1 Supplementary Information of

# 2 Global Nitrous Oxide Budget 1980-2020

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#### 5 Extended methodology 6

#### SI-1 NMIP-2: global Nitrogen/N<sub>2</sub>O Model Inter-comparison Project phase 2

9 The NMIP2 is a follow-up model intercomparison project of NMIP (*Tian et al.*, 2018), which provides 10 estimates of N<sub>2</sub>O emissions from natural and agricultural soils and covers the time period 1850-2020. Eight 11 process-based Terrestrial Biosphere Models (TBMs) participate in NMIP-2. In general, N<sub>2</sub>O emissions 12 from soil are regulated at two levels, which are the rates of nitrification and denitrification in the soil and 13 soil physical factors regulating the ratio of N<sub>2</sub>O to other nitrous gases (*Davidson et al.*, 2000). For N input 14 to land ecosystems, all eight models considered N fertilizer use, atmospheric N deposition and biological 15 fixation, but five models considered manure as N input. For vegetation processes, all models included 16 dynamic algorithms in simulating N allocation to different living tissues and vegetation N turnover, and simulated plant N uptake using the "Demand and Supply-driven" approach. For soil N processes, all eight 17 18 models simulated N leaching according to water runoff rate; however, models are different in representing 19 nitrification and denitrification processes and the impacts of soil chemical and physical factors. The 20 differences in simulating nitrification and denitrification processes are one of the major uncertainties in 21 estimating N<sub>2</sub>O emissions. Model characteristics in simulating major N cycling processes associated with 22 N<sub>2</sub>O emissions in each participating model are briefly described in Table SI-1.

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24	Table SI-1. M	odel char	acteristics i	n simulatin	g major l	N cycling p	rocesses

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	CLASS IC	DLEM	ELM	ISAM	LPX- Bern	O-CN	ORCHIDEE	VISIT
Open C cycle	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes
C-N coupling	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes
N pools <sup>b</sup>	(3, 1, 3)	(6,6,8)	(6,4,5)	(6,4,4)	(4,3,8)	(9,6,9)	(9,6,9)	(4,1,4)
Demand and supply– driven plant N uptake	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes
N allocation <sup>c</sup>	Dynami	Dynami	Dynam	Dynam	Dynami	Dynami	Dunamia	Dynam
	c	c	ic	ic	c	c	Dynamic	ic
Nitrification	f(T, SWC, C <sub>NH4</sub> )	f(T, SWC, C <sub>NH4</sub> )	f(T, SWC, pH, rh, C <sub>NH4</sub> )	f(T, SWC, C <sub>NH4</sub> )	f(T, SWC, C <sub>NH4</sub> )	<i>f(T,</i> <i>SWC,</i> <i>pH,</i> <i>C</i> <sub>NH4</sub> )	f(T, SWC, pH, C <sub>NH4</sub> )	<i>f(T,</i> <i>SWC,</i> <i>pH,</i> <i>C</i> <sub>NH4</sub> )
Denitrificatio n	f(T, SWC, C <sub>NO3</sub> )	f(T, SWC, clay, rh, C <sub>NO3</sub> )	f(T, SWC, pH, rh, C <sub>NO3</sub> )	$f(T, SWC, C_{NO3})$	$f(T, SWC, R_{mb}, C_{NO3})$	f(T, SWC, pH, R <sub>m</sub> b, C <sub>NO3</sub> )	f(T, SWC, pH, denitrif ier, C <sub>NO3</sub> )	f(SWC, rh, C <sub>NO3</sub> )
Mineralizatio n, immobilizati on	f(C:N)	f(C:N)	f(C:N)	f(C:N)	f(C:N)	f(C:N)	f(C:N)	f(C:N)
N leaching	f(runoff , $C_{NO3}$ , $C_{NH4}$	f(runoff , $C_{NO3}$ , $C_{NH4}$	f(runoff , $C_{NO3}$ )	<i>f(runof</i> <i>f,</i> C <sub>NO3</sub> , C <sub>NH4</sub> )	f(runoff , $C_{NO3}$ )	f(runoff , $C_{NO3}$ , $C_{NH4}$	f(runoff, C <sub>NO3</sub> , C <sub>NH4</sub> )	f(runoff , C <sub>NO3</sub> )
NH <sub>3</sub> volatilization	$f(C_{NH3})$	f(T, SWC,	No	$f(C_{NH3})$	f(T, SWC,	$f(C_{NH3})$	f(SWC, pH, C <sub>NH4</sub> )	f(T, SWC,

		рН, С <sub>NH4</sub> )			рН, С <sub>NH4</sub> )			рН, С <sub>NH4</sub> )
Plant N	Dynami	Dynami	Dynam ic	Dynam	Dynami	Dynami	Dynamic	Dynam
N resorption	Fixed	f(C:N)	Fixed	f(C:N)	Fixed	Fixed	Fixed	Fixed
N fixation	$f(N_{limit})$	f(T, SWC, C <sub>NH4</sub> , C <sub>NO3</sub> )	f(T, C:N)	f(ET)	Implied by mass balance	$f(N_{limit})$	Fixed	f(ET)
N fertilizer use	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes
Manure N use	No	Yes	No	Yes	No	Yes	Yes	Yes
N deposition	Yes	Yes	Yes	Yes	Yes	Yes	Yes	Yes

<sup>a</sup> "Open" denotes that excess N can be leached from the system.

26 <sup>b</sup> Numbers of N pools (vegetation pools, litter pools, soil pools).

27 <sup>c</sup> Dynamic denotes time-varied N allocation ratio to different N pools.

28 *T: soil temperature, SWC: soil water content, clay: soil clay fraction, ET: evapotranspiration, denitrifier:* 

29 soil denitrifier biomass, rh: soil heterogeneous respiration, Nlimit: N limitation of vegetation growth,

30  $C_{NO3}$ : soil  $NO_3^-$  content,  $C_{NH4}$ : soil  $NH_4^+$  content.

31

32 All NMIP2 models are driven by consistent input datasets (i.e., climate, atmospheric CO<sub>2</sub> concentration, 33 land cover change, irrigation, atmospheric N deposition, mineral N fertilization, and manure N application 34 and deposition) and implemented consistent simulation experiments (SH0 - SH12; Table A4). Nitrogen 35 inputs data used in NMIP2 simulations are from History of anthropogenic Nitrogen inputs (HaNi) dataset 36 (*Tian et al.*, 2022), which takes advantage of different data sources in a spatiotemporally consistent way to 37 generate a set of high-resolution (5 arcminutes) gridded N input products from 1850 to 2020. HaNi data set 38 shows that the total anthropogenic N inputs to global terrestrial ecosystems increased from 29.05 Tg N yr 39 <sup>1</sup> in the 1860s to 267.23 Tg N yr<sup>-1</sup> in the 2010s, with the dominant N source changing from atmospheric N 40 deposition (before the 1900s) to manure N (the 1910s-2000s), and to synthetic fertilizer in the 2010s (Fig. 41 B3). The climate data used to run historical simulations is the half degree CRU-JRA2.2 6-hourly forcing 42 over 1901- 2020 (https://catalogue.ceda.ac.uk/uuid/4bdf41fc10af4caaa489b14745c665a6). Annual CO<sub>2</sub> 43 concentration during 1850-2020 were derived from ice core CO<sub>2</sub> data and NOAA annual 44 observations(https://www.esrl.noaa.gov). Historical distribution of cropland, pasture, rangeland and 45 irrigation during 1850-2020 were from Land-Use Harmonization 2 (LUH2) dataset (Hurtt et al., 2020). The 46 original dataset of LUH2 is at a resolution of 0.25° x 0.25° longitude/latitude. We aggregated all geo-47 referenced input data into a consistent spatial resolution of 0.5° x 0.5° longitude/latitude to run NMIP2 models.

48 49

50 NMIP2 models perform a subset of 13 simulations (SH0-SH12) to quantify N<sub>2</sub>O emissions from both 51 agricultural and natural soils during the study period, and to disentangle the effects of multiple 52 environmental factors on soil N<sub>2</sub>O emissions. The SH1 results were taken as the "best estimates" of soil 53 N<sub>2</sub>O emissions because they include the effects of all driving factors that models can take into account. In 54 the SH0 simulation, driving forces were kept constant at the level in 1850 over the entire simulation period 55 (1850-2020). According to previous N<sub>2</sub>O budget studies, atmospheric N<sub>2</sub>O growth rate and Monte-Carlo 56 method, we suggest the following criteria for the  $N_2O$  budget inclusion (Table A6), and the criteria for 57 carbon components are consistent with TRENDY. By comparing results from factorial simulation 58 experiments (SH0 - SH12), we attribute changes in soil N<sub>2</sub>O emissions to seven natural and anthropogenic 59 factors, namely, climate (CLIM, including precipitation, humidity, temperature and photosynthetic active

60 radiation changes), atmospheric CO<sub>2</sub> concentration (CO<sub>2</sub>), land cover change (LCC), irrigation (IRRI),

61 atmospheric N deposition (NDEP), mineral N fertilizer use (NFER), and manure N use in cropland 62 (MANN). In order to understand soil  $N_2O$  emissions dynamics caused by crop cultivation, we further 63 separate the global and regional N<sub>2</sub>O emissions into those derived from cropland soils and those from soils 64 of other land ecosystems. In this study, we attribute the impact of a single factor on cropland N<sub>2</sub>O emissions. 65 Five models (DLEM, ISAM, O-CN, ORCHIDEE, and VISIT) considered the effects of manure N application in cropland, therefore, we use these five models' results to calculate the manure N effect (SH1-66 67 SH2). Meanwhile, we used results from all the eight models (i.e., CLASSIC, DLEM, ELM, ISAM, LPX-68 Bern, O-CN, ORCHIDEE, and VISIT) to calculate the effects of synthetic N fertilizer use (SH1-SH3) and 69 atmospheric N deposition (SH1-SH4). The effect of N deposition in natural ecosystems (SH1-SH4) and the 70 effects of CO<sub>2</sub> (SH1-SH7) and climate (SH1-SH8) on global terrestrial ecosystems are calculated from the 71 eight NMIP2 models mentioned above.

#### 72 73 74

Carbon criteria	N <sub>2</sub> O criteria
(1) Steady state after spin-up, diagnosed from SH0 run: steady-state defined as an offset < $0.10$ PgC yr <sup>-1</sup> , drift < $0.05$ PgC yr <sup>-1</sup> per century (i.e. first is the average over 1850-2020, second is the slope x 100).	(1) Steady state after spin-up, diagnosed from SH0 run: drift < 0.2 Tg N <sub>2</sub> O-N yr <sup>-1</sup> per century (i.e. the slope x 100).
(2) Net annual land flux is a carbon sink over the 1990s and 2000s as constrained by global atmospheric and oceanic observations (Keeling & Manning, 2014), diagnosed from SH3 run.	(2) Inside the present-day (2007-2016) land emission range: 7- 13 Tg N <sub>2</sub> O-N yr <sup>-1</sup> , diagnosed from SH1 run. The upper limit was calculated using the maximum total N <sub>2</sub> O emissions minus the minimum estimates of other sources, and the lower limit was calculated using the minimum total N <sub>2</sub> O emissions minus the maximum estimates of other sources. The range of total emissions was estimated by a one-box model using atmospheric N <sub>2</sub> O growth rate, and the range of the sum of other sources was calculated by a Monte-Carlo method using estimates from Tian et al. (2020).
	(3) Inside the pre-industrial land emission range: 3 to 9 Tg $N_2O$ -N yr <sup>-1</sup> , diagnosed from SH1 run. This range is derived from the pre-industrial atmospheric burden/N <sub>2</sub> O lifetime minus ocean and river/ coastal/estuary emissions (Michael J. Prather et al., 2015).

