Biogeosciences Discuss., 10, 5749–5780, 2013 www.biogeosciences-discuss.net/10/5749/2013/ doi:10.5194/bgd-10-5749-2013 © Author(s) 2013. CC Attribution 3.0 License.



This discussion paper is/has been under review for the journal Biogeosciences (BG). Please refer to the corresponding final paper in BG if available.

Modeling microbial exchanges between forms of soil nitrogen in contrasting ecosystems

M. Pansu¹, D. Machado², P. Bottner³, and L. Sarmiento⁴

 ¹IRD, UMR Eco&Sol (Supagro, Cirad, Inra, IRD), Place Viala, Montpellier, France
 ²Laboratorio de Investigación en Análisis Químico Industrial y Agropecuario, Departamento de Química, Facultad de Ciencias, Universidad de los Andes, Mérida, Venezuela
 ³CEFE-CNRS Deceased 15 October 2006, Montpellier, France
 ⁴Instituto de Ciencias Ambientales y Ecológicas, Facultad de Ciencias, Universidad de los Andes, Merida, Venezuela

Received: 16 January 2013 - Accepted: 8 March 2013 - Published: 25 March 2013

Correspondence to: M. Pansu (marc.pansu@ird.fr)

Published by Copernicus Publications on behalf of the European Geosciences Union.



Abstract

Although nitrogen (N) is often linked to carbon (C) in organic molecules, C is carried from the atmosphere to the soil through plants while N is carried from the soil to plants by microbial transformations. Many schemes have been proposed to describe the mi ⁵ crobial conversion between organic and inorganic forms of N but current models do not fully represent the microbial control over these conversions. This study followed the transfer of ¹⁵N between plant materials, microorganisms, humified compartments and inorganic forms in 6 very different ecosystems along an altitudinal transect. The microbial conversion of the ¹⁵N forms appeared to be strongly linked to that found previously for ¹⁴C forms since the parameters and relationships defined for C were appropriate for modeling the N cycle. The only difference was in the flows between microbial and inorganic forms. The CO₂-C loss was modeled using the equation for microbial respiration. Inorganic N appears also closely associated with microorganisms, which, depending

- on their C : N ratio and those of the available substrates, regulate the N mineralization and immobilization processes. Applications at earth scale can use the approximation that the microbial C : N ratio does not vary with time, but for this study, microorganisms
- cannot be treated always as homeostatic as their C: N ratio can decrease during incubation and increase with altitude when C storage increases. The MOMOS model has been validated for the C cycle, and it also appears to be valid for microbial conversion
- ²⁰ of N forms. It uses a relatively small number of well-defined, climate-dependent parameters, and it should fill a gap in the range of current models based on a direct microbial control for describing C and N flows in ecosystems.

1 Introduction

Nitrogen (N) in living plants represents about 5 % of the global N stock: it is adsorbed by plant roots mostly in mineral forms in small quantities in soil, where more than 90 % of N is in organic form (Lin et al., 2000; Pansu and Gautheyrou, 2006). Microbial exchanges



play a major role in the N cycle that must be considered in conjunction with the carbon (C) cycle. Mechanistic models are required to give an accurate prediction of all the transfers of N between organic and inorganic compartments of various stabilities. Manzoni and Porporato (2009) classified the published N models as SIMP for sim-

- ⁵ plified formulations, MIT for mineralization/immobilization turnover mechanisms which assume a transfer of organic to inorganic N pools before microbial assimilation, DIR for direct microbial assimilation of all available organic N, MIX for models combining DIR and MIT principles and PAR for a parallel DIR/MIT scheme including direct assimilation, ammonium production by microorganisms and then microbial assimilation of the ammonium produced (Barraclough, 1997). With increasing knowledge of the mecha-
- nisms, the types of models available have changed from 60 % SIMP and 40 % MIT in 1970 to 5 % SIMP, 7 % MIT, 5 % MIX, 17 % PAR, and 66 % DIR in 2010.

