in 966 changes in high-O2 production pathway. (d) Change in low-O2 production pathway of N2O (in 967 mgN  $m^{-2}$  yr<sup>-1</sup>) in the upper 1500m between 2080-2100 to 1985-2005 averaged time periods. 968 Hatched areas indicate regions where oxygen minimum zones ( $O_2 \le 5 \mu mol L^{-1}$ ) expand. (e) 969 Volume (in  $10^6$  km<sup>3</sup>) of hypoxic (black,  $O_2 \le 60 \ \mu mol \ L^{-1}$ ) and suboxic (red,  $O_2 \le 5 \ \mu mol \ L^{-1}$ ) 970 areas in the 1851 to 2100 period in NEMO-PISCES historical and future RCP8.5 simulations.
- 971





- 974  $\;$  Fig 6: (a) Anomalies in  $N_2O$  inventory (in TgN) from 1851 to 2100 in P.TEMP (blue) and
- 975 P.OMZ (red) using the historical and future RCP8.5 simulations in the upper 1500m. Change
- 976 in vertically integrated  $N_2O$  concentration (in mgN m<sup>-2</sup>) in the upper 1500m using NEMO-
- 977 PISCES model mean from the averaged 2080-2100 to 1985-2005 time periods in future
- 978 RCP8.5 and historical scenarios respectively in (b) P.TEMP and (c) P.OMZ. Hatched areas
- 979 indicate regions where the annual mean mixed layer depth is reduced by more than 5m in
- 980 2080-2100 compared to 1985-2005.





- 986 Fig. 7: Change in the whole water column in  $N_2O$  sea-to-air flux (blue), high- $O_2$  production
- 987 pathway (red), low-O<sub>2</sub> production pathway (orange), total N<sub>2</sub>O production (yellow) and N<sub>2</sub>O
- 988 inventory (green) for P.OMZ from the averaged 2080-2100 to present 1985-2005 averaged
- 989 time period in the NEMO-PISCES historical and future RCP8.5 simulations (based on Mikaloff-
- 990 Fletcher et al. (2006) oceanic regions).





2

# 995996Figure 8; Distribution of O2 concentration in NEMO-PISCES 1985 to 2005 averaged time

# Jorge 5/8/15 8:02 PM

**Deleted:** Fig. 8: Box model results, analyzing the effect of changes in ocean circulation by reducing the mixing coefficient ( $\mu$  in %) and changes in biogeochemistry by reducing export of organic matter (in %) separately in N<sub>2</sub>O sea-to-air emissions and N<sub>2</sub>O inventory in 2100. (a) Constant regimes in percentage of the historical N<sub>2</sub>O sea-to-air flux: 95% pink, 90% blue, 85% cyan and 80% green, and (b) Constant regimes in percentage of the historical N<sub>2</sub>O concentration in the deep: 90% pink, 110% blue, 125% cyan and 150% green.



997period (black) compared to the oxygen-corrected World Ocean Atlas (red) from Bianchi et al.998(2012). Interval widths are  $O_2$  concentrations at steps of 5 µmol L<sup>-1</sup>.



999

1000

1019

**1016** Figure 9; Averaged  $O_2$  concentration between 200-600m depth (in  $\mu$ mol L<sup>-1</sup>) (left) and export

1017 of carbon (in mmolC  $m^{-2} d^{-1}$ ) (right) in (a) WOA2005\* and Dunne et al. (2007), (b) CMIP5

1018 model mean historical simulations over the 1985-2005 time period and (c) NEMO-PISCES for

Jorge 5/8/15 8:01 PM Deleted: 10

1020 a. WOA2005\* and Dunne et al., 2007

the present  $1985\mathchar`-2005$  time period.



1022 b. CMIP5 model mean

c. NEMO-PISCES



1023 1024



## **1027** SUPPLEMENTARY MATERIAL

1028

**1029** The  $O_2$  modulation fuction  $f(O_2)$  in P.OMZ is defined as,

$$f(O_2) = \begin{cases} \frac{O_2}{O_2^{*1}} & O_2 < O_2^{*1} \\ 1 & O_2^{*1} < O_2 < O_2^{*2} \\ 0.7 \cdot exp - 0.5(O_2 - O_2^{*2})/O_2^{*2} + \\ 0.3 \cdot exp - 0.05(O_2 - O_2^{*2})/O_2^{*2} & O_2 \ge O_2^{*2} \end{cases}$$

1030

1031 where  $O_2^{*1}$  is 1 µmol L<sup>-1</sup> and  $O_2^{*2}$  is 5 µmol L<sup>-1</sup>. The shape of the function is shown in Fig. S1. 1032

1033 Fig. S1: Oxygen modulating function  $f(O_2)$  in the low-O<sub>2</sub> production pathway term included in

1034 P.OMZ from Goreau et al. (1980).





Fig. S2: Vertically integrated (a) high-O\_2 and (b) low-O\_2 production pathways (in gN  $m^{\text{-2}}\ yr^{\text{-1}})$ 

in  $\ensuremath{\text{P.OMZ}}$  for the averaged 1985 to 2005 historical simulation.



#### 1042 . 1043

## Jorge 5/8/15 8:01 PM

**Deleted:** Fig. S3: Diagram of the box model. N<sub>2</sub>O inventory is separated into surface and deep concentrations above and below 100m. The fraction of N<sub>2</sub>O outgassed to the atmosphere (k), mixing ratio (v) between deep and surface and the rate of N<sub>2</sub>O production from the export of organic matter to depth (e) regulate the N<sub>2</sub>O budget in the ocean interior.