## Table SI-2. Criteria for the N<sub>2</sub>O budget inclusion

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# 76

#### 77 Brief description of algorithms associated with N<sub>2</sub>O flux in each NMIP2 model:

#### 78 1: CLASSIC

- 79 The representation of nitrogen cycling in CLASSIC is described in Asaadi and Arora (2021) and Kou
- Giesbrecht and Arora (2022). N<sub>2</sub>O production due to both nitrification and denitrification are represented. 80 N<sub>2</sub>O loss during nitrification ( $I_{N_2O}$ ; g N m<sup>-2</sup> d<sup>-1</sup>) is represented with the following equation: 81

82 
$$I_{N_2O} = \eta_{N_2O} f_I(T_{0.5}) f_I(\psi) N_{NH_4}$$

- (1) $\eta_{N_0,0}$  is a coefficient (d<sup>-1</sup>),  $f_I(T_{0,5})$  is a dimensionless scalar that depends on soil temperature averaged over 83
- 84 the top 0.5m soil depth  $(T_{0.5})$ ,  $f_I(\psi)$  is a dimensionless scalar that depends on soil matric potential  $(\psi)$ , and
- 85  $N_{NH_4}$  is the soil ammonium pool (g N m<sup>-2</sup>).

86 N<sub>2</sub>O loss during denitrification ( $E_{N_2O}$ ; g N m<sup>-2</sup> d<sup>-1</sup>) is represented with the following equation:

87  $E_{N_2O} = \mu_{N_2O} f_E(T_{0.5}) f_E(\theta) N_{NO_3}$  (2)

88  $\mu_{N_2O}$  is a coefficient (d<sup>-1</sup>),  $f_E(T_{0.5})$  is a dimensionless scalar that depends on soil temperature averaged over

the top 0.5m soil depth  $(T_{0.5})$ ,  $f_E(\theta)$  is a dimensionless scalar that depends on soil moisture  $(\theta)$ , and  $N_{NO_3}$ is the soil nitrate pool (g N m<sup>-2</sup>).

#### 92 **2: DLEM**

The nitrogen cycle scheme in DLEM2.0 (*Xu et al.*, 2017; *Yang et al.*, 2015; Tian et al. 2020) are similar as
DLEM1.0 (*Lu and Tian*, 2013; *Tian et al.*, 2012b; *Tian et al.*, 2010; *Tian et al.*, 2011; *Xu et al.*, 2011),
However, the N<sub>2</sub>O emission schemes in DLEM2.0 (*Xu et al.*, 2017) have been modified based on *Chatskikh et al.* (2005) and *Heinen* (2006).

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91

 $R_{nit} = k_{nit\_max} f(T1) f(WFPS) C_{NH4}$ (3)

$$100 R_{den} = k_{den\_max} f(T2) f(WFPS) C_{NO3} (4)$$

101 102 where  $R_{nit}$  is the daily nitrification rate (g N/m<sup>2</sup>/d);  $R_{den}$  is the daily denitrification rate (g N/m<sup>2</sup>/d); 103 f(T1) and f(T2) are the impact function of daily soil temperature on nitrification and denitrification, 104 respectively; f(WFPS) is the impact function of water-filled pore space (WFPS) on nitrification, 105 denitrification and N<sub>2</sub>O diffusion;  $k_{nit\_max}$  is the maximum fraction of NH<sub>4</sub><sup>+</sup>-N that is converted to NO<sub>3</sub><sup>-</sup>-106 N or gases (0-1);  $k_{den\_max}$  is the maximum fraction of NO<sub>3</sub><sup>-</sup>-N that is converted to gases (0-1);  $C_{NH4}$  and 107  $C_{NO3}$  are the soil NH<sub>4</sub><sup>+</sup>-N and NO<sub>3</sub><sup>-</sup>-N content (g N/m<sup>2</sup>). N<sub>2</sub>O from denitrification and nitrification processes 108 are calculated as,

$$R_{N20} = (R_{nit} + R_{den})f(T3)(1 - f(WFPS))$$
(5)

111

where  $R_{N20}$  is the daily N<sub>2</sub>O emission rate (g N/m<sup>2</sup>/d); f(T3) is the impact function of daily soil temperature on N<sub>2</sub>O diffusion rate from soil pores. The calculation methods for these functions and parameters were described in detail in *Xu et al.* (2017) and *Yang et al.* (2015).

#### 116 **3: ELM**

117 The nitrogen dynamics in ELM is simulated based on the theory of equilibrium chemistry approximation 118 (Zhu et al., 2016). Plants, soil microbes, and abiotic factors such as mineral surfaces coexist in the same 119 soil environment and vie for a diverse array of nutrients, including NH4+, NO3-. Due to the limited 120 availability of these nutrients, intense competitive interactions are expected. The competition of those 121 limited resources is represented by consumer-substrate networks, therefore, the uptake of nutrient substrate 122 by each consumer is dependent on the relative competitiveness of one consumer over the others. Nutrient 123 consumers' competitiveness is parametrized with kinetic parameters (Zhu et al., 2016). As a result, neither 124 plan nor soil microbes get the first priority to access nutrient substrates. Instead, when the potential nutrient 125 demands (from all nutrient consumers) exceed the supply at a given time step, the allocation of limited 126 nutrients among the consumers affects their performance (e.g., plant growth, soil organic matter 127 accumulation, nitrification, denitrification rates). ELM adopts a multiple-consumer-multiple-substrate 128 competition network (Zhu et al., 2016; Zhu et al., 2019) to simulate (1) nitrogen uptake facilitated by 129 nitrogen carrier enzymes, (2) binding of a nutrient substrate to a particular enzyme precludes it from 130 attaching to other enzymes, and (3) rates and affinities of consumers for different substrates. After the 131 nutrient competition has been resolved, scaling terms ( $f(ECA_{nit})$  and  $f(ECA_{den})$ ) will be applied to the 132 potential nitrification and denitrification processes:

134 
$$R_{nit} = k_{nit\_max} f(\theta) f(T) (1 - f(0)) f(ECA_{nit}) C_{NH4}$$
(6)

- 135
- 136  $R_{den} = min (f(deomp), f(C_{NO3}))f(ECA_{den})$
- 137 138 where  $k_{nit\_max}$  is the maximum nitrification rate,  $f(\theta)$ , f(T), f(0) are soil moisture, temperature, and 139 oxygen scalars, respectively. f(deomp) and  $f(C_{NO3})$  are carbon limited and NO3- limited denitrification 140  $f(C_{NO3})$  are carbon limited and NO3- limited denitrification

(7)

140 rates (Del Grosso et al., 2000).

141

#### 142 **4: ISAM**

143 ISAM model contains detailed calculations of the terrestrial ecosystem's organic and mineral N cycle (Yang 144 et al., 2009). The major N processes in ISAM include biological fixation, leaching, mineralization and 145 immobilization, plant uptake, nitrification, and denitrification. The soil biogeochemistry module of ISAM 146 shares the same ten soil layers (to 3.5 m depth) as the soil biogeophysics and calculates the vertical transport 147 of SOC and N (Shu et al., 2020; Yang et al., 2009). N<sub>2</sub>O emission in ISAM N<sub>2</sub>O is produced as a byproduct 148 of nitrification and denitrification (Xu et al., 2021). N<sub>2</sub>O module explicitly accounts for the vertical transport 149 of C, N, and O<sub>2</sub> within every soil layer for both saturated and unsaturated soil conditions by accounting for 150 the process of oxygen diffusing into the soil from the atmosphere and the soil oxygen supply. The model 151 also explicitly accounts for the effects of anoxic and oxic environments on nitrification (N<sub>ni</sub>, Eq. 6) and 152 denitrification (N<sub>de</sub>, Eq. 7). Both environments are calculated based on the fraction of anoxic soil depending 153 on soil  $O_2$  concentration, which is non-linearly correlated with the chemical pathways forming  $N_2O$ .

155 
$$N_{ni} = NH_4^+ \times (1 - e^{-F_{te_m} \times F_{sm_m} \times r_{ni}}) \times F_{pH_m ni} \times R_d$$
(8)

157 
$$N_{de} = NO_3^- \times r_{de} \times Rh \times F_{pH_m_de} \times R_d$$
(9)

158

159 where  $NH_4^+$  and  $NO_3^-$  are ammonium and nitrate pool sizes;  $F_{te\_m}$  is temperature modifier;  $F_{sm\_m}$  is soil 160 moisture modifier;  $r_{ni}$  and  $r_{de}$  are base nitrification and denitrification rates;  $F_{pH\_m\_ni}$  and  $F_{pH\_m\_de}$  are pH 161 modifiers for nitrification and denitrification;  $R_d$  is relative soil anoxic fraction; Rh (= 1- $R_d$ ) is heterotrophic

162 respiration.163

164 Under anoxic soil conditions, N<sub>2</sub>O is produced through denitrification, while under oxic soil conditions, 165 more  $N_2O$  is produced from nitrification. The model accounts for soil  $NH_4^+$  volatilization at the soil surface 166 when  $NH_4^+$  in  $NH_4^+$ -containing fertilizers (e.g., urea) is converted to ammonia gas, depending upon pH 167 (Huang and Gerber, 2015). The soil  $NH_4^+$  volatilization in the model is also affected by the anoxic 168 condition, which increases under a higher temperature and relatively lesser soil anoxic condition. The model 169 accounts for the impacts of pH on nitrification, denitrification, and volatilization rates (Li et al., 2000; Xu-170 *Ri and Prentice*, 2008). We prescribe the soil pH from the Global Soil Dataset for Earth System Modeling 171 dataset (GSDE) (Shangguan et al., 2014).

#### 173 **5: LPX-Bern**

The implementation of nitrogen dynamics in LPX-Bern is based on the work of *Xu-Ri and Prentice* (2008). Nitrogen uptake by plants is governed by their demand and the availability of nitrogen in two soil pools representing ammonium and nitrate. Nitrogen from deposition and fertilization are added to these inorganic soil pools. Losses include ammonium volatilization, nitrate leaching as well as N<sub>2</sub>O and NO production during nitrification and N<sub>2</sub>O, NO and N<sub>2</sub> production during denitrification. Aerobic nitrification of ammonium is dependent on soil temperature ( $T_{soil}$ ) and indirectly on soil water content due to the partitioning of wet and dry soil:

181

$$\begin{array}{l} 182 \qquad R_{nit} = max_{nit}f_1(T_{soil})C_{NH4,dry} \tag{10} \\ 183 \end{array}$$

- where  $max_{nit} = 0.92 \ day^{-1}$  is the daily maximum nitrification rate at 20°C.
- 185 Anaerobic denitrification of nitrate in wet soil depends on labile carbon availability and soil temperature: 186

187 
$$R_{den} = R_{mb} / (R_{mb} + K_{mb}) f_2(T_{soil}) C_{NO3,wet} / (C_{NO3,wet} + K_n)$$
(11)

192

189 The parameters  $K_{mb}$  and  $K_n$  are taken from *Xu-Ri and Prentice* (2008) and  $R_{mb}$  is the microbiotical soil 190 respiration. The amount of nitrogen lost as N<sub>2</sub>O due to nitrification and denitrification is modelled as a 191 function of soil temperature, water content and the respective process rate.

#### 193 **6: O-CN**

194 The treatment of inorganic soil nitrogen dynamics in O-CN follows largely *Xu-Ri and Prentice* (2008). O-195 CN (*Zaehle and Friend*, 2010) considers N losses to NH<sub>3</sub> volatilisation, NOx, N<sub>2</sub>O and N<sub>2</sub> production and 196 emission, as well as NH<sub>4</sub> and NO<sub>3</sub> leaching. Inorganic nitrogen dynamics in the soil are tightly coupled to 197 plant uptake and net mineralization. The anaerobic volume fraction of the soil is estimated by an empirical 198 function of the fractional soil moisture content (*Zaehle et al.*, 2011). The fraction of ammonium in the 199 aerobic part of the soil is subject to nitrification, according to:

200

206

208

$$\begin{array}{l} 201 \qquad R_{nit} = vmax_{nit}f(T1)f(pH1)C_{NH4}\\ 202 \end{array}$$

where f(pH1) is the soil pH response functions for nitrification (*Li et al.*, 1992; *Xu-Ri and Prentice*, 2008), and  $vmax_{nit}$  is the maximum daily nitrification rate under 20°C and favourable pH conditions (*Xu-Ri and Prentice*, 2008).

