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Seasonal Variability of Tropical Wetland CH₄ emissions: the role of the methanogen-available carbon pool

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Abstract

We develop a dynamic methanogen-available carbon model (DMCM) to quantify the role of the methanogen-available carbon pool in determining the spatial and temporal variability of tropical wetland CH_4 emissions over seasonal timescales. We fit DMCM parameters to satellite observations of CH_4 columns from SCIAMACHY CH_4 5 and equivalent water height (EWH) from GRACE. Over the Amazon river basin we find substantial seasonal variability of this carbon pool (coefficient of variation = 28 ± 22 %) and a rapid decay constant ($\phi = 0.017 \, \text{day}^{-1}$), in agreement with available laboratory measurements, suggesting that plant litter is likely the prominent methanogen carbon source over this region. Using the DMCM we derive global CH₄ emissions for 10 2003–2009, and determine the resulting seasonal variability of atmospheric CH_4 on a global scale using the GEOS-Chem atmospheric chemistry and transport model. First, we estimate tropical emissions amount to 111.1 Tg CH₄ yr⁻¹ of which 24 % is emitted from Amazon wetlands. We estimate that annual tropical wetland emissions have increased by $3.4 \text{ Tg CH}_4 \text{ yr}^{-1}$ between 2003 and 2009. Second, we find that the model 15

¹⁵ Creased by 3.4 Fg CH₄ yr ¹⁶ between 2003 and 2009. Second, we find that the model is able to reproduce the observed seasonal lag between CH_4 concentrations peaking 1–3 months before peak EWH values. We also find that our estimates of CH_4 emissions substantially improve the comparison between the model and observed CH_4 surface concentrations (r = 0.9). We anticipate that these new insights from the DMCM represent a fundamental step in parameterising tropical wetland CH_4 emissions and guantifying the seasonal variability and future trends of tropical CH_4 emissions.

1 Introduction

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Wetlands are the single largest source of methane (CH₄) into the atmosphere and account for 20–40 % of the global CH₄ source (Denman et al., 2007), of which tropical wetlands account for 50–60 % of this global wetland CH₄ source (e.g. Cao et al., 1996; Bloom et al., 2010). Tropical wetland biogeochemistry is poorly understood compared



to boreal peatlands (Mitsch et al., 2010), resulting in large inter-model discrepancies of the magnitude and distribution of tropical wetland CH₄ emission estimates (Riley et al., 2011). Tropical climate variability (e.g., resulting in widespread droughts, Lewis et al., 2011) can lead to large year to year variations in tropical wetland CH₄ emissions and subsequently the global CH₄ budget (Hodson et al., 2011). An improved quantitative

⁵ Subsequently the global CH₄ budget (Hodson et al., 2011). An improved quantitative understanding of the magnitude, distribution, and variation of tropical wetland CH_4 emissions is therefore essential to further understanding of the global CH_4 cycle. Here, we parameterise tropical wetland CH_4 emissions, and hence introduce a predictive capability that can be used to determine future emissions and to help quantify global CH_4 climate feedbacks.

In wetlands and rice paddies, methanogenesis (the biogenic production of CH_4) occurs as the final step of anoxic organic matter decomposition (Neue et al., 1997). Factors influencing methanogenesis rates include substrate availability, soil pH, temperature, water table position and redox potential (Whalen, 2005). Wetland vegetation type

- and aquatic herbivore activity can also affect the transport of CH_4 between the soil and atmosphere (Joabsson et al., 1999; Dingemans et al., 2011). On a global scale, seasonal variations in wetland CH_4 fluxes are mostly determined by temporal changes in wetland water volume and soil temperature (Walter et al., 2001; Gedney et al., 2004), and from seasonal changes in wetland extent and wetland water table depth (Ringeval
- et al., 2010; Bloom et al., 2010). Recent work that used SCIAMACHY lower tropospheric CH₄ column concentrations and Gravity Recovery And Climate Experiment (GRACE) equivalent water height (EWH) retrievals shows that the seasonality of wetland CH₄ emissions can be largely explained by seasonal changes in surface temperature and water volume (Bloom et al., 2010). The Amazon and Congo river basins
 were the only major exceptions in this study, where CH₄ concentrations peaked sev-
- eral weeks before EWH, highlighting our incomplete understanding of the processes controlling tropical wetland CH_4 emissions over seasonal timescales.