Organic N transformations have often been modeled by considering compartments with different C:N ratios (e.g. van Veen and Ladd, 1985; Bradbury et al., 1993; Carter

- et al., 1993; Dou and Fox, 1995; Quemada and Cabrera, 1995; Richter and Benbi, 1996; Franko, 1996; Mueller et al., 1998; Garnier et al., 2001; Nicolardot et al., 2001; Pansu et al., 2003, 2004; Neill and Gignoux, 2006), but Todd-Brown et al. (2012) considered that "current global models do not represent direct microbial control over decomposition" and a new generation of models is required. An important aspect is re-
- ²⁰ lated to the stoichiometry of decomposers (Sterner and Elser, 2002). Microbial biomass (MB) has often been considered homeostatic, i.e. with a composition independent of that of the substrates used, implying that assumptions are made to maintain a constant MB C:N ratio, but other models and experimental data (Bottner et al., 2006) allow the C:N ratio of MB to change with time in response to the substrate C:N ratio and changes in the microbial communities during decomposition.

This work deals with N dynamics along an altitudinal transect previously used to validate the MOMOS-C model (Pansu et al., 2010). The aim was to predict the conversion of the ¹⁵N labeled forms simultaneously with the conversion of ¹⁴C labeled forms, assuming that MB can assimilate some N from labile and stable molecules of plant



and microbial origin as well as some N from the soil inorganic N pool (see above PAR scheme). This raised three questions:

1 Can it be considered that microbial enzymatic assimilation rates are the same for C and N?

⁵ 2 Can it be considered that C transfers by microbial respiration and mortality cause simultaneous transfers of N into labile humus and inorganic forms to balance the MB C : N ratio? Can the assimilation of inorganic N be modeled to sustain microbial activity in the case of an N deficit during conversion of organic forms?

3 Can it be considered that the microbial biomass is homeostatic or does it have 10 a C:N ratio that varies through incubation periods and is different in ecosystems at different altitudes?

2 Materials and methods

2.1 The experimental sites

The experiment was carried out in six sites (Table 1) along an altitudinal transect in Venezuela, from 65 to 3968 m a.s.l., covering a large bioclimatic gradient that com-15 prised tropical rainforest (A(65)), natural savanna (A(165)), seasonal montane forest (A(780)), cloud forest (A(1800)) and Andean páramo (alpine vegetation) at two heights (A(3400) and A(3968)). The sites have been described in previous publications (Couteaux et al., 2002; Pansu et al., 2010). This altitudinal transect is characterized by contrasting conditions of temperature, annual precipitation and its seasonal distri-20 bution, and soil characteristics. The long-term mean annual air temperature ranged from 5.5 °C at A(3968) to 27.4 °C at A(65), the mean annual precipitation ranged from 790 mm at A(3968) to 1992 mm at A(1800). Soils were acid in all sites but particularly in the two páramo soils. The soils were loam at A(3400) and sandy loams at the other sites. The savanna soil at site A(165) contained the highest amount of sand and 25 the lowest amount of organic matter, both water holding capacity (WHC) and cation



exchange capacity (CEC) were lower than any of the other sites. The mountain soils A(1800) and A(3400) contained the highest amounts of fine particles and organic matter and had the highest WHC and CEC. The three other soils had intermediate WHC and CEC. The carbon content in the 0–10 cm soil layer varied from $13 g(C) kg^{-1}$ at A(165) to more than $100 g(C) kg^{-1}$ (soil) at A(1800) and A(3400). The quality of the soil organic matter was also variable, with C : N ratios ranging from 13 at A(65) and A(780)

through 15 at A(165) up to 17 to 22 at the highest sites A(1800) to A(3968).