(12)

207 Gross denitrification of the fraction of nitrate under anoxic conditions is modelled as:

209 
$$R_{den} = R_{mb} / (R_{mb} + K_{mb}) f(T2) f(pH2) C_{NO3} / (C_{NO3} + K_n)$$
(13)  
210

where f(pH2) is the soil pH response functions for denitrification (*Li et al.*, 1992; *Xu-Ri and Prentice*, 2008), *R<sub>mb</sub>* is the soil microbial respiration rate, and *K<sub>mb</sub>* and *K<sub>n</sub>* parameters taken from *Li et al.* (1992).

 $\begin{array}{ll} \text{214} & \text{The } N_2 O \text{ production from nitrification and denitrification is then calculated as:} \\ \text{215} \end{array}$ 

216 
$$R_{N20} = a_{nit} f(T1) R_{nit} + b_{den} f(T2) f(pH3) R_{den}$$
(14)  
217 (14)

where  $a_{nit}$  and  $b_{denit}$  are fraction loss constants, f(pH3) is a pH-modifier changing the degree of denitrification producing N<sub>2</sub>O versus NO<sub>x</sub> or N<sub>2</sub> (*Zaehle et al.*, 2011). Emissions of volatile compounds are simulated using the empirical emission of *Xu-Ri and Prentice* (2008).

#### **222 7: ORCHIDEE**

Modeling of the mineral N dynamics by the ORCHIDEE model originates from the formulations used in the O-CN (*Zaehle and Friend*, 2010). It is composed of five pools for ammonium/ammoniac, nitrate, NOx, nitrous oxide, and di-nitrogen forms. N<sub>2</sub>O production in both nitrification and denitrification processes are represented.

227

The potential daily rate of nitrification,  $R_{nit}$ , occurs only on the aerobic fraction of the soil and is a function of temperature, pH, and ammonium concentration ( $C_{NH4}$ ):

$$R_{nit} = (1 - f(WFPS))f(T1)f(pH1)k_{nit}C_{NH4}$$
(15)

233 where  $k_{\text{nit}}$  is the reference potential NO3<sup>-</sup> production per mass unit of ammonium.

## 235 8: VISIT

The nitrogen cycle scheme of VISIT is composed of three organic soil nitrogen pools (microbe, litter, and humus), two inorganic soil nitrogen pools (ammonium and nitrate), and vegetation pools. Fertilizer is considered as an input to the ammonium and nitrate pools at a fixed ratio, and manure as an input into the litter organic nitrogen pool. N<sub>2</sub>O emissions through nitrification and denitrification are estimated using the scheme developed by *Parton et al.* (1996). Nitrification-associated N<sub>2</sub>O emission ( $R_{nit,N20}$ ) is evaluated as follows,

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280

243 
$$R_{nit,N20} = f(WFPS)f(pH1)f(T1)(K_{max} + F_{max}f(NH4))$$
 (16)  
244

where  $K_{max}$  is the soil-specific turnover coefficient;  $F_{max}$  is the parameter of maximum nitrification gas flux; and  $f(NH_4)$  is the effect of soil ammonium on nitrification. Denitrification-associated N<sub>2</sub>O emission  $(R_{den,N2O})$  is evaluated by the following equation:

249 
$$R_{den,N20} = R_{den}(1 + R_{N2/N20})$$
 (17)  
250

$$R_{den} = min (f(NO_3), f(CO_2)) \times f(WFPS)$$

where  $R_{N2/N2O}$  is the fractionation coefficient, which is also a function of WFPS, soil nitrate, and heterotrophic respiration,  $f(NO_3)$  is the maximum denitrification rate in high soil respiration rate condition,  $f(CO_2)$  is the maximum denitrification rate in high NO<sub>3</sub><sup>-</sup> levels, and f(WFPS) is the effect of WFPS on denitrification rate.

258 N<sub>2</sub>O production by nitrification ( $R_{N2O,nit}$ , g N-N<sub>2</sub>O/m<sup>2</sup>/d) is expressed as a function of the potential daily rate 259 of nitrification ( $R_{nit}$ , g N-NO<sub>3</sub><sup>-</sup>/m<sup>2</sup>/d), temperature and the water content as shown in *Zhang et al.* (2002). 260

$$261 \qquad R_{N20,nit} = f(WFPS)f(T1)R_{nit}p_{N20,nit}$$

where  $p_{N20,nit}$  (g N-N<sub>2</sub>O (g N-NO<sub>3</sub><sup>-</sup>)<sup>-1</sup>) is the reference N<sub>2</sub>O production per mass unit of NO<sub>3</sub><sup>-</sup> produced by nitrification. The denitrification occurs on the anaerobic fraction of the soil which is computed as a function of the water-filled porosity (*f*(*WFPS*)) and is controlled by temperature, pH, soil NO concentration and denitrifier microbial activity (*a<sub>microb</sub>*, g m<sup>-2</sup>) (*Li et al.*, 2000).

$$\begin{array}{l} 268 \\ 269 \end{array} R_{N2O,den} = f(WFPS)f(T2)f(pH)f(NO)p_{N2O,den}a_{microb} \tag{20}$$

where f(NO) is a Michaelis-Menten shape function and  $p_{N2O,den}$  is the reference N<sub>2</sub>O production per mass unit of denitrifier microbes.

## 273 SI-2 The FAOSTAT inventory

- The FAOSTAT emissions data (*FAO*, 2022) are computed at Tier 1 following *IPCC* (2006), Vol. 275
- 276 4. The overall equation is as follows:

Direct emissions are estimated at the country level, using the formula:

$$Emission = A * EF$$
(21)

(18)

(19)

where emission represents kg N yr<sup>-1</sup>; A represents the amount of N in the following items (annual synthetic N applications/manure applied to soils/manure left on pasture/manure treated in manure management

- systems/crop residue/biomass burned amount) in kg N yr<sup>-1</sup>; EF = Tier 1, default IPCC emission factors, 283 284 expressed in kg N/kg N. 285 286 Indirect emissions are estimated at the country level, using the formula: 287 288 *Emission* =  $A_{v\&l} * EF$ (22)289 290 where emission represents kg N yr<sup>-1</sup>; A vel represents the fraction of manure/synthetic N fertilizers that 291 volatize as NH<sub>3</sub> and NO<sub>x</sub> and are lost through runoff and leaching in kg N yr<sup>-1</sup>; EF = Tier 1, default IPCC 292 emission factors, expressed in kg N/kg N. 293 294 Synthetic N fertilizers: N<sub>2</sub>O from synthetic fertilizers is produced by microbial processes of nitrification 295 and denitrification taking place on the addition site (direct emissions), and after volatilization/redeposition 296 and leaching processes (indirect emissions). 297 298 Manure management: The term manure includes both urine and dung (i.e., both liquid and solid material) 299 produced by livestock. N<sub>2</sub>O is produced directly by nitrification and denitrification processes in the manure, 300 and indirectly by nitrogen (N) volatilization and redeposition processes. 301 Manure applied to soils: N<sub>2</sub>O is produced by microbial processes of nitrification and denitrification taking 302 place on the application site (direct emissions), and after volatilization/redeposition and leaching processes 303 (indirect emissions). 304 305 Manure left on pastures: N<sub>2</sub>O is produced by microbial processes of nitrification and denitrification taking 306 place on the deposition site (direct emissions), and after volatilization/redeposition and leaching processes 307 (indirect emissions). 308 309 Crop Residue: N<sub>2</sub>O emissions from crop residues consist of direct and indirect emissions from nitrogen (N) 310 in crop residues left on agricultural fields by farmers and from forages during pasture renewal (following 311 the definitions in the IPCC guidelines (IPCC, 2006)). Specifically, N<sub>2</sub>O is produced by microbial processes 312 of nitrification and denitrification taking place on the deposition site (direct emissions), and leaching 313 processes (indirect emissions). 314 315 Cultivation of organic soils: The FAOSTAT domain "Cultivation of organic soils" contains estimates of 316 direct N<sub>2</sub>O emissions associated with the drainage of organic soils – histosols – under cropland and grazed 317 grassland. 318 319 Burning-savanna: N<sub>2</sub>O emissions from the burning of vegetation biomass in the land cover types: Savanna, 320 Woody Savanna, Open Shrublands, Closed Shrublands, and Grasslands, Burning-crop residues: N<sub>2</sub>O produced by the combustion of a percentage of crop residues burnt on-site. Burning-biomass: N2O 321 322 emissions from the burning of vegetation biomass in the land cover types: Humid tropical forests, other 323 forests, and organic soils. 324 325 SI-3 The EDGAR v7.0 inventory 326 The new online version, EDGAR v7.0 (https://edgar.jrc.ec.europa.eu/dataset ghg70) incorporates a full 327 differentiation of emission processes with technology-specific emission factors and additional end-of-pipe 328 abatement measures and as such updates and refines the emission estimates. The emissions are modelled 329 based on the latest scientific knowledge and available global statistics primarily from International Energy
  - Agency (*IEA*, 2021) for energy related sectors, FAO statistics (*FAO*, 2022) for agriculture, which were
  - 331 complemented for the rest of sectors with United States Geological Survey (USGS), International Fertiliser
  - 332 Association (IFA), Gas Flaring Reduction Partnership (GGFR)/U.S. National Oceanic and Atmospheric

Administration (NOAA) and World Steel Association (worldsteel) recent statistics; the methods are those recommended by *IPCC* (2006). Official data submitted by the Annex I countries to the United Nations Framework Convention on Climate Change (UNFCCC) and to the Kyoto Protocol are used to some extent, particularly regarding control measures implemented since 1990 that are not described by international statistics. A fast-Track approach was used to extend the N<sub>2</sub>O emission time series for the latest years up to 2021 (*Crippa et al.*, 2021; *Crippa et al.*, 2022).

339

340 The  $N_2O$  emission factors for direct soil emissions of  $N_2O$  from the use of synthetic fertilizers, from manure

341 used as fertilizers, and from crop residues are taken from *IPCC* (2006), which updated the default IPCC 342 emission factor in the IPCC Good Practice Guidance (2000) with a 20% lower value. N<sub>2</sub>O emissions from 343 the use of animal waste as fertilizer are estimated considering both the loss of N that occurs from manure 344 management systems before manure is applied to soils and the additional N introduced by bedding material 345 (*Janssens-Maenhout et al.*, 2019). N<sub>2</sub>O emissions from fertilizer use and CO<sub>2</sub> from urea fertilization are 346 estimated based on IFA and FAO recent statistics.

347

N<sub>2</sub>O emissions from manure management are based on the distribution of manure management systems from Annex I countries reporting to the UNFCCC, *Zhou et al.* (2007) for China and *IPCC* (2006) for the rest of the countries.

350 re

Different N<sub>2</sub>O emission factors are applied to tropical and non-tropical regions. N and dry matter content of agricultural residues are estimated from the cultivation area and yield for 24 crop types from *FAO* (2022) and using emission factors of *IPCC* (2006).

355

Indirect N<sub>2</sub>O emissions from leaching and runoff of nitrate are estimated from N input to agricultural soils. Leaching and runoff are assumed to occur in all agricultural areas except non-irrigated dryland regions, which are identified with maps of FAO Geonetwork (https://www.fao.org/land-water/databases-andsoftware/geonetwork/en/). The fraction of N lost through leaching and runoff is based on the study of *Van Drecht et al.* (2003). The updated emission factor for indirect N<sub>2</sub>O emissions from N leaching and run-off from the *IPCC* (2006) guidelines is selected, while noting that it is 70% lower than the mean value of the 1996 IPCC Guidelines and the IPCC Good Practice Guidance *IPCC* (1996; 2000).

363

Indirect N<sub>2</sub>O emissions from atmospheric deposition of N of NO<sub>x</sub> and NH<sub>3</sub> emissions from non-agricultural sources, mainly fossil fuel combustion, are estimated using N in NO<sub>x</sub> and NH<sub>3</sub> emissions from these sources as activity data, based on EDGAR v7.0 database for these gases. The same emission factor from *IPCC* (2006) is used for indirect N<sub>2</sub>O from atmospheric deposition of N from NH<sub>3</sub> and NOx emissions, as for agricultural emissions (*Janssens-Maenhout et al.*, 2019).