In this paper we focus on the seasonal lag between CH_4 emissions and flooding over the Amazon river basin area (Oki and Sud, 1998). We use SCIAMACHY CH_4 retrievals



and GRACE EWH (both described in Sect. 2.2) to determine the seasonal lag between wetland CH₄ emissions and wetland water volume. Figure 1 shows that seasonal flooding of the Amazon basin occurs typically 1–3 months after the peak CH₄ concentrations, and to a lesser extent the lag persists throughout tropical wetland areas. In

- Sect. 2, we test the hypothesis that this lag is related to the depletion of methanogenavailable carbon during the onset of the tropical wet season by explicitly accounting for this carbon pool in a parameterised model of tropical wetland CH₄ emissions (Bloom et al., 2010). We optimise model parameters by fitting them to SCIAMACHY CH₄ column and GRACE EWH measurements, and use the resulting model to estimate global
- wetland emission estimates. In Sect. 3 we compare our results to previous estimates of wetland CH₄ emissions and to decomposition rates of methanogen-available carbon in anaerobic environments. Finally, we use our estimated emissions to drive the GEOS-Chem atmospheric chemistry model as an approach to test the consistency between our emission estimates and observed variations of atmospheric CH₄ concentration. We appelude the paper in Cent. 4
- ¹⁵ We conclude the paper in Sect. 4.

2 Process-based model and application

Here, we introduce a methanogen-available carbon pool (C_{μ}) that typically originates from labile plant litter, recalcitrant organic matter decomposition and root exudates (e.g. Wania et al., 2010). Typically soil carbon pool decay constants are more than an order of magnitude lower than those of leaf litter (Sitch et al., 2003; Wania et al., 2010). Therefore, if C_{μ} originates mostly from the slow-decomposing recalcitrant carbon pool, then variations in C_{μ} over seasonal timescales are likely to be small. Conversely, if C_{μ} is drawn from leaf litter, then large variations in C_{μ} abundance may arise as a result of rapid litter decomposition in the tropics. Miyajima et al. (1997) measured CH₄ accumulation of anaerobic decomposition of incubated tropical withered tree leaves over a 200 day period. These observations show a rapid decrease in decomposition rates over the incubation period. Bianchini Jr. et al. (2010) found similar results from



dried and ground anaerobic decomposition of *Oxycaryum cubense* at 20°C: following a 20-day lag (where no emissions were observed) CH_4 produced from organic carbon decomposition peaked after a 50-day period, and then rapidly decreased. On a tropical river-basin scale, flooded areas expand at the onset of the wet season and engulf newly available plant litter: as a result, CH_4 emissions from plant litter may peak before the height of the water table. The occurrence of anaerobic CH_4 emissions from litter decomposition within sub-seasonal timescales raises the question as to whether C_{μ} significantly varies in time.

2.1 Model description

¹⁰ We base our model on previous work (Bloom et al., 2010) that describes the temporal variability of wetland emissions $F_{CH_4}^t$ (mg CH₄ m⁻² day⁻¹) as a function of EWH and surface temperature:

$$F_{CH_{a}}^{t} = k(\Gamma_{w}^{t} + D_{\alpha})Q_{10}(T_{s}^{t})^{\frac{T_{s}^{t}}{10}},$$
(1)

where at time *t* (days), Γ_w^t is the EWH, T_s^t is the surface temperature (K), D_α is the equivalent depth of the wetland soil (m), $Q_{10}(T_s^t)$ is the temperature dependence function implemented by Gedney et al. (2004), and *k* is a scaling constant (mg $CH_4 m^{-2} day^{-1}$) accounting for all temporally constant factors (e.g. Gedney et al., 2004).