2.2 Experimental design and data collection

5

Plots with herbaceous vegetation were selected at each site to minimize the variability due to the effect of soil cover on microclimate conditions, mainly on soil temperature. For the savanna and alpine ecosystems, areas with natural vegetation were selected, but for the forest ecosystems, plots on managed grassland were selected. ¹⁴C and ¹⁵N labeled straw was mixed with soil, from the top 0–10 cm layer at each of the sites, in 14 × 15 cm porous bags. The top part of the bags had a 1 mm mesh to allow the pas-

- sage of plant roots and mesofauna and the mesh of the bottom part was 0.1 mm to minimize losses by gravity. The soil weight per bag was adjusted to reproduce the natural bulk density for a volume of 210 cm^3 . The labeled straw was obtained by growing the wheat in a labeling chamber with controlled temperature, radiation, humidity and CO₂ concentration. The wheat was grown from seed to maturity in four months in a ¹⁴C
- labeled atmosphere with a ¹⁵N labeled, NPK + micro-nutrient solution. The straw, containing 392 mgCg⁻¹ and 12.33 mgNg⁻¹ (C:N ratio of 31.79), was roughly ground to < 5 mm particles. The N and ¹⁵N composition of the soil in the bags is given in Table 2. The added carbon and nitrogen from the straw ranged from 1.2% (A(3400)) to 13.8% (A(165)) of the native C and 0.70% (A(3400)) to 6.60% (A(165)) of the native N in the soil.

40 bags containing the labeled straw and soil were buried 5 cm deep along four parallel lines in each experimental plot (10 samples at different times × 4 replicates for each sample at each site, making a total of 240 soil bags). When the plots were set



up (23 November to 6 December 1994), the soil bags were moistened with de-ionized water to the midpoint between WHC and the water content at wilting point (WCWP) (Table 1). On each sampling date, one bag from each line of the four lines at each site was selected at random to measure soil water content, total ¹⁴C and ¹⁵N and ¹⁴C

- and ¹⁵N in the microbial biomass and inorganic N stock. The soil bags were left in the soil for 18 months at the two lowest sites (A(65) and A(165)) 24 months at A(780), 31 months at A(1800) and 38 months at the two highest sites (A(3400) and A(3968)). The first samples were taken one month after setting up the experiment and the sampling interval increased with time to 6 months at the end of the experiment for the highest sites. After collection, the soil bags were stored refrigerated for no more than three
- days before analysis.

Total N was determined by Kjeldahl digestion by boiling in concentrated sulfuric acid with a potassium sulfate-copper sulfate-grey selenium catalyst for 2 h at 400 °C. The ammonia in the solution was then distilled with sodium hydroxide into a standard H_2SO_4

¹⁵ solution and the excess H₂SO₄ was determined by back titration with an NaOH standard solution. After titration, the distillate was acidified to pH between 3 and 4 to avoid N losses and evaporated to obtain ammonium sulfate crystals that were analyzed for ¹⁵N abundance using mass spectrometry.

The MB N was determined by fumigation-extraction (Brookes, 1985). After homoge-²⁰ nization, a fresh soil sample equivalent to 30 g dry soil was fumigated with alcohol free chloroform for 18 h. The fumigated sample and an equivalent control soil sample were treated with 150 mL of 0.5 mol (K_2SO_4) L⁻¹ solution for 30 min and centrifuged. The extracts were digested, titrated, crystallized and analyzed for ¹⁵N as for total N. The N labeled part of the microbial biomass was calculated as the difference between the

²⁵ labeled N in the fumigated and control samples, corrected by a K_N factor of 0.54 (Joergensen and Mueller, 1996). An aliquot of the extracted solution from the unfumigated samples was used to determine the total inorganic N and ¹⁵N abundance (ammonium and nitrate separately).