369

The uncertainties for EDGAR N<sub>2</sub>O emissions estimated by *Solazzo et al.* (2021) are based primarily on the uncertainties in emissions factors and activity data statistics from the *IPCC* (2006). Globally, these emissions are accurate within an interval of  $\pm 113$  for energy, -12% to  $\pm 16\%$  for industrial processes and product use, -225 to  $\pm 302$  for agriculture, -159% to 203% for waste and  $\pm 112\%$  for others; the most uncertain emissions are those related to N<sub>2</sub>O from waste and agriculture.

375

# 376 SI-4 The UNFCCC inventory (need description of UNFCCC)

377 The UNFCCC collects detailed data on GHG emissions from its parties. Following extensive guidance

developed by IPCC (Buendia et al., 2019; Eggleston et al., 2006), parties to the convention prepare national

379 GHG inventories, including emissions (and sinks) of N<sub>2</sub>O. All anthropogenic activities are covered, in

380 agriculture both direct and indirect N<sub>2</sub>O emissions are included. While IPCC basically provides emission

- 381 factor approaches, parties are encouraged to take account of national specificities, use national factors and
- 382 data, wherever available, or develop emission models, with adequate scientific proof provided.

Combustion-related emissions and emissions from industrial processes may take advantage of emission monitoring or specific plant operation conditions, if provided. Emission processes that are not associated

- 385 with anthropogenic activities are also not covered in the inventories.
- 386

387 Obligations and quality of data provided differ strongly by country category. High scrutiny is put on GHG 388 inventories from countries listed in Annex-I of the convention (Annex-I countries include most European 389 countries, U.S. and Canada, Australia and New Zealand, and Japan). Annex-I countries are obliged to 390 provide annual national inventories in considerable detail and have to be very transparent also in terms of 391 methodology used and underlying information. Each year, time-series of emissions and underlying data 392 since 1990 (in a few cases, alternative base years are used) up to the pre-previous year are freshly provided 393 in April each year (e.g., in April 2023 data up to the year 2021 had to be provided), leading to a 394 homogeneous data series. Reports and emission data are provided (to UNFCCC, and to all users from the 395 UNFCCC web site at https://unfccc.int/reports) in standardized format such that they can be transferred to 396 databases. National results are routinely being checked and evaluated by expert teams in form of specific 397 internal and external audits to assure data quality and consistency.

398

399 National information is highly relevant also for non-Annex I countries to the UNFCCC and is being 400 collected and distributed by UNFCCC as well. Requirements are much less stringent, however, as parties 401 are expected to provide data only according to their own capabilities and the support they get from other 402 countries. The so-called Biannual Update Reports are to be prepared every other year only. While in 403 principle following the same IPCC guidance, commitments to format, timing, and quality assessment are 404 by far less stringent, and the own ambition level of the respective party (country) may determine much of 405 the outcome. In any case, self-reporting of a country always also means the party is willing to take the 406 responsibility of the emissions reported.

407

408 The "EDGAR/UNFCCC" dataset used in this paper utilizes the database for Annex-1 countries for 409 emissions from fossil-fuel consumption, industrial processes, waste and wastewater, and merges with the 410 respective set derived from EDGARv7.0 for all remaining countries.

411

#### 412 SI-5 The SRNM model: Flux upscaling model

The SRNM model (Wang et al., 2020) was applied to simulate direct cropland-N<sub>2</sub>O emissions. In SRNM, N<sub>2</sub>O emissions were simulated from N application rates using a quadratic relationship, with spatially variable model parameters that depend on climate, soil properties, and management practices. The original version of SRNM was calibrated using field observations only from China (Zhou et al., 2015). In this study, we used the global N<sub>2</sub>O observation dataset to train it to create maps of gridded annual emission factors of N<sub>2</sub>O and the associated emissions at 5-minute resolution from 1901 to 2014(Cui et al., 2021). The gridded EF and associated direct cropland-N<sub>2</sub>O emissions are simulated based on the following equation:

420 421

424

$$E_{ijt} = \alpha_{ij}N_{ijt}^2 + \beta_{ij}N_{ijt} + \varepsilon_{ijt}, \ \forall i$$
(23)

423 where

$$\alpha_{ij} \sim N\left(\sum_{k} (x_k \lambda_{ijk}), \sigma_{ijk}^2\right), \ \beta_{ij} \sim N\left(\sum_{k} (x_k \phi_{ijk}), \sigma_{ijk}'^2\right)$$
(24)

$$\lambda_{ijk} \sim N(\boldsymbol{\mu}_{ijk}, \boldsymbol{\omega}_{ijk}^2), \ \boldsymbol{\phi}_{ijk} \sim N(\boldsymbol{\mu}_{ijk}', \boldsymbol{\omega}_{ijk}'^2), \ \boldsymbol{\varepsilon}_{ijt} \sim N(0, \tau^2)$$
(25)

426 and *i* denotes the sub-function of N<sub>2</sub>O emission (*i*=1, 2, ..., *I*) that applies for a sub-domain division W<sub>i</sub> of 427 six climate or soil factors, *j* represents the type of crop (*j*=1-2, 1 for upland crops and 2 for paddy rice), *k* is 428 the index of climate or soil factors (k=1-6, i.e., soil pH, clay content, SOC, BD, the sum of cumulative 429 precipitation and irrigation, mean daily air temperature). W<sub>i</sub> denotes a set of the range of multiple  $x_k$ .  $E_{ijt}$ 430 denotes direct N<sub>2</sub>O emission flux (kg N ha<sup>-1</sup> yr<sup>-1</sup>) estimated for crop type *j* in year *t* in the *i*th sub-domain,

431  $N_{ijt}$  is N application rate (kg N ha<sup>-1</sup> yr<sup>-1</sup>), and  $a_{ij}$  and  $b_{ij}$  are defined as summation of the product of  $x_k$  and  $l_{ijk}$ 432 over k. The random terms l and f are assumed to be independent and normally distributed, representing the

433 sensitivity of a and b to  $x_k$ . e is the model error. m and mc are the mean effect of  $x_k$  for a and b, respectively.

434 s, sc, w, wc, and a restandard deviations. Optimal sub-domain division, associated parameters mean

435 values and standard deviations were determined by using the Bayesian Recursive Regression Tree version

436 2 (BRRT v2), constrained by the extended global cropland-N<sub>2</sub>O observation dataset. The detailed 437 methodological approach of the BRRT v2 is described by Zhou et al (2015).

438

# 439 Global cropland N<sub>2</sub>O observation dataset

440 We aggregated cropland  $N_2O$  flux observation data from 180 globally distributed observation sites from 441 online databases, on-going observation networks, and peer-reviewed publications (Figure SI-1). Chamber-442 based observations were only included in this dataset. These data repositories are as follows: the 443 NitroEurope, CarbonEurope, GHG-Europe (EU-FP7), GRACEnet, TRAGnet, NANORP, and 14 meta-444 analysis datasets (Decock, 2014; Harris et al., 2014; Helgason et al., 2005; Hénault et al., 2005; Hickman 445 et al., 2014; Kim et al., 2013a; Kim et al., 2013b; Lehuger et al., 2011; Leppelt et al., 2014; Rochette and Janzen, 2005; Sacks et al., 2010; Shcherbak et al., 2014; Stehfest and Bouwman, 2006; Walter et al., 2015). 446 447 Four types of data were excluded from our analysis: (i) observations without a zero-N control for 448 background N<sub>2</sub>O emission, (ii) observations from sites that used controlled-release fertilizers or nitrification 449 inhibitors, (iii) observations not covering the entire crop growing season, (iv) observations made in 450 laboratory or greenhouse. We then calculated cropland-N<sub>2</sub>O emissions as the difference between observed 451 N<sub>2</sub>O emission (E) and background N<sub>2</sub>O emission (E<sub>0</sub>). Values of EF were estimated for each nonzero N 452 application rate ( $N_a$ ) as direct cropland-N<sub>2</sub>O emission divided by  $N_a$ : EF = (E - E<sub>0</sub>)/ $N_a$ . This yielded a 453 global dataset of direct cropland-N<sub>2</sub>O emissions, N-rate-dependent N<sub>2</sub>O EFs and fertilization records from 454 each site (i.e., 1,052 estimates for upland crops from 152 sites and 154 estimates for paddy rice from 28 455 sites), along with site-level information on climate, soils, crop type, and relevant experimental parameters. 456 Total numbers of sites and total measurements in the dataset were more than doubled those for previous 457 datasets of N<sub>2</sub>O EF. The extended global N<sub>2</sub>O observation network covered most of fertilized croplands, 458 representing a wide range of environmental conditions globally. For each site in our dataset, the variables 459 included four broad categories: N<sub>2</sub>O emissions data, climate data (cumulative precipitation and mean daily 460 air temperature), soil attributes (soil pH, clay content, SOC, BD), and management-related or experimental 461 parameters (N application rate, crop type). More details on global cropland N<sub>2</sub>O observation dataset can be

462 found in *Cui et al.* (2021).



Figure SI-1 Global observation dataset of  $N_2O$  EF for direct soil emissions. Green area indicates the harvested areas of all crops derived from the Earthstat. Sites are indicated in different colors for maize, wheat, rice, and other crops.

468

#### 469 Gridded input datasets:

470 The updated SRNM model was driven by many input datasets, including climate, soil properties, 471 agricultural management practices (e.g., fertilization, tillage, irrigation), as well as the historical distribution 472 of cropland. Cumulative precipitation and mean daily air temperature over the growing season were 473 acquired from the CRU TS V4.06 climate dataset (0.5-degree resolution) (Harris et al., 2014), where 474 growing season in each grid cell was identified following Sacks et al. (2010) The patterns of SOC, clay 475 content, BD, and soil pH were acquired from the HWSD v1.2 ((Berdanier and Conant, 2012), 1-km 476 resolution). Both climate and soil properties were re-gridded at 5-arc-minute spatial resolution using a first-477 order conservative interpolation widely used in the CMIP5 model intercomparison (Yang et al., 2017). The 478 annual cropland area at 5-arc-minute spatial resolution from 1961 to 2020 was obtained from the History 479 Database of the Global Environment (HYDE 3.2.1) (Goldewijk et al., 2017).

480

481 For fertilization, crop-specific N fertilizer inputs (including synthetic N fertilizers, crop residues and 482 manure), fertilizer types, and placement during 1961-2020 were obtained from Adalibieke et al., (2023). 483 The frequency (i.e., one or multiple times) of N fertilization were the same as *Cui et al.* (2021) and we 484 assumed that the frequency remained constant during the study period. For tillage, the fraction of tillage by 485 crop during 1961-2020 was obtained from Adalibieke et al., (2023), which was constructed with the country 486 and province (or state) level no-tillage area data during 1961-2020 and downscaled to grid with the method 487 of Porwollik et al. (2019). For irrigation, the History Database of the Global Environment (HYDE version 488 3.2) (Goldewijk et al., 2017) and the MIRCA2000 dataset (Portmann et al., 2010) were used to compile the 489 global crop-specific irrigation proportion data from year 1961 to 2020. Categories of cropland in HYDE 490 provided new distinctions with irrigated and rain-fed crops (upland crops, other than rice), irrigated and 491 rain-fed rice during 1960-2017. The national-level dataset of "Agricultural area actually irrigated" was 492 obtained from (FAO, 2022), which was used to scale the baseline year 2015 maps of irrigated area from 493 HYDE over the period 2016-2020. The area of irrigated upland crops from HYDE was first disaggregated 494 into 21 crops based on the irrigated proportion from MIRCA2000 for per grid cell. We assumed an even 495 share of irrigated area by each upland crop during the period 1961-2020, like MIRCA2000. Finally, the 496 crop-specific irrigated area was masked by reporting harvested area, then the irrigated proportion of each

497 crop can be calculated as the crop-specific irrigated area divided by the physical area of each crop. For rice,
498 we further divided irrigated rice into continuously and intermittently flooded systems as provided by *Cui*499 *et al.* (2021), and we assumed that the irrigation proportion was kept the same during the study period.