Equation (1) assumes an inexhaustible source of methanogen-available carbon. ²⁰ Here we account for the potential seasonal changes in C_{μ} by substituting k with $\phi_0 C_{\mu}^t$, where ϕ_0 (day⁻¹) is the temperature, water and carbon independent decay constant of wetland methanogenesis, and C_{μ}^t is the value of C_{μ} (mg CH₄ m⁻²) at time *t*:

$$F_{CH_4}^t = \phi_0 C_{\mu}^t (\Gamma_w^t + D_{\alpha}) Q_{10} (T_s)^{\frac{T_s^t}{10}}.$$



(2)

To determine temporal changes in C_{μ} , we define C_{μ}^{t+1} in terms of C_{μ}^{t} :

$$C^{t+1}_{\mu} = C^t_{\mu} + N_{\mu} \Delta t - F^t_{CH_4} \Delta t,$$

where Δt is the time interval, $F_{CH_4}^t$ is the carbon loss due to emitted CH₄ (Eq. 2), N_{μ} is the net influx of carbon available for methanogenesis from plant litter, root exudates, and breakdown of complex polymers from the recalcitrant carbon pool. We assume N_µ is temporally constant, and we assume wetland carbon stocks are in quasi-equilibrium on annual timescales, hence $\overline{N_{\mu}} = \overline{F_{CH_4}^t}$. Note that when ϕ_0 is small, the equilibrium $C_{\mu} \gg N_{\mu}\Delta t$. In this case, $C_{\mu}^{t+1} \simeq C_{\mu}^t$ and Eq. (2) converges to Eq. (1) (Bloom et al., 2010), which assumes $\phi_0 C_{\mu}$ is constant over seasonal timescales. In order to compare derived decay constants with observed and model values (e.g. Miyajima et al., 1997; Wania et al., 2010), we determine the annual mean decay constant of wetlands areas as $\overline{\phi} = \overline{F_{CH_4}^t} / \overline{C_{\mu}^t}$ (day⁻¹). Equations (2) and (3) constitute the dynamic methanogenavailable carbon model (DMCM).

2.2 Data

- ¹⁵ For the sake of brevity we only include a brief description of the datasets for our analysis and refer the reader to dedicated papers. Solar backscatter data from the Scanning Imaging Absorption Spectrometer for Atmospheric Cartography (SCIAMACHY) instrument onboard Envisat is used to retrieve the mean column concentrations of CH₄ in the atmosphere (Frankenberg et al., 2005). The spatial resolution of CH₄ retrievals is
- 30 km × 60 km, and the Envisat orbital geometry ensures global coverage at 6-day intervals. CH₄ retrievals are only achievable in daytime cloud-free conditions. The Gravity Recovery and Climate Experiment (GRACE) is a twin satellite system from which the Earth's gravity field is retrieved at 10-day intervals. Tides, atmospheric pressure and wind are included in the applied corrections on GRACE gravity retrievals: the remaining temperatural unstation in CRACE.
- ²⁵ maining temporal variation in GRACE gravity is dominated by terrestrial water variability



(3)

(Tapley et al., 2004). We incorporate SCIAMACHY CH_4 concentrations, GRACE EWH and NCEP/NCAR daily $1.9^{\circ} \times 1.88^{\circ}$ temperature re-analyses (Kalnay et al., 1996) into a process-based model following Bloom et al. (2010). We use the 2003–2008 SCIA-MACHY column CH_4 retrievals (Frankenberg et al., 2008), and the CNES GRACE EWH $1^{\circ} \times 1^{\circ}$ 10-day resolution product (Lemoine et al., 2007): we aggregate all three datasets to a daily $3^{\circ} \times 3^{\circ}$ horizontal grid (see Bloom et al., 2010).

2.3 Global parameter optimisation

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We implement the DMCM on a global 3° × 3° grid for the period 2003–2009. We drive the DMCM using the aggregated daily values of T_s^t and Γ_w^t . We spin up the DMCM using 2003 T_s^t and Γ_w^t values until it reaches an annual equilibrium ($\overline{N_{\mu}} = \overline{F_{CH_4}^t}$). In contrast to Bloom et al. (2010), we supplement the $Q_{10}(T_s)$ function with a gradual linear cut-off for temperatures for 0°C < T_s^t < -10°C, and when T_s^t < -10°C, $F_{CH_4}^t$ = 0 as a first order approximation to wintertime CH₄ emission inhibition in boreal wetlands. As the Q_{10} function never reaches zero, this supplementary constraint will effectively suppress winter-time CH₄ emissions, which is broadly consistent with our current understanding of CH₄ emissions in boreal wetlands.