For all compartments (total soil, microbial biomass and mineral nitrogen) the percentage of the N in the samples that had come from the N added in the straw (%Ndff) was calculated as:

$$\% \text{Ndff} = \frac{\% E_{\text{comp}}}{\% E_{\text{straw}}} \cdot 100$$

5

10

15

where $\% E_{comp}$ is the atom percent excess of the tracer in the compartment and $\% E_{straw}$ is the atom percent excess in the straw. From this, the amount of N from the straw in each compartment was calculated as:

 $mg N = \frac{\% Ndff \cdot N_{total}}{100}$

where N_{total} is the total N in the sample in mg.

Following the death of our colleague, Pierre Bottner, we recovered most of the experimental data, except the last results for BM and inorganic ¹⁵N, where only the first five results were available for each site.

All ¹⁵N labeled data and ¹⁴C labeled data (Pansu et al., 2010) are expressed as a fraction of the labeled N and C added at the start of the experiment. As the labeled inorganic N in the soil bags was very low compared to the total labeled ¹⁵N, the total labeled ¹⁵N was considered to be the labeled organic ¹⁵N, the difference between the ¹⁵N added in the straw and the organic ¹⁵N being the production of inorganic ¹⁵N which was assumed to have been lost mostly by root uptake, leaching and gaseous losses through the porous soil bags.

The soil water content was measured in each soil bag using four 5 g replicates that were dried at 105 °C for 24 h. Other soil analyses were performed using standard methods (Pansu and Gautheyrou, 2006).

25 2.3 The decomposition model MOMOS

As carbon and nitrogen are closely associated in living organisms, it was assumed that the nitrogen cycle could be modeled in MOMOS-N in the same way as the carbon



cycle in MOMOS-C (Pansu et al., 2004, 2010), using the subscript e (either C or N) to differentiate each element in the model.

MOMOS (Fig. 1) was defined as a five compartment model centered on the activity of soil microbial biomass (MBe) that grows by assimilation of labile (VLe) and stable

- ⁵ (VSe) fractions of plant necromass (NC) as well as labile (HLe) and stable (HSe) fractions of humus. The microbial mortality regulates humus formation. The only process which is considered more of a chemical process than a biological process is humus stabilization from HLe to HSe. The only difference between the C and N models is in the outputs from MBe to inorganic forms of C (CO₂-C) and N (NH₄-N) or possibly in the outputs from MBe to inorganic forms of C (CO₂-C) and N (NH₄-N) or possibly
- ¹⁰ inputs from inorganic N into MB_N. MOMOS has only seven first order kinetic parameters (dimension day⁻¹) and does not need the partitioning coefficients used in other decomposition models. Using the assumption that enzymatic assimilation rates from organic matter, are the same for C and N (see question 1 in the introduction), the best fit parameter values previously found for the C cycle are all used to describe the N cy-
- ¹⁵ cle for each of the 6 ecosystems (Table 3). All the C and N parameters are conditioned by functions of the soil temperature and water content ranging from 0 to 1, as in the general MOMOS equation:

$$\dot{x}_{e} = f(T)f(\theta)\mathbf{A}_{e}x_{e} + B_{e}$$

²⁰ where x_e is the vector of the state variables (¹⁴C or ¹⁵N content of the compartments), \dot{x}_e is the vector of the derivatives of x_e , A_e is the model parameter matrix for each organic element, B_e is a vector determining the external C and N inputs (see Pansu et al., 2009 for C inputs from living roots, $B_e = 0$ for ¹⁴C and ¹⁵N labeled data in this experiment) and f(T) is an exponential function of temperature (Pansu et al., 2010):

25
$$f(T) = Q_{10}^{(T-T_{opt})/10}$$

where T is the actual daily temperature of soil (0–10 cm layer) set equal to the air temperature; T_{opt} is the optimum decomposition temperature set to 28 °C, a temperature often used to perform laboratory experiments under optimum conditions (Thuriès



(1)