499 500

# 500 SI-6 Global N flow in aquaculture

502 We applied the IMAGE-GNM aquaculture nutrient budget model for shellfish and finfish (Bouwman et al., 503 2013; Bouwman et al., 2011) to calculate the nutrient flows in aquaculture production systems. These flows 504 comprise feed inputs, retention in the fish, and nutrient excretion. Individual species within crustaceans, 505 seaweed, fish and molluscs are aggregated to the International Standard Statistical Classification of Aquatic 506 Animals and Plants (ISSCAAP) groups (FAO, 2022), for which production characteristics are specified. 507 Feed and nutrient conversion rates are used for each ISSCAAP group to calculate the feed and nutrient 508 intake based on production data from FAO (FAO, 2020). Feed types include home-made aquafeeds and 509 commercial compound feeds with different feed conversion ratios that also vary in time due to efficiency 510 improvement; in addition, the model accounts for algae in ponds, that are often fertilized with commercial 511 fertilizers or animal manure, consumed by omnivore fish species like carp. A special case is the filter-512 feeding bivalves that filter seston from the water column, and excrete pseudofeces, feces and dissolved 513 nutrients. Based on production data and tissue/shell nutrient contents, the model computes the nutrient 514 retention in the fish. Using apparent digestibility coefficients, the model calculates outflows in the form of 515 feces (i.e., particulate nutrients) and dissolved nutrients. Finally, nutrient deposition in pond systems and 516 recycling are calculated. For computing the N<sub>2</sub>O emissions, we consider the amount of N released to the 517 environment, i.e., the difference between N intake and N in the harvested fish, which includes all the 518 nutrient excretion. Since in pond cultures part of that N is managed, we made the amount of N recycling 519 explicit, as well as ammonia emissions from ponds. This is to avoid double counting when computing N2O 520 emissions from crop production.

521

# 522 SI-7 Continental Shelves N<sub>2</sub>O fluxes

523 N<sub>2</sub>O emissions from the global ocean do not include the contribution from continental shelves and are added 524 here using the extended mask of Laruelle et al. (2017) to delineate the coastal ocean. This mask excludes 525 estuaries and inland water bodies, while its outer limit is set 300 km away from the shoreline. Within this 526 coastal ocean domain, gridded N<sub>2</sub>O emissions were calculated using one data-driven estimate and three 527 high-resolution model estimates from two distinct models, all interpolated on the same 0.25° x 0.25° grid. 528 Models and data are each covering different time-periods and only one climatology is provided, keeping 529 the original timespan of each product: 1988-2017 for the observation-based product that relied on a random-530 forest (RF) algorithm to interpolate N<sub>2</sub>O data (Yang et al., 2020) from the MEMENTO database (MEM-531 RF) (Kock and Bange, 2015), 1998-2018 for the estimate relying on the high-resolution configuration 532 (Berthet et al., 2019) of the global ocean-biogeochemical component of CNRM-ESM2-1 (CNRM-0.25°), 533 1998-2013 and 2006-2013 for the estimates relying on the ECCO-Darwin model running at 1/3° (ECCO-534 Darwin1) and  $1/6^{\circ}$  (ECCO-Darwin2), respectively. The resulting climatology can be considered as broadly 535 representative of the last 2-3 decades. Each product is further described as follows:

536 537 M

# **537 MEM-RF**

The N<sub>2</sub>O air-sea flux reconstruction by *Yang et al.* (2020) is based on a synthesis of over 158,000 observations of N<sub>2</sub>O mixing ratio, partial pressure, and concentration in the surface ocean from the MEMENTO database (<u>https://memento.geomar.de</u>) (*Kock and Bange*, 2015) and additional cruises (Dataset S1) (*Yang et al.*, 2020). N<sub>2</sub>O measurements are converted to surface N<sub>2</sub>O mixing ratio anomalies using observations from the NOAA atmospheric flask dataset (*Hall et al.*, 2007), and extrapolated to a 0.25degree resolution global monthly climatology using an ensemble of 100 random forest realizations. The

random forest algorithm predicts N<sub>2</sub>O mixing ratio anomalies based on their relationship to oceanographic

545 predictors that include hydrographic variables, nutrients, oxygen, chlorophyll, net primary production, and

- 546 seafloor depth. Reconstructed mixing ratio climatologies are used to estimate air-sea fluxes by applying a
- 547 commonly used gas exchange parameterization (*Wanninkhof*, 2014). Two formulations of piston velocity
- are adopted: one based on a quadratic dependence on wind speed (*Wanninkhof*, 2014), and one that
- 549 explicitly accounts for bubble-mediated fluxes (*Liang et al.*, 2013). Sea ice cover, surface temperature, 550 salinity and atmospheric pressure are taken from ERA5 reanalysis (*Hersbach et al.*, 2017). Calculations are
- 551 performed with two high-resolution wind products (ERA5 and CCMP) that are available at 0.25, 6-hourly
- resolution for the period from 1988 to 2017, yielding four permutations of the piston velocity. The resulting
- 553 ensemble of 400 global N<sub>2</sub>O air-sea flux estimates is averaged in time to obtain monthly mean
- 554 climatologies. A description of the dataset and methods is presented in *Yang et al.* (2020). The code used
- to produce these datasets is archived on a public GitHub repository at https://github.com/yangsi7/mapping-
- 556 ocean-n2o (DOI: 10.5281/zenodo.3757194). 557

# 558 CNRM-0.25°

559 N<sub>2</sub>O fluxes have been inferred from the global ocean-biogeochemical component of CNRM-ESM2-1 (Séférian et al., 2019) run at 0.25° horizontal resolution with 75 vertical levels in the ocean. This high-560 561 resolution configuration is described in Berthet et al. (2019) and is based on the NEMOv3.6 oceanic model 562 (Madec, 2008), the multi-category sea ice model GELATOV6 (Salas v Mélia, 2002) and the PISCESv2-gas 563 model for marine biogeochemistry (Aumont et al., 2015), which includes an updated version of (Martinez-564 Rey et al., 2015) for the marine N<sub>2</sub>O module. The simulation was first spun-up during 300 years under 565 preindustrial conditions and then has been forced by the OMIP2-compliant JRA55-do-1-5 atmospheric reanalysis (Tsujino et al., 2020; Tsujino et al., 2018) considering the historical evolution of CO<sub>2</sub> and N<sub>2</sub>O 566 567 in the atmosphere since the year 1850. Boundary conditions for nitrogen deposition and riverine inputs are 568 prescribed from monthly climatologies. The suboxic production of N<sub>2</sub>O uses the oxygen-dependent 569 formulation of Jin and Gruber (2003) and is enhanced at low oxygen concentrations. This formulation 570 encompasses N<sub>2</sub>O production during remineralization, nitrification and grazing, as well as a sink term 571 corresponding to N<sub>2</sub>O consumption under anoxic conditions by denitrification. The oceanic N<sub>2</sub>O partial 572 pressure is computed based on the surface N<sub>2</sub>O concentration and the N<sub>2</sub>O solubility in the ocean. Sea-to-573 air N<sub>2</sub>O fluxes are then computed using the standard gas exchange parameterization of *Wanninkhof* (1992; 574 2014). 575

# 576 ECCO-Darwin & ECCO2-Darwin

577 To generate global air-sea fluxes of nitrous oxide  $(N_2O)$  from the global ocean we have used the ECCO-578 Darwin Model (Carroll et al., 2020). The ECCO-Darwin model is based on MITgcm and it has a nominal 579 horizontal resolution of 1/3 of a degree with 50 vertical levels where in the top 100 meters the model is 580 vertically resolved with 10-meter grid boxes. The ECCO-Darwin model is forced with an atmospheric 581 forcing corresponding to the 1992-present optimized with adjoint technique in order to realistically 582 represent the observed physical climate phenomena such as El Nino, the Pacific Decadal Oscillation, the 583 North Pacific "Warm Blob", etc. A more detailed description of the model forcing and the Darwin 584 biogeochemical model configuration used in this study can be found in Carroll et al. (2020).

585

586 The Darwin biogeochemical/ecological model (Carroll et al., 2020; Manizza et al., 2019) used for this 587 study carries 33 biogeochemical tracers to explicitly represent the cycle of carbon, oxygen, phosphorus, 588 silica, and iron in the global ocean. For this particular version of the model, we implemented a 589 parameterization of the oceanic cycle of  $N_2O$  using the scheme of Nevison et al. (2003) based on the oceanic 590 oxygen cycle previously represented in ECCO2-Darwin model (Ganesan et al., 2020). The air-sea gas flux 591 of N<sub>2</sub>O was parameterized according to Wanninkhof (1992). In addition, a thermal- only N<sub>2</sub>O tracer (a tracer 592 in which biogeochemical sources and sinks are suppressed but with the same solubility as  $N_2O$ ) was also 593 added to the model to isolate the process of ocean ventilation affecting the N<sub>2</sub>O concentration in the ocean

594 at seasonal time scales as done in *Manizza et al.* (2012). The ECCO-Darwin numerical simulation was run

for the 1992-2014 period, but we discarded the inclusion of the output relative to the 1992-1996 period in our analysis due to the model adjustment in this initial part of our numerical simulation.

597

# 598 SI-8 Open Ocean N<sub>2</sub>O fluxes

599 N<sub>2</sub>O is produced in the open ocean by microbial activity during organic matter cycling in the subsurface 600 ocean, and its production pathways are influenced by the local environmental oxygen level. In the oxic ocean N<sub>2</sub>O is produced as a byproduct during the oxidation of ammonia to nitrate, mediated by ammonia 601 602 oxidizing bacteria and archaea. N<sub>2</sub>O is also produced and consumed in sub- oxic and anoxic waters through 603 the action of marine denitrifiers during the multi-step reduction of nitrate to gaseous N. The oceanic  $N_2O$ 604 distribution therefore displays significant heterogeneity with background levels of 10-20 nmol/l in the well-605 oxygenated ocean basins, high concentrations (> 40 nmol/l) in hypoxic waters, and N<sub>2</sub>O depletion in the 606 core of ocean oxygen minimum zones (OMZs).

607

For this synthesis open ocean  $N_2O$  emissions to the atmosphere were compiled from four global ocean biogeochemistry models/Earth System models that simulate the production, consumption and circulation of oceanic  $N_2O$  (Table 6).  $N_2O$  flux exchange between ocean and atmosphere is derived using gas-exchange parameterizations applied to modeled surface ocean  $N_2O$ . Versions of the four submitting models also participated in the previous  $N_2O$  budget synthesis (*Tian et al.*, 2020a). Model details and updates to the

- 613 previous N<sub>2</sub>O budget synthesis are summarized below.
- 614 The models differ in aspects of physical configuration (e.g., spatial resolution), meteorological forcing 615 applied at the ocean surface, and in their parameterizations of ocean biogeochemistry; specific details on
- 616 individual models are provided in the publications listed in Table 1. Towards this N<sub>2</sub>O budget synthesis,
- 617 modelling groups reported grid-resolved ( $1^{\circ}\times1^{\circ}$  horizontal resolution) monthly estimates of ocean-
- 618 atmosphere N<sub>2</sub>O fluxes for the period 1980-2020 (or for as many years as possible in that period).
- 619

# 620 U. Bern: Bern-3D

621 N<sub>2</sub>O fluxes are derived from the Bern-3D Earth System Model of Intermediate Complexity which includes 622 a prognostic marine biogeochemistry model (based on (Parekh et al., 2008) and (Tschumi et al., 2011)). 623 Configuration of the model for simulation of  $N_2O$  is outlined in *Battaglia and Joos* (2018). Model 624 simulations were run at a native resolution of horizontal resolution of 41 by 40 grid cells and 32 625 logarithmically scaled vertical layers. Modifications of the biogeochemistry model relevant for the  $N_2O$ 626 cycle include the assignment of organic matter remineralization to aerobic and anaerobic pathways 627 dependent on mean grid-cell dissolved oxygen level. N2O is produced by nitrification as a product of 628 remineralization rate and a specified N<sub>2</sub>O yield which has a functional form dependent on oxygen level (see 629 details in (Battaglia and Joos, 2018)). N<sub>2</sub>O consumption by denitrification processes is represented by a 630 first-order kinetics formulation which also includes a dependence on local oxygen level to account for the 631 relative importance of denitrification-related N<sub>2</sub>O production and consumption processes in each gridcell. 632 Measurements of dissolved N<sub>2</sub>O (surface and interior) from the MEMENTO database (*Kock and Bange*, 633 2015) together with an ensemble of model runs are used to constrain the model parameters governing N<sub>2</sub>O 634 production and consumption mechanisms. From a pre-industrial equilibrium state the model is forced by 635 historical CO2 emissions, non-CO2 radiative forcing, and land-use changes. N<sub>2</sub>O in the atmosphere is 636 prescribed based on historical data.