We apply the DMCM globally in order to determine (i) the magnitude of ϕ and C_{μ} in the tropics within each 3° × 3° gridcell (ii) the potential of C_{μ} temporal variability on extra-tropical wetland environments, and (iii) CH₄ emissions from wetlands and rice paddies at a global scale. We determine the global distribution and seasonal variability of wetland CH₄ emissions by optimising parameters ϕ_0 and D_{α} at each gridcell by minimising the following cost function (*J*):

$$J = \sum_{t=1}^{n} (\kappa * \Delta F_{\mathsf{CH}_4}^t - \Delta S_{\mathsf{CH}_4}^t)^2,$$

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(4)

where $\Delta S_{CH_4}^t$ denotes the SCIAMACHY CH₄ variability after we remove the interannual trend (represented as a 2nd order polynomial); $F_{CH_4}^t$ is derived from Eqs. (2) and (3); and the conversion factor κ (ppm kg⁻¹ CH₄ m⁻² day⁻¹) relates CH₄ emissions to the equivalent column concentration in the lower troposphere (e.g. Bloom et al., 2010).

- ⁵ We then implement the global $Q_{10}(T_s)$ optimisation approach of Bloom et al. (2010). Like other top-down parameter optimisation methods of global wetland CH₄ emissions (Gedney et al., 2004; Bloom et al., 2010), our method is unable to distinguish between the seasonality of CH₄ emissions from wetlands and rice paddies due to the concurring fluxes over seasonal timescales, although we anticipate varying fertilisation and irriga-
- tion practices will also influence the seasonality in rice paddy CH₄ emissions (Conen et al., 2010). We hence distinguish the sources spatially (Bloom et al., 2010) for which we have more confidence in the distribution of rice paddies. Finally, we use the IPCC global wetland and rice paddy CH₄ emissions median of 227.5 Tg CH₄ yr⁻¹ (Denman et al., 2007) as a base value for 2003 emissions.

¹⁵ We propagate the following uncertainties through our global wetland and rice paddy CH₄ emissions estimation (Bloom et al., 2010): (i) SCIAMACHY CH₄ observation errors; (ii) the uncertainty of the linear fit between $F_{CH_4}^t$ and $S_{CH_4}^t$; (iii) the uncertainty $\sigma_{\kappa} = \pm 16\%$ associated with κ ; and (iv) a global wetland and rice paddy uncertainty of $\pm 58 \text{ Tg CH}_4 \text{ yr}^{-1}$ (Denman et al., 2007).

20 3 Results and discussion

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Over the Amazon river basin we find wetland CH₄ fluxes coinciding with small values of C_{μ} , resulting in a highly variable C_{μ} over seasonal timescales. Assuming an annual mean inundated fraction of 3.3% (Prigent et al., 2007), the median CH₄ flux over a flooded area is 1.06 Mg C ha⁻¹ (369 mg CH₄ m⁻² day⁻¹). The median Amazon wetland $C_{\mu} = 0.16$ Mg C ha⁻¹ with a range of 0.02–7.89 Mg C ha⁻¹ (5th–95th percentile). The large spatial variability of C_{μ} is consistent with the complexity of methanogenesis rates



in wetlands (Neue et al., 1997; Whalen, 2005). Large temporal changes of C_{μ} are observed in the Amazon river basin where the mean C_{μ} coefficient of variation (c_{ν}) is $28 \pm 22\%$ over the period 2003–2009. When we allow C_{μ} to vary in extra-tropical regions we find a median of $c_{\nu} < 0.1\%$, and as a result the relatively small C_{μ} variability