(2)

et al., 2002) and just above the mean annual temperature of the warmer sites A(65) and A(165) of this study; Q_{10} is the factor by which the rate increases with a 10 °C increase in temperature. This was set to 2.2 for all sites for this study (see discussion in Pansu et al., 2010); $f(\theta)$ is the response function to soil moisture expressed as a fraction of the WHC (Table 1, see discussion in Pansu et al., 2010):

$$f(\theta) = MIN\left(\frac{\theta}{WHC}, 1\right)$$

The soil water content θ was predicted using the SAHEL model (Penning de Vries et al., 1989). This model calculates the daily water content for each soil layer using meteorological data (daily minimum and maximum temperature, precipitation and latitude), WHC (Table 1) and plant cover as inputs. SAHEL was calibrated for each site using the water content of the soil in the soil bags and then daily water content values for the 0-10 cm layer were generated (Pansu et al., 2004). Meteorological data for the period over which the experiment was carried out was collected for each site from the nearest weather station or estimated using local or archive data, a transition probability matrix and climate corrections as described in Pansu et al. (2010)

The model matrices A_{C} and A_{N} are:

$$\mathbf{A_{C}} = \begin{bmatrix} -k_{VL} & 0 & 0 & 0 & 0 \\ 0 & -k_{VS} & 0 & 0 & 0 \\ k_{VL} & k_{VS} & -\left(q_{CO_{2}} + k_{MB}\right) & k_{HL} & k_{HS} \\ 0 & 0 & k_{MB} & -\left(k_{HL} + k_{HLS}\right) & 0 \\ 0 & 0 & 0 & k_{HLS} & -k_{HS} \end{bmatrix} \text{ and }$$
$$\mathbf{A_{N}} = \begin{bmatrix} -k_{VL} & 0 & 0 & 0 \\ 0 & -k_{VS} & 0 & 0 & 0 \\ k_{VL} & k_{VS} & -\left(f(x_{C,MB}x_{N,MB})/f(T)f(\theta)x_{N,MB} + k_{MB}\right) & k_{HL} & k_{HS} \\ 0 & 0 & k_{MB} & -\left(k_{HL} + k_{HLS}\right) & 0 \\ 0 & 0 & 0 & k_{HLS} & -k_{HS} \end{bmatrix}$$

(3)

Jiscussion Paper

Discussion Paper

Jiscussion Papel

20

5

The vectors x_{c} and x_{N} of the C and N concentrations in each compartment are:

$$\boldsymbol{x}_{C} = \begin{bmatrix} \boldsymbol{x}_{C,VL} \\ \boldsymbol{x}_{C,VS} \\ \boldsymbol{x}_{C,MB} \\ \boldsymbol{x}_{C,HL} \\ \boldsymbol{x}_{C,HS} \end{bmatrix} \qquad \boldsymbol{x}_{N} = \begin{bmatrix} \boldsymbol{x}_{N,VL} \\ \boldsymbol{x}_{N,VS} \\ \boldsymbol{x}_{N,MB} \\ \boldsymbol{x}_{N,HL} \\ \boldsymbol{x}_{N,HS} \end{bmatrix}$$

and the C: N ratios of each compartment are: C : $N_i = \frac{x_{C,i}}{x_{N,i}}$. For each incubation period, the derivative of C is¹:

$$\dot{C} = \sum_{i=1}^{5} \dot{x}_{i,C} = -f(T)f(\theta)q_{CO_2}x_{C,MB}$$

5

where $q_{\rm CO_2}$ is the metabolic quotient of the microbial biomass:

$$q_{\rm CO_2} = k_{\rm resp} \frac{x_{\rm MB}}{C_{\rm MB}^0}$$

where k_{resp} is the respiration coefficient, (dimension day⁻¹) scaled by C_{MB}^{0} , the biomass at steady state (estimated on untreated soil without recent addition of substrate. In this case, it was estimated from the values of MB-¹⁴C measured at the end of incubation).