637

# 638 CNRM: CNRM-ESM2-1

639 N<sub>2</sub>O fluxes are provided by the CNRM-ESM2-1 Earth System model. The ocean model component is based

on NEMO Version 3.6 (*Madec et al.*, 2017) and coupled to the GELATO sea ice model (*Salas y Mélia*,

641 2002) Version 6 and the marine biogeochemical model PISCESv2-gas (Aumont et al., 2015). The spatial

- model resolution follows the eORCA1L75 grid, with a nominal horizontal resolution of 1° and with higher
- 643 resolution in the tropics (increasing to  $\sim (1/3)^{\circ}$ ). The model has 75 vertical levels with higher resolution

towards the ocean surface. The simulations were forced at the surface by the atmospheric state of JRA55-

645 do v1.5.0 (*Tsujino et al.*, 2018). Atmospheric N2O concentration is given as annual means as specified by

646 CMIP6 protocols and is linearly interpolated in time. Parameterization of the marine N<sub>2</sub>O cycle follows that 647 of *Martinez-Rev et al.* (2015) with some modifications. N<sub>2</sub>O production is driven by an oxygen-dependent

647 of *Martinez-Rey et al.* (2015) with some modifications. N<sub>2</sub>O production is driven by an oxygen-dependent 648 yield of N<sub>2</sub>O, which encompasses production from denitrification and nitrification processes. This

649 formulation also assumes a constant background yield representing N<sub>2</sub>O production by nitrification and a

650 consumption of N<sub>2</sub>O in suboxic conditions. Originally implemented by *Martinez-Rev et al.* (2015), the

- marine  $N_2O$  parameterization has benefited from a recoding and an improved calibration presented in
- 652 *Berthet et al.* (2023). Further details of the model biogeochemistry and configuration are provided by 653 *Séférian et al.* (2019) and *Berthet et al.* (2019).
- 654
- 655 UVic2.9

656 N<sub>2</sub>O model fluxes are derived from the UVic2.9, Earth System Model of Intermediate Complexity with

657 prescribed monthly climatological winds (*Kalnay et al.*, 1996) and ice sheets (*Peltier*, 2004), configuration 658 outlined in *Landolfi et al.* (2017). Oceanic subsurface N<sub>2</sub>O production is parameterized following (*Zamora* 

659 *and Oschlies*, 2014), as a function of  $O_2$  consumption with a linear  $O_2$  dependency, inherently including 660 both nitrification and denitrification. In  $O_2$ -deficient waters (<4 mmol m<sup>-3</sup>), denitrification becomes a sink

661 of N<sub>2</sub>O consumed at a constant rate. The gradient driving the air-sea N<sub>2</sub>O gas exchange, is computed online 662 based on departure of the surface ocean concentration from the saturation value using the solubility

663 coefficients of *Weiss and Price* (1980) and time-varying prescribed atmospheric N<sub>2</sub>O concentrations

664 (historical and RCP8.5). The model was spun-up for 6000 years with preindustrial boundary conditions

- 665 (solar and volcanic and aerosol forcing, fixed atmospheric CO<sub>2</sub> of 280 ppm and N<sub>2</sub>O of 276 ppb, and
- 666 667

# 668 UEA: NEMO-PlankTOM10.2

preindustrial atmospheric N deposition).

669 N<sub>2</sub>O model fluxes are derived from the NEMO-PlankTOM10.2 ocean model. The physical circulation 670 component is NEMO v3.1 (Madec, 2008), with horizontal resolution of 2° longitude, and a variable 671 latitudinal resolution (average  $\sim 1^{\circ}$ ) with higher resolution in the tropics and polar regions. The model has 672 30 vertical layers, with variable resolution ranging from 10m in the upper 100m to 500m at depths of 5000 673 m. The biogeochemical component relies on the marine ecosystem model PlankTOM10, which includes 674 representation of 10 plankton functional types (Le Quéré et al., 2016). It has been extended by Buitenhuis 675 et al. (2018) to include nitrogen cycle processes, and prognostic and diagnostic models of  $N_2O$  production. 676 N<sub>2</sub>O is produced from nitrification and denitrification pathways, with oxygen dependent yields employed 677 to account for varying production and consumption processes in hypoxic waters. Nitrogen cycle parameters 678 are optimized using ocean databases of dissolved N2O (MEMENTO, Kock and Bange (2015)) nitrification 679 rates (Yool et al., 2007), and surface ammonium concentrations (Johnson et al., 2015; Paulot et al., 2015). 680 Further details on model configuration are provided in (Buitenhuis et al., 2018).

681

# 682 SI-9 Net N<sub>2</sub>O emission from land cover change

683 This section mainly involves the calculation of post-deforestation  $N_2O$  emissions, deforestation induced 684  $N_2O$  reduction and  $N_2O$  emissions from forest regrowth (afforestation or reforestation). The methods 685 include both bookkeeping and process-based modeling.

686

# 687 a. Deforestation area, crop/pasture expansion and secondary forests

The LUH2 v2h (land use harmonization, http://luh.umd.edu) land use data was used to derive the deforestation area and its partition between crops and pastures during 1860–2020. LUH2 categorizes forest lands into forested primary land and potentially forested secondary land, while croplands are divided into

- 691 C3 annual crops, C3 perennial crops, C4 annual crops, C4 perennial crops, and C3 N-fixing crops.
- 692

693 In the empirical computation of deforestation induced N<sub>2</sub>O emissions, all sub-classes within each land use 694 type were treated the same. Thus, only the annual transition area from forests to croplands or managed 695 pasture was needed. In the process-based estimates, the DLEM model was improved to further account for 696 the classifications of primary forests, secondary forests (further partitioned into established and newly 697 converted by an age threshold of 15 years), croplands/pastures /rangelands (further partitioned into 698 established and newly converted by an age threshold of seven nine years). Each land use type can be divided 699 into several different plant functional types (PFTs). Specifically, within a grid cell, cropland can only be 700 dominated by only one crop type. The pastures and rangelands can be either C3 type or C4 type. To generate 701 the historical spatial distribution of PFTs, a potential vegetation map and the accompanied composition 702 ratio map of each natural PFT acquired from the Synergetic Land Cover Product (SYNMAP) were jointly 703 used with LUH2 v2h. 704

#### 705 **b. Methods**

706 The bookkeeping method was mainly applied to the tropical areas, where forests generally have high  $N_2O$ 707 emissions. Specifically, the average tropical forest N<sub>2</sub>O emission rate of 1.974 kg N<sub>2</sub>O-N ha-1 yr-1 was 708 adopted as the baseline. Two logarithmic response curves of soil N2O emissions (normalized to the baseline) after deforestation were developed:  $y = -0.31 \ln(x) + 1.53$  and  $y = -0.454 \ln(x) + 2.21$ . This form of the 709 response functions can effectively reproduce the short-lived increase in soil N<sub>2</sub>O emissions after initial 710 711 forest clearing and the gradually declining emission rates of converted crops/pastures (Melillo et al., 2001; 712 Verchot et al., 1999). Using these two curves and the baseline, we kept track of the N<sub>2</sub>O reduction of tropical 713 forests and the post-deforestation crop/pasture N<sub>2</sub>O emissions at an annual timescale.

714

715 There are not many studies on N<sub>2</sub>O emissions from secondary tropical forests that regrowth after crop or 716 pasture abandonment. Sullivan et al. (2019) lumped together all forms of N "gas loss" including NO and 717 N<sub>2</sub>O for secondary forests across the tropics and the results showed gas loss gradually increases with age 718 since the regrowth of secondary forest. Keller and Reiners (1994) also reported a gradual recovery of soil 719 nitrate and soil emissions of N<sub>2</sub>O and nitric oxide (NO) during 20 years of secondary forest succession, 720 which however did not return to the level of the primary forests. In this study, using field observations from 721 Davidson et al. (2007) and Keller and Reiners (1994), we regressed normalized N<sub>2</sub>O emissions (relative to 722 a reference mature forest) of secondary tropical forests with their ages as y=0.0084x + 0.2401 (R<sup>2</sup> = 0.44; 723 unit of x is year). The derived y values were multiplied by tropical forest  $N_2O$  emissions estimated by 724 NMIP2 models that do not distinguish secondary forests from primary forests.

724

The DLEM model was run with varying climate and CO<sub>2</sub> with other factors held constant to estimate forest baseline emissions and unfertilized crop/pasture emissions from 1860-2020. The climate data were acquired from CRUJRA, which is a fusion of the CRU and JRA reanalysis products at a spatial resolution of  $0.5^{\circ} \times$ 0.5° and a daily time-step. The atmospheric CO<sub>2</sub> data were obtained from NOAA GLOBLVIEW-CO2 dataset (https://www.esrl.noaa.gov), which are derived from atmospheric and ice core measurements. In the tropical area, both estimates from the DLEM model and the bookkeeping method were adopted, whereas in extra-tropical area, we only adopted the DLEM outputs.

733

## 734 SI-10 Inland water, estuaries, and coastal vegetation

735

# 736 a. Dynamic Land Ecosystem Model-Terrestrial/Aquatic Continuum (DLEM-TAC)

To quantify N<sub>2</sub>O emissions from global inland waters (rivers, lakes, and reservoirs), we use a process-based
coupled terrestrial-aquatic model, which builds up on the Dynamic Land Ecosystem Model (DLEM).
DLEM-TAC is a fully distributed, process-based land surface model which couples the major land
processes (terrestrial hydrology, plant phenology and physiology, soil biogeochemistry) and aquatic
dynamics (lateral transport and in-stream biogeochemistry) (*Pan et al.*, 2021; *Tian et al.*, 2015; *Tian et al.*,

742 2020b; Yao et al., 2020). The land component of DLEM-TAC explicitly simulates the carbon, nitrogen,

and water fluxes between plants, soil, and atmosphere, and the surface and drainage runoff and nitrogen
load from the land module are used as input for the aquatic module. The simulated nitrogen load
includes dissolved inorganic nitrogen (DIN), dissolved organic nitrogen (DON), particulate organic
nitrogen (PON), and runoffs, sewers as the initial condition of the aquatic module.

747

748 DLEM-TAC aquatic module calculated lateral water transport and the associated aquatic biogeochemical 749 processes by adopting a scale-adaptive scheme. The water transport scheme, which coupled hillslope flow, 750 subnetwork flow, and main channel flow, simulated the water transport processes within grid cells. In the 751 aquatic module, lakes and reservoirs were linked with small streams and large rivers, forming a river-lake-752 reservoir corridor (Wollheim et al., 2008)). Particularly, lakes that are linked to small streams are typically 753 small in size and are defined as small lakes, while those linked to large rivers are usually had large size and 754 are defined as large lakes; similarly, reservoirs that are linked to main channels are considered as large 755 reservoirs, while those that are linked to small streams are considered as small reservoirs. The incoming 756 flow of a linked river-lake-reservoir corridor drains to lakes first, and the outflow rate of lakes and reservoirs 757 is determined based on the predefined residence time obtained from the global lake dataset (Lehner et al., 758 2011; Messager et al., 2016; Yao et al., 2022). The aquatic N module was developed based on the scale 759 adaptive water transport scheme, including lateral transport, decomposition of organic matter, particle 760 organic matter deposition, nitrification, and denitrification. The detailed description could be found in the 761 previous studies (Pan et al., 2021; Tian et al., 2020b; Yao et al., 2020).

762

Following our previous work referring to the development of water transport and biogeochemistry, we developed an inland water N<sub>2</sub>O module within the aquatic biogeochemical component of the DLEM framework (*Yao et al.*, 2020). The net fluxes of dissolved N<sub>2</sub>O (including physical and biogeochemical processes) in the main channel (high-order streams) and subnetwork (small rivers) are estimated as:

(26)

$$(\Delta M N2O) / \Delta t = Fa + Ywater + D - R - E$$

where M\_N2O is the total mass of dissolved N<sub>2</sub>O in the main channel or subnetworks (g N),  $\Delta t$  is the time step, Fa is advective N<sub>2</sub>O fluxes (g N d<sup>-1</sup>), Ywater is the N<sub>2</sub>O production within the water column (g N d<sup>-1</sup>), D is the dissolved N<sub>2</sub>O from rainfall to rivers (i.e. deposition) (g N d<sup>-1</sup>) with an initial concentration equal to the atmospheric equilibrium N<sub>2</sub>O concentration, R is the flux from N<sub>2</sub>O reduction (g N d<sup>-1</sup>) to nitrogen gas, and E is the riverine N<sub>2</sub>O efflux (g N d<sup>-1</sup>) through the air-water interface.