- ⁵ does not influence the seasonality of CH₄ emissions outside the tropics. For rice paddy areas in southeast Asia we find a median of $c_v = 4.8$ %. We acknowledge that due to the varying rice cultivation practices around the world (Conen et al., 2010), the effects of rice paddy irrigation and the timing of fertilisation on C_{μ} cannot be captured by the DMCM approach.
- ¹⁰ To determine whether our derived values for C_{μ} and $\overline{\phi}$ are relevant to tropical ecosystems, we compare them against laboratory measurements of anaerobic decomposition of withered leaves from a wetland region in Narathiwat, Thailand (Miyajima et al., 1997). We simulate CH₄ production from C_{μ} at each model gridcell for a 200-day period without fresh carbon input (N_µ=0), and we use innundated fraction observations (Prigent ¹⁵ et al., 2007) to determine the flux magnitude over flooded areas only. Figure 2 shows the cumulative CH₄ production over a 200-day period for (i) simulated decomposition
- from derived $\overline{\phi}$ and C_{μ} values over the Amazon, (ii) simulated decomposition derived $\overline{\phi}$ and C_{μ} values over boreal wetlands, and (iii) upscaled withered leaf mineralisation rates by Miyajima et al. (1997) using a median of 17.5 Mg C ha⁻¹ fine and coarse woody debris (Malhi et al., 2009). For boreal and tropical C_{μ} decomposition, the median cumulative CH₄ emissions, 68 % confidence interval, and mean decay constants ($\overline{\phi}$) are shown. For the withered leaf mineralisation rates, we show the mean fitted decay constant ($\overline{\phi}$) and the range and median cumulative CH₄ emissions.

The top-down parameter estimation of $\overline{\phi}$ and C_{μ} suggest plant litter C_{μ} is a funda-²⁵ mental component of tropical CH₄ emission seasonality. Our top-down estimation of anaerobic decomposition rates for tropical wetland CH₄ emissions compare favourably with laboratory measurements of anaerobically produced CH₄: while the magnitude of tropical C_{μ} decomposition is more than a factor of two smaller than laboratory



measurements (Miyajima et al., 1997), the mean decay constant $\overline{\phi}_{Amazon} = 0.017 \text{ day}^{-1}$ compares well to $\overline{\phi}_{leaf} = 0.011 \text{ day}^{-1}$ for withered leaf decomposition. The larger laboratory measurements (Miyajima et al., 1997) are partially explained by an incubation temperature of 35 °C (cf. a mean surface temperature in the Amazon basin of 23 °C), and the lack of observations for coarse woody debris decomposition. As a result of relatively high $\overline{\phi}$ values, measured leaf decomposition and model CH₄ emissions both show a significant reduction of CH₄ emission rates throughout the 200-day period. In contrast, the boreal decay constant ($\overline{\phi}_{Boreal} = 0.0003 \text{ day}^{-1}$) indicates relatively constant CH₄ emission rates throughout the 200-day period.

¹⁰ Table 1 shows a comparison between observed and model decay constants derived from a variety of methods. The range of $\overline{\phi}_{Amazon}$ values are within the order of magnitude of leaf and wetland macrophyte decay constants (Miyajima et al., 1997; Longhi et al., 2008; Wania et al., 2010). We believe that $\overline{\phi}_{Amazon}$ is an indicator for the cumulative decay constant of the rapid anaerobic decomposition of root exudates, plant litter decomposition, and the contribution of recalcitrant carbon pools. For a more detailed $\overline{\phi}_{Amazon}$ comparison with observed and model decay constant values, an estimation of the overall $\overline{\phi}$ in wetland CH₄ production from bottom-up process-based models (e.g. Wania et al., 2010) is needed.

Figure 3 shows the total CH₄ flux over the central branch of the Amazon river (0° N– 6° S, 80° W–40° W). The temporal changes in C_{μ} result in a significantly different timing for CH₄ emissions over the tropics in comparison to the Bloom et al. (2010) water volume and temperature dependence approach. While in the dry season the minimum CH₄ fluxes coincide with the lowest GRACE EWH, peak CH₄ fluxes occur during the rising water phase. The DMCM optimisation predicts that the accumulation of carbon in the dry season results in higher C_{μ} values at the onset of the wet season. This carbon pool is then rapidly depleted during the wet season. As a result, CH₄ emission within the water column has been proposed as a mechanism explaining reduced CH₄



emissions during the peak of the wet season (Mitsch et al., 2010), although this would result in a second CH_4 peak at the end of the wet season. The absence of this peak in our analysis suggests this process plays only a minor role in the seasonality of tropical wetland CH_4 emissions.