For each incubation period, the derivative of the total organic N is the negative of the ⁵ derivative of total inorganic N and is expressed by:

$$\dot{N} = \sum_{i=1}^{5} \dot{x}_{i,N} = -f(x_{C,MB}, x_{N,MB})$$

¹The Eq. (5) previously given for MOMOS-C (Pansu et al., 2010) had an optimum \dot{C} which must be multiplied by $f(T)f(\theta)$ to give a \dot{C} adjusted for weather conditions. qCO_2 on the right-hand scale of Fig. 3c–8c of Pansu et al. (2010) must be changed to $f(T)f(\theta)qCO2$.

where positive values of the function $f(x_{C,MB}, x_{N,MB})$ correspond to N mineralization of microbial N and negative values correspond to microbial immobilization of inorganic N. As the simulation concerned only the ¹⁴C and ¹⁵N introduced in the straw, the initial conditions for C and N compartments of microbial origin were set to zero. If C_0 is the ⁵ amount of added ¹⁴C (= 1 for these data scaled by the ¹⁴C input) and f_S is its stable fraction, the initial conditions for the ¹⁴C simulation were given by:

$$x_{C,VL}(0) = (1 - f_S)C_0, x_{C,VS}(0) = f_SC_0, x_{C,MB}(0) = x_{C,HL}(0) = x_{C,HS}(0) = 0$$
(8)

The stable fraction $f_{\rm S}$ was estimated as that of the stable compartment of the TAO (Transformation of Added Organic materials) model (Thuriès et al., 2002) between $f_{\rm S}$ and biochemical composition of straw, which gave $f_{\rm S} = 0.14$. If $\eta_{\rm NC}$ is the C: N ratio of labeled NC, and $\eta_{\rm VS}$ the C: N ratio of the stable fraction of NC, the initial conditions for the ¹⁵N simulation were given by:

$$\begin{aligned} x_{\rm N,VL}(0) &= \left(\frac{1}{\eta_{\rm NC}} - \frac{f_{\rm s}}{\eta_{\rm VS}}\right) C_0, x_{\rm N,VS}(0) = \frac{f_{\rm s}}{\eta_{\rm VS}} C_0, \\ x_{\rm N,MB}(0) &= x_{\rm N,HL}(0) = x_{\rm N,HS}(0) = 0 \end{aligned}$$

The function $f(x_{C,MB}, x_{N,MB})$ of Eq. (7) was defined in terms of η_{MB}^{lim} , the target value for the C : N ratio of the MB (η_{MB}). Two assumptions were tested:

1. an MB C: N ratio being constant throughout incubation:

15

20
$$f(x_{C,MB}, x_{N,MB}) = x_{N,MB} - \frac{x_{C,MB}}{\eta_{MB}^{lim}}$$
 (10)

For this function, there is only one parameter to be fitted for each site, $\eta_{\text{MB}}^{\text{lim}}$, all the other parameters being those fitted for ¹⁴C simulations (Table 3).



(9)

2. an MB C: N ratio which fell linearly with incubation time from η_{MB}^{max} to η_{MB}^{min} at time t_c then constant at η_{MB}^{min} :

$$f(x_{C,MB}, x_{N,MB}) = x_{N,MB} - \frac{x_{C,MB}}{\eta_{MB}^{max} - (\eta_{MB}^{max} - \eta_{MB}^{min})\frac{t}{t_c}} \text{ for } t \le t_c, \text{ otherwise}$$
$$f(x_{C,MB}, x_{N,MB}) = x_{N,MB} - \frac{x_{C,MB}}{\eta_{MB}^{inf}} \text{ for } t > t_c$$

5

For this function three parameters must be fitted for each site: η_{MB}^{max} , η_{MB}^{min} and t_c .