773

774 Input data. The driving data of DLEM-TAC include the climate variables, atmospheric CO2 concentration, 775 land use change, nitrogen (N) deposition, N fertilizer, and manure application with a spatial resolution of 776  $0.5^{\circ} \times 0.5^{\circ}$ . The daily climate variables (precipitation, mean temperature, maximum temperature, minimum 777 shortwave radiation) were obtained from the CRUNCEP temperature. and dataset 778 (https://vesg.ipsl.upmc.fr) for 1901-2019. Climate data of each year during 1850-1900 was randomly 779 sampled from 1901-1920. Annual atmospheric CO2 concentration from 1900-2019 was obtained from the 780 NOAA GLOBALVIEW-CO2 dataset (https://www.esrl.noaa.gov). The annual land use change data was 781 derived from a potential natural vegetation map (synergetic land cover product) and a prescribed cropland 782 area dataset from the history database of the global environment v.3.2 (HYDE 3.2, ftp://ftp.pbl.nl/hyde). 783 The data of N fertilizer, manure N application, and N deposition data was obtained from (*Tian et al.*, 2022). 784

- In the aquatic module, the required channel dataset included channel slope, channel width, and channel length generated from the Hydroshed dataset (*Lehner et al.*, 2008) and DDM30 dataset (*Döll and Lehner*, 2002). The flow direction and distance data were obtained from the Dominant River Tracing (DRT) dataset. For modeling water dynamics in lakes and reservoirs, we generated 0.5 grid level surface water area,
- vpstream area, volume, depth, and average residence time for lakes based on the Hydrolakes dataset

790 (*Messager et al.*, 2016), while the GRanD database provided the same information for reservoirs (*Lehner et al.*, 2011).

792

793 Simulation protocol. DLEM-TAC simulations include three steps: equilibrium run, spin-up run and two 794 transit runs, one with dam operation close, and another one with dam operation open. First, the equilibrium 795 run is required to obtain the initial and steady condition of carbon, nitrogen, and water pool at the pre-796 industrial level in each grid cell (Thornton and Rosenbloom, 2005). In this step, we held all the driving 797 forces such as climate data, atmospheric CO<sub>2</sub> concentration, land use data, and nitrogen additions consistent 798 with the first year's data we used in the simulation. Second, we conducted a 30-year spin-up run by 799 randomly selecting climate data within the 1850s (*Tian et al.*, 2012a). This step can alleviate the disturbance 800 of driving data changes in the transit run. Then we conduct the natural flow simulation with the dam model 801 temporarily closed, and all the driving forces change over time. After the natural flow simulation, we set 802 up a management flow simulation with the dam module open, specifically the dam module needs natural 803 flow in the previous run as model input.

# 805b. Mechanistic Stochastic Modeling of N2O emissions from large rivers, lakes, reservoirs, and806estuaries:

807

804

808 To calculate the cascading loads of TN and TP delivered to each water body along the river-reservoir-809 estuary continuum, we spatially routed all reservoirs from the GRanD database (Lehner et al., 2011), with 810 river networks from Hydrosheds 15s (Lehner et al., 2008) and, at latitudes above 50°N, Hydro1K 811 (http://edc.usgs.gov/products/elevation/gtopo30/hydro/), which were in turn connected to estuaries as 812 represented in the "Worldwide Typology of Nearshore Coastal Systems" of Dürr et al. (2011). In addition, 813 the global database HydroLAKES (Messager et al., 2016) was used to topologically connect 1.4 million lakes with a minimum surface area of 0.1 km<sup>2</sup> within the river network. Note that besides natural lakes, 814 815 HydroLAKES includes updated information on 6,796 reservoirs from the GRanD database, which was used 816 in the study of Maavara et al. (2019). In order to estimate the TN and TP loads to each water body, we then 817 relied on a spatially explicit representation of TN and TP mobilization from the watershed into the river 818 network (see (Maavara et al., 2019) for details (Bouwman et al., 2009; Van Drecht et al., 2009).

819

820 For the estimation of N<sub>2</sub>O emission, we applied two distinct model configurations, respectively named DS1 821 and DS2 in Maavara et al. (2019). DS1 estimates N2O emissions from denitrification and nitrification based 822 on an EF of 0.9%, which is in the mean of published values (Beaulieu et al., 2011), and the assumption that 823  $N_2O$  production equals  $N_2O$  emissions (*Maavara et al.*, 2019). For DS2, the reduction of  $N_2O$  to  $N_2$  during 824 denitrification if N<sub>2</sub>O is not evading sufficiently rapidly from the water body is considered. The fluxes in 825 the model represent lumped sediment-water column rates and were resolved at the annual timescale. The 826 use of water residence time as an independent variable in both the mechanistic model and the upscaling 827 process introduces an important kinetic refinement to existing global N<sub>2</sub>O emission estimates. Rather than 828 applying an average EF (directly scaling N<sub>2</sub>O emissions to N inputs) to all water bodies, the use of water 829 residence time explicitly adjusts for the extent of N<sub>2</sub>O production and emission that is kinetically possible 830 within the timeframe available in a given water body. Simulated N<sub>2</sub>O emission rates were evaluated against 831 UNFCCC measurement-based upscaling methods applied to reservoirs (Deemer et al., 2016) and rivers 832 (Hu et al., 2016) as well as a UNFCCC observation-driven regional estimate of lake N<sub>2</sub>O emissions based 833 on literature data (Lauerwald et al., 2019).

834

## 835 c. Meta analysis-based N<sub>2</sub>O emissions from large rivers

B36 Data from 70 published studies between 1998 and 2016 that provided N<sub>2</sub>O emission from streams and rivers

837 were compiled by *Hu et al.* (2016). The N<sub>2</sub>O emission factors (EF = N<sub>2</sub>O /DIN) and emission rates (ER =  $\frac{1}{2}$ 

838 EF \* DIN load, kg N<sub>2</sub>O-N yr<sup>-1</sup>) were calculated for each studied river. Exploratory multiple regression 839 analyses were conducted to predict EF and ER using various combinations of factors (N concentrations, loads, yields, DOC: DIN, discharge, and watershed area) and different functions. Among them, DIN yield (kg N yr<sup>-1</sup> km<sup>-2</sup>) was identified as the best predictor of EF and ER. Using the optimal model and DIN loads for 6400 global rivers calculated by the NEWS2-DIN-S model (*McCrackin et al.*, 2014), we estimated global riverine N<sub>2</sub>O emissions (*Hu et al.*, 2016).

844

#### 845 d. Stream and river N<sub>2</sub>O emissions combining machine-learning and model-based upscaling

846 Marzadri et al. (2021) developed a novel approach that combines a data-driven Random Forest Machine 847 Learning (RM-ML) model with a physically-based upscaling model to predict near global (neglecting Arctic and Antarctic areas) riverine N<sub>2</sub>O emissions flux (F\*N<sub>2</sub>O given by the ratio between the flux of N<sub>2</sub>O, 848 849 FN<sub>2</sub>O, and the in-stream flux of dissolved inorganic nitrogen species FDIN) from both surface (i.e. water 850 column) and subsurface (i.e. benthic zone and hyporheic zone) riverine environments. In particular, to 851 capture the local scale processes responsible for N<sub>2</sub>O emissions and to provide estimations at different 852 spatial scales (from local reach up to the global scale) the model compute two different denitrification 853 Damköhler numbers (given by the ratio between a characteristics time of transport and a characteristics 854 time of denitrification (Marzadri et al., 2021; Marzadri et al., 2017)) starting from the hydro-morphological 855 and biogeochemical information provided by the RM-ML model. Model results at the local reach scale 856 shows that nearly 50% of the riverine N<sub>2</sub>O emissions comes from small streams (i.e. widths lower than 10 857 m, although they represent only the 13% of the total riverine surface area worldwide) while at the large 858 scale predict a near-global annual riverine N<sub>2</sub>O emissions around 72.8 GgN<sub>2</sub>O-N/yr.

859

#### 860 e. Meta-analysis based N<sub>2</sub>O emissions from estuaries and coastal vegetation

861 N2O emissions from estuaries and coastal vegetated ecosystems were obtained from a meta-analysis of 862 observed N<sub>2</sub>O fluxes (Rosentreter et al., 2023). In brief, the meta-data analysis relies on a categorization of 863 estuaries into 'tidal systems and deltas', 'lagoons', and 'fjords'. Water-air N<sub>2</sub>O fluxes from 123 estuary 864 sites globally were then compiled from peer-reviewed publications until the end of 2020. Coastal vegetation 865 comprises 'mangrove', 'salt marsh', and 'seagrass' ecosystems and N<sub>2</sub>O sediment-water-air fluxes were 866 compiled from 55 sites globally from peer-reviewed publications until the end of 2020. A non-parametric 867 bootstrapping method (1000 iterations of the median of samples) was used to resample N<sub>2</sub>O fluxes per unit 868 area for each estuary and coastal vegetation type in each of the 18 regions using the 'boot' function in the 869 package 'boot' in R software. Results from the bootstrapping output (minimum, Q1, median, mean, Q3, 870 maximum) were then scaled to the surface area of each estuary and coastal vegetation type in each of the 871 18 RECCAP regions. If an ecosystem type had less than three sites in a region, we applied the global 872 statistics of this type in this region. Note that the meta-data analysis only provides flux assessments at the 873 scale of the 18 regions.

874

#### 875 SI-11 Atmospheric inversion models

876 Emissions were estimated using four independent atmospheric inversion frameworks (see Table 1). The 877 frameworks all used a Bayesian inversion method. The approach used here finds the maximum posteriori 878 (MAP), or optimal, estimate of emissions, that is, those, which when coupled to a model of atmospheric 879 transport, provide the best agreement to observed  $N_2O$  mixing ratios while being guided by their prior 880 probability. In this particular case, where both the likelihood and prior probability are assumed to be 881 distributed normally, the optimal emissions are equivalent to those that minimize the cost function,

882

$$J(\mathbf{x}) = \frac{1}{2} \left( \mathbf{x} - \mathbf{x}_{\mathbf{b}} \right)^{\mathrm{T}} \mathbf{B}^{-1} \left( \mathbf{x} - \mathbf{x}_{\mathbf{b}} \right) + \frac{1}{2} \left( \mathbf{y} - H(\mathbf{x}) \right)^{\mathrm{T}} \mathbf{R}^{-1} \left( \mathbf{y} - H(\mathbf{x}) \right)$$
(27)

883 884

885 where **x** and  $\mathbf{x}_b$  are, respectively, vectors of the multivariate means of the posterior and prior emission 886 distributions, **B** is the prior error covariance matrix for emissions, **y** is a vector of observed N<sub>2</sub>O mixing 887 ratios, **R** is the observation error covariance matrix, and  $H(\mathbf{x})$  is the model of atmospheric transport (for 888 details on the inversion method see (*Tarantola*, 2005)). The optimal emissions,  $\mathbf{x}$ , were found by solving 889 the first order derivative of equation (21):

- 890
- 891

891 
$$J'(\mathbf{x}) = \mathbf{B}^{-1} (\mathbf{x} - \mathbf{x}_{b}) + (H'(\mathbf{x}))^{\mathrm{T}} \mathbf{R}^{-1} (\mathbf{y} - H(\mathbf{x})) = 0$$
892 (28)

893 where  $(H'(\mathbf{x}))^{\mathrm{T}}$  is the sensitivity of the atmospheric observations to emissions, derived from an adjoint 894 model of transport. In frameworks INVICAT, PyVAR-CAMS and GEOS-Chem, equation (5b) was solved 895 using a variational approach (Thompson et al., 2014; Wells et al., 2015; Wilson et al., 2014), which uses a 896 descent algorithm and computations involving the forward and adjoint models. In framework MIROC4-897 ACTM (Patra et al., 2018), equation (22) was solved directly by computing a transport operator, H from integrations of the forward model, such that Hx is equivalent to H(x), and taking the transpose of H (*Patra* 898 899 et al., 2022).