By globally integrating the DMCM method we estimated tropical wetlands emit 5 111.1 Tg CH₄ yr⁻¹, where Amazon wetlands account for 26.2 Tg CH₄ yr⁻¹ (24%). Table 2 shows our estimates are within the range of other independent Amazon wetland emission CH_4 estimates. Figure 4 shows the zonal profile of our top-down approach with the associated uncertainty estimates. We capture three main features of global wetland and rice paddy emissions, i.e. peaks over the tropics, subtropics and lower 10 mid-latitudes (mainly due to rice), and boreal latitudes, in agreement with previous studies (Bloom et al., 2010; Fung et al., 1991; Riley et al., 2011). In comparison to our previous work (Bloom et al., 2010) we find a slight reduction in boreal wetland emissions (3.2%), primarily due to the introduction of a gradual cut-off in methanogenesis rates under 0°C (Sect. 2.3). During 2003–2008, the global change in CH_4 15 wetland emissions amounts to an increase of 7.7 Tg CH₄ yr⁻¹, mostly as a result of boreal wetlands (3.1 Tg CH₄ yr⁻¹) and tropical wetlands (3.4 Tg CH₄ yr⁻¹), while there is also a significant increase of 1.1 Tg CH_4 yr⁻¹ from mid-latitude wetlands. The increase in Southern Hemisphere extra-tropical wetland emissions $(0.13 \text{ Tg CH}_{4} \text{ yr}^{-1})$ did not significantly contribute to the CH_4 wetland emissions growth during 2003–2008. 20

Finally, we use our wetland and rice CH₄ emission estimates to drive the GEOS-Chem global 3-D atmospheric chemistry and transport model (described and evaluated by Fraser et al., 2011) allowing us to test consistency between our emissions to surface measurements of CH₄ concentrations. We sample the model at the time and geographical location of the surface CH₄ measurements from the GasLab, AGAGE and ESRL networks (Francey et al., 1996; Prinn et al., 2000; Cunnold et al., 2002; Dlugokencky et al., 2009). Figure 5 shows model and observed CH₄ concentration anomalies (i.e., minus the mean trend) for the northern and Southern Hemispheres. We chose to remove the interannual trend from all CH₄ concentrations to compare the



seasonality of model and surface measurements of CH₄. We show that the DMCM approach better describes the observed seasonality in both hemispheres ($r_{NH} = 0.9$, $r_{SH} = 0.9$), and the amplitude of the Southern Hemisphere seasonality is largely improved in comparison to the GEOS-Chem runs using Fung et al. (1991) and Bloom ⁵ et al. (2010) CH₄ emissions.

4 Concluding remarks

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Understanding the temporal controls of temperature, water volume and carbon content of wetlands is crucial in determining the global and regional seasonal cycle of wetland CH_4 emissions. We show that incorporating a temporally variable methanogenavailable carbon pool, C_{μ} , in our top-down approach results in a significant improvement in describing the temporal behaviour of tropical and global CH_4 emissions.

By implementing our dynamic methanogen-available carbon model (DMCM) on a global scale we determine the effects of a seasonally variable C_{μ} on the seasonality of wetland CH₄ emissions in the Amazon river basin. We find a median decay constant

- ¹⁵ of $\overline{\phi}_{Amazon} = 0.017 \text{ day}^{-1}$ over the Amazon river basin. Seasonal changes in C_{μ} in the tropics largely explain the seasonal lag between SCIAMACHY observed CH₄ concentrations and GRACE equivalent water height. The relatively high seasonal variability in C_{μ} (mean $c_{\nu} = 28$ %) over the Amazon river basin results in peak CH₄ emissions occurring mostly 1–3 months prior to the peak water height period: in contrast, the median
- ²⁰ boreal C_{μ} variability is $c_{\nu} < 0.1$ %. We show a substantial improvement in simulating surface concentrations when using the GEOS-Chem ACTM with our wetland and rice CH₄ emission estimates (r = 0.9). These improvements in the magnitude and temporal dynamics of tropical CH₄ emissions will ultimately help constrain global inverse modelling efforts.
- We anticipate that this work will lead to further and more detailed parameterisation of tropical wetland CH_4 emissions, and we expect our tropical wetland CH_4 emission parameterisation will reduce the uncertainty in forecasting future changes in wetland CH_4 emissions.