Two alternative simulation strategies were also tested: (a) simulated values of MB ¹⁵N and inorganic ¹⁵N were limited to positive values and (b) negative simulated values for MB ¹⁵N and inorganic ¹⁵N were allowed. The strategy (b) was used for the assumption 1 of a constant MB C : N ratio (although the model did not calculate many negative values, except at the start of simulation), both strategies were tested for the assumption 2 of a decreasing C : N ratio. Simulated negative values did not, of course, indicate that the ¹⁵N content was really negative but that ¹⁴N could replace ¹⁵N to supply the nitrogen requirement.

¹⁵ For each incubation period, the model assumes that $^{15}_{inorg}$ N, the inorganic ¹⁵N remaining in the porous soil bags is the mineralized ¹⁵N (–N, Eq. 7) less the ¹⁵N lost from the bag by plant uptake, leaching or gaseous losses, using a loss rate k_1 :

$${}^{15}_{\text{inorg}} \dot{N} = -\dot{N} (1 - k_{\text{I}}) \text{ if } {}^{15}_{\text{inorg}} N > 0, \text{ otherwise } {}^{15}_{\text{inorg}} \dot{N} = 0 \text{ if } {}^{15}_{\text{inorg}} N \le 0$$
(12)

²⁰ The Powell optimization method was used to estimate the values of η_{MB}^{lim} , η_{MB}^{max} , η_{MB}^{min} , t_c , and k_l for the six experimental sites. The values of the other parameters (Table 3) remained unchanged from MOMOS-C calibration (Pansu et al., 2004) and validation (Pansu et al., 2010). The model was developed using VENSIM 5.6b (http://www.vensim.com).



(11)

2.4 Accuracy tests

The significance of the MOMOS simulations compared to the mean of measured values was tested by:

$$F = \frac{\sum_{i=1}^{n} (\bar{y}_i - \bar{y})^2 / (n-1)}{\sum_{i=1}^{n} (\bar{y}_i - \hat{y}_i)^2 / (n-p)}$$

10

20

where i = 1, ..., n is the number of sampling occasions (n = 11), p the number of model parameters which were specifically adjusted to predict the total production of inorganic ¹⁵N (p = 1 for assumption 1, p = 3 for assumption 2), \bar{y}_i the measured total remaining ¹⁵N at *i*, \hat{y}_i the corresponding MOMOS predicted value with assumption 1 or 2 and \bar{y} is the mean of the data series for each site. A graphical representation (Figs. 2 to 7) shows whether the predicted values were within or outside the confidence intervals of the corresponding data series.

F tests were performed using RSS_{T} the residual sum of squares between the measured values and the values predicted by MOMOS for assumptions 1 and 2 (Table 4):

15
$$F_{y_{A12}} = \frac{\text{RSS}_{A1}}{\text{RSS}_{A2}} = \frac{\sum_{i=1}^{n} (\bar{y}_i - \hat{y}_{iA1})^2 / (n-1)}{\sum_{i=1}^{n} (\bar{y}_i - \hat{y}_{A2})^2 / (n-3)}$$
 (14)

where \hat{y}_{A1} and \hat{y}_{iA2} were the predicted values for assumptions 1 and 2, respectively. An *F* value (Eq. 11) greater than $F_{(n,n-p)}^{0.05}$ indicates that assumption 1 must be rejected at 5% significance level: RSS_{A1} was significantly greater than RSS_{A2} and so assumption 1 predictions were significantly less accurate than assumption 2 predictions. A non significant *F* test (Eq. 14) meant the two assumptions did not give significantly different predictions.



(13)

3 Results

3.1 Mineralization of added ¹⁵N

The model assumed that mineralized ¹⁵N was the difference between added ¹⁵N and the remaining ¹⁵N. The largest part of this mineralized ¹⁵N was exported from the porous soil bags by root absorption, water leaching or gaseous losses, since mineral ¹⁵N remaining in bags represented only 1–3% of the mineralized ¹⁵N (Figs. 2–7). The data showed a decrease in ¹⁵N mineralization rates from low altitude sites to higher sites. About 63% of the added ¹⁵N was mineralized at the lowest sites A(65) and A(165), 57% at A(780), 47% at A(1800), 25% at A(3400) and 31% at A(3968). The ¹⁴C mineralization at the end of incubation (Pansu et al., 2010) was higher at about 80% of the added ¹⁴C at the lowest sites A(65) and A(1800) and 45% at the highest sites A(3400) and A(3968), which indicated the same trend for both ¹⁴C and ¹⁵N, all values being well predicted by MOMOS.