900

901 Each of the inversion frameworks used a different model of atmospheric transport with different horizontal 902 and vertical resolutions (see Table 1). The transport models TOMCAT and LMDz5, used in INVICAT and PvVAR-CAMS respectively, were driven by ECMWF ERA-5 and ERA-Interim wind fields respectively, 903 904 MIROC4-ACTM was driven by JRA-55 wind fields, and GEOS-Chem was driven by MERRA-2 wind 905 fields. While INVICAT, PyVAR-CAMS, and GEOS-Chem optimized the emissions at the spatial 906 resolution of the transport model, MIROC4-ACTM optimized the error in the emissions aggregated into 84 907 land and ocean regions. All frameworks optimized the emissions with monthly temporal resolution. The 908 transport models included an online calculation of the loss of N<sub>2</sub>O in the stratosphere due to photolysis and 909 oxidation by  $O(^{1}D)$  resulting in mean atmospheric lifetimes of between 118 and 129 years, broadly 910 consistent with recent independent estimates of the lifetime of 116±9 yr (Prather et al., 2015)).

911

912 All inversions used N<sub>2</sub>O measurements of discrete air samples from the National Oceanic and Atmospheric 913 Administration Carbon Cycle Cooperative Global Air Sampling Network (NOAA). In addition, discrete 914 measurements from the Commonwealth Scientific and Industrial Research Organisation network (CSIRO) 915 as well as in-situ measurements from the Advanced Global Atmospheric Gases Experiment network 916 (AGAGE), the NOAA CATS network, and from individual sites operated by University of Edinburgh (UE), 917 National Institute for Environmental Studies (NIES), the Finnish Meteorological Institute (FMI) and the 918 Japan Meteorological Agency (JMA) were included in INVICAT, PyVAR-CAMS and GEOS-Chem. 919 Measurements from networks other than NOAA were corrected to the NOAA calibration scale, NOAA-920 using the results of the WMO Round Robin inter-comparison experiment 2006A. 921 (https://www.esrl.noaa.gov/gmd/ccgg/wmorr/), where available. For AGAGE and CSIRO, which did not 922 participate in the WMO Round Robins, the data at sites where NOAA discrete samples are also collected 923 were used to calculate a linear regression with NOAA data, which was applied to adjust the data to the 924 NOAA-2006A scale. For the remaining CSIRO sites where there were no NOAA discrete samples, the 925 mean regression coefficient and offset from all other CSIRO sites were used. The inversions used the 926 discrete sample measurements without averaging, and hourly or daily means of the in-situ measurements, 927 depending on the particular inversion framework.

928

929 Each framework applied its own method for calculating the observation space uncertainty, the square of 930 which gives the diagonal elements of the observation error covariance matrix R. The observation space 931 uncertainty accounts for measurement and model representation errors and is equal to the quadratic sum of 932 these terms. Typical values for the observation space uncertainty were between 0.3 and 0.5 ppb for all 933 inversion frameworks.

934

935 Prior mean emissions were based on estimates from terrestrial biosphere and ocean biogeochemistry models 936 as well as from inventories. INVICAT, PyVAR-CAMS and GEOS-Chem used the same prior estimates for

937 emissions from natural and agricultural soils from the model OCN v1.1 (Zaehle et al., 2011) and for biomass 938 burning emissions from GFEDv4.1s. For non-soil anthropogenic emissions (namely those from energy, 939 industry and waste sectors), INVICAT, PyVAR-CAMS, and GEOS-Chem used EDGAR v5. MIROC4-940 ACTM used the VISIT model (Inatomi et al., 2010; Ito et al., 2018) for emissions from natural soils and 941 EDGAR 4.2 for all anthropogenic emissions, including agricultural waste burning, but did not explicitly 942 include a prior estimate for wildfire emissions.

943

944 For the prior mean estimate of ocean fluxes, INVICAT, PyVAR-CAMS and GEOS-Chem used the 945 prognostic version of the PlankTOM-v10.2 model (Buitenhuis et al., 2018) with a global total source 2.5 946 TgN yr<sup>-1</sup>. Prior uncertainties were estimated in all the inversion frameworks for each grid cell (INVICAT, 947 PyVAR-CAMS and GEOS-Chem) or for each region (MIROC4-ACTM) and the square of these 948 uncertainties formed the diagonal elements of the prior error covariance matrix **B**. INVICAT, PvVAR-949 CAMS and GEOS-Chem estimated the uncertainty as proportional to the prior value in each grid cell, but 950 MIROC4-ACTM set the uncertainty uniformly for land regions at 1 Tg N yr<sup>-1</sup> and for ocean regions at 0.5 951 Tg N vr<sup>-1</sup>. INVICAT also included off-diagonal covariances in B corresponding to a spatial correlation 952 between flux uncertainties of 500 km and assumed a semi-exponential distribution of uncertainties so as to 953 restrict the possibility of negative fluxes.

954

#### 955 956 SI-12 Atmospheric N<sub>2</sub>O Observation Networks

957 The NOAA Network: For atmospheric  $N_2O$  observations from the NOAA network (Dutton et al. 2023), we 958 used global mean mixing ratios from the NOAA Global Monitoring Laboratory (GML) (combined dataset 959 based on measurements from five different measurement programs [HATS old flask instrument, HATS 960 current flask instrument (OTTO), the Carbon Cycle and Greenhouse Gases (CCGG) group Cooperative 961 Global Air Sampling Network (https://www.esrl.noaa.gov/gmd/ccgg/flask.php), HATS in situ (RITS 962 program), and HATS in situ (CATS program)]. CCGG provides uncertainties with each measurement (see 963 site files: <u>ftp://aftp.cmdl.noaa.gov/data/greenhouse\_gases/n2o/flask/surface/</u>). The CCGG measurements 964 for N<sub>2</sub>O analysis from more than 50 sites globally was changed to tunable infrared laser direct absorption 965 spectroscopy (TILDAS) in mid-2019 from gas chromatography. About 40 sites of them (mostly Marine 966 Boundary Laver sites) are used to calculate CCGG monthly mean global N<sub>2</sub>O levels. Monthly mean 967 observations from different NOAA measurement programs are statistically combined to create a long-term NOAA/ESRL GML dataset. Uncertainties (1 sigma) associated with monthly estimates of global mean 968 969 N<sub>2</sub>O, are ~1 ppb from 1977–1987, 0.6 ppb from 1988–1994, 0.3–0.4 ppb from 1995–2000, and 0.1 ppb 970 from 2001-2017. NOAA data are generally more consistent after 1995, with standard deviations on the 971 monthly mean mixing ratios at individual sites of ~0.5 ppb from 1995–1998, and 0.1–0.4 ppb after 1998. 972 A detailed description of these measurement programs and the method to combine them are available via 973 https://www.esrl.noaa.gov/gmd/hats/combined/N2O.html.

974 THE AGAGE network: The Advanced Global Atmospheric Gases Experiment (AGAGE) global network 975 (and its predecessors ALE and GAGE) (Prinn et al., 2018) has made continuous high-frequency gas 976 chromatographic (GC) measurements with electronic capture detection (ECD) of  $N_2O$  at five globally 977 distributed sites since 1978. Improved GC/ECD methods have been deployed over time resulting in N<sub>2</sub>O 978 measurement precision of about 0.35% in ALE, 0.13% in GAGE (Prinn et al., 1990) and 0.05% in AGAGE 979 (Prinn et al., 2008; 2018). We used the global mean of AGAGE N<sub>2</sub>O measurements during 1980–2020 980 which are reported on the Scripps Institution of Oceanography SIO-16 scale. Further information on 981 AGAGE stations, instruments, calibration, uncertainties and access to data is available at the AGAGE Data 982 website: https://www.osti.gov/dataexplorer/biblio/dataset/1841748.

983

984 The CSIRO network: The CSIRO flask network (Francey et al., 2003) consists of nine sampling sites 985 distributed globally and has been in operation since 1992. Flask samples are collected approximately every 986 two weeks and shipped back to CSIRO GASLAB for analysis. Samples were analyzed by gas 987 chromatography with electron capture detection (GC-ECD). One Shimadzu gas chromatograph labelled 988 "Shimadzu-1" (S1) has been used over the entire length of the record and the measurement precision for

- 989 N<sub>2</sub>O from this instrument is about 0.1%. N<sub>2</sub>O data from the CSIRO global flask network are reported on
- 990 the NOAA-2006A N<sub>2</sub>O scale and are archived at the World Data Centre for Greenhouse Gases (WDCGG:
- 991 https://gaw.kishou.go.jp/). Nine sites from the CSIRO network were used to calculate the annual global
- 992 N<sub>2</sub>O mole fractions. Smooth curve fits to the N<sub>2</sub>O data from each of these sites were calculated using the 993
- technique outlined in Thoning et al. (1989), using a short-term cut-off of 80 days. The smooth curve fit data 994 were then placed on an evenly spaced latitude (5 degree) versus time (weekly) grid using the Kriging
- 995 interpolation technique. Finally, the gridded data were used to calculate the global annual values.
- 996
- 997 Table SI-3 Factors used to convert  $N_2O$  in various units (by convention Unit 1=Unit 2 × conversion)

Unit 1	Unit 2	Conversion
Tg N <sub>2</sub> O (teragrams of N <sub>2</sub> O)	Tg N (teragrams of nitrogen)	1.57
Tg N (teragrams of nitrogen)	g N (grams of nitrogen)	10 <sup>-12</sup>
Tg N (teragrams of nitrogen)	ppb (parts per billion)	4.79

- 999 Table SI-4 Atmospheric N<sub>2</sub>O dry mole fraction measured by different observing networks during 2000-
- 1000 2022.

ppb	NOAA	AGAGE	CSIRO	Min	Max
2000	315.58	316.18	315.48	315.48	316.18
2001	316.33	316.95	316.12	316.12	316.95
2002	316.99	317.54	316.67	316.67	317.54
2003	317.64	318.26	317.31	317.31	318.26
2004	318.24	318.99	317.99	317.99	318.99
2005	318.98	319.71	318.83	318.83	319.71
2006	319.93	320.39	319.58	319.58	320.39
2007	320.59	321.16	320.34	320.34	321.16
2008	321.54	322.11	321.45	321.45	322.11
2009	322.24	322.91	322.22	322.22	322.91
2010	323.04	323.77	323.08	323.04	323.77
2011	324.21	324.68	324.09	324.09	324.68
2012	325.01	325.65	324.99	324.99	325.65
2013	325.92	326.61	325.89	325.89	326.61
2014	327.06	327.66	326.93	326.93	327.66
2015	328.13	328.52	327.99	327.99	328.52
2016	328.94	329.36	328.77	328.77	329.36
2017	329.75	330.37	329.68	329.68	330.37

2018	330.87	331.53	330.90	330.87	331.53
2019	331.85	332.35	331.66	331.66	332.35
2020	333.06	333.48	332.70	332.70	333.48
2021	334.33	334.81	334.03	334.03	334.81
2022	335.71	336.09	335.57	335.57	336.09

1002

1003 Table SI-5: Uncertainty in future projections of atmospheric N<sub>2</sub>O dry mole fraction.

ppb	SSP	1-1.9	SSP	1-2.6	SSP2	2-4.5	SSP.	3-7.0	SSP	4-3.4	SSP	4-6.0	SSP	5-8.5
Year	Min	Max												
2020	330.4	331.1	330.4	331.1	331.0	331.6	331.4	332.0	331.2	331.5	331.2	331.4	331.2	331.9
2030	335.1	336.9	335.5	337.0	337.6	339.6	339.5	342.2	337.5	338.6	338.8	339.6	339.5	341.1
2040	336.2	341.1	336.8	342.0	343.2	347.3	347.9	353.4	340.5	345.7	346.3	349.2	349.2	350.7
2050	336.2	344.6	337.8	345.7	348.5	354.3	356.1	364.9	343.3	353.3	353.5	359.2	359.4	361.2

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1005	<b>References:</b>
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