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Table 1. Model and observed decay constants for organic matter decomposition in anaerobic environments.

	Decay Constant (yr ⁻¹)	Study
Amazon Wetlands ($\overline{\phi}_{Amazon}$)	2.6-9.6 ^ª (median=5.9)	This Study : Top-down wetland CH_4 emission parameter optimization
Withered Leaves (35 °C)	4.0	Miyajima et al. (1997) : Decay constant from anaerobic tropical leaf CH ₄ mineralisation
Wetland Macrophyte Decomposition	1.0–5.5	Longhi et al. (2008) ^b : Measured decomposition rates in Paluda di Ostiglia, Italy
Soil Carbon Pool (10°C) Leaf Litter (10°C) Root Exudates (10°C)	0.001–0.03 0.35 13	Wania et al. (2010) : Bottom-up CH_4 Emissions from Northern Peatlands

^a 68 % confidence interval

^b Mass-loss decomposition rates



Discussion Paper BGD 9, 387-409, 2012 Seasonal variability of tropical wetland CH₄ emissions **Discussion** Paper A. A. Bloom et al. **Title Page** Abstract Introduction Conclusions References **Discussion** Paper Tables Figures 14 Back Close **Discussion** Paper Full Screen / Esc **Printer-friendly Version** Interactive Discussion

Table 2. Estimates of total annual Amazon river basin wetland CH_4 emissions (Tg CH_4 yr⁻¹).

Study	Amazon Wetland CH_4 Emissions (Tg CH_4 yr ⁻¹)
Melack et al. (2004) Fung et al. (1991) Riley et al. (2011) Bloom et al. (2010)	22 5.3 58.9 ^a 20.0
This study	26.2±9.8

^a High tropical fluxes by Riley et al. (2011) are a result of anomalously high predicted net primary productivity in the Community Land Model (CLM version 4)



Fig. 1. Top: The timing (day of year) of peak CH_4 concentrations from SCIAMACHY (left), peak equivalent water height (EWH) from GRACE (middle), and the peak CH_4 concentration lead over tropical South America (right). Bottom: Normalised anomaly of GRACE EWH, mean flood fraction (Prigent et al., 2007) and mean CH_4 concentrations (including 1-standard deviation envelope) over the main branch of the Amazon river (0°–6° S, 40°–80° W).











Fig. 3. Daily wetland CH₄ emissions for 2003–2009 (blue) and GRACE equivalent water height (green) over the central branch of the Amazon river ($0^{\circ}-6^{\circ}$ S, $40^{\circ}-80^{\circ}$ W).





Fig. 4. Zonal profile of CH_4 emissions from wetlands and rice paddies: top-down approach (blue); Fung et al. (1991), wetlands only (red); Riley et al. (2011) wetland and rice paddy emissions (orange); Bloom et al. (2010) wetland and rice paddy CH_4 emissions (green). Riley et al. (2011) attribute their elevated tropical fluxes to anomalously high predicted net primary productivity in the Community Land Model (CLM version 4).





Fig. 5. Hemispheric mean observed and model methane anomalies from surface concentration measurements, 2003–2008. Surface concentration measurements (black) are from the GasLab, AGAGE and ESRL networks (Francey et al., 1996; Prinn et al., 2000; Cunnold et al., 2002; Dlugokencky et al., 2009). The GEOS-Chem global 3-D chemistry transport model (Fraser et al., 2011) is driven by wetland CH_4 emission estimates from Fung et al. (1991) (blue), Bloom et al. (2010) (red), and our new top-down approach (green).