3.2 Prediction using constant MB C: N ratio

¹⁵ The predicted values of ¹⁵N mineralization (Table 4) using a constant MB C:N ratio during incubation, assumption 1 (Eq. 10) corresponded with the measured values only for low altitude sites at 1 % significance level for A(65) and A(165) and 5 % significance level for A(780). Predicted values were within the 95 % confidence interval of the measured data for A(165) (Fig. 3), slightly overestimated between 360 and 500 d
²⁰ incubation for A(65) (Fig. 2) and after 5 months incubation for A(780). For A(1800), the values were slightly underestimated for the first three months and 20 % overestimated after 8 months of incubation. This effect was larger for the two higher sites where the predicted values agreed with the measurements only for the first year of incubation, after 3 yr of incubation the overestimate was 35 % for A(3968) and 50 % for A(3400).



3.3 Prediction of ¹⁵N mineralization using variable MB C: N ratio

Assumption 2 (Eq. 11) gave more accurate predictions than assumption 1, especially at high altitude. The predicted values of ¹⁵N mineralization (Table 4) corresponded with the measured values at 1 % significance level for 5 of the 6 sites. For A(3400), the predicted values were not significantly close to the mean of the measurements.

In five cases, the predicted values were significantly closer when assumption 2 was used, at 1 % significance level for A(65), A(780), and A(1800) and at 5 % significance level for A(3400) and A(3968). There was no significant difference for A(165). Using the strategy (b), which allowed negative values for inorganic ¹⁵N and MB ¹⁵N, gave closer predicted values for the two highest sites, at 1 % significance level for A(3968) and 5 % significance level for A(3400).

For A(65), A(780) and A(1800) (Figs. 2, 4, 5), the predicted values, using variable MB C:N ratio, truncating negative values (strategy (a)), were within the 95% confidence intervals of all the measurements. For A(3400) (Fig. 6), 10 of the 11 predicted values

¹⁵ were within the 95 % confidence intervals for both strategy (a) (curve 2) and strategy (b) (curve 3). For A(3968) (Fig. 7), 9 of the 11 predicted values were within the 95 % confidence intervals of the measurements when strategy (a) was used (curve 2). All predicted values were within the 95 % confidence intervals when strategy (b) was used (curve 3).

20 3.4 Prediction of other N labeled compartments

Unfortunately measurements were only available for the first five periods for MB ¹⁵N and inorganic-¹⁵N in the soil bags (see material and methods) and the model fit was not significant.

For A(65) (Fig. 2), 4 of the 5 MB ¹⁵N values were slightly over-estimated using assumption 1 (curve 1) and 3 were overestimated using assumption 2 (curve 2). However, over the same period, the MB ¹⁴C values were slightly underestimated. For A(165) (Fig. 3), MB ¹⁵N values were overestimated while the MB ¹⁴C predicted values were



always within the 95 % confidence intervals. For A(780) (Fig. 4), the last MB ¹⁵N value was slightly underestimated (but within its wide confidence interval) while the 3 previous values were underestimated. However, over the same period, the MB ¹⁴C values were underestimated. For A(1800) (Fig. 5), 3 values of MB ¹⁵N were accurately predicted while one was underestimated and one was overestimated. For A(3400) (Fig. 6), both the MB ¹⁵N and the MB ¹⁴C values were overestimated. For A(3968) (Fig. 7), 2 of the 5 MB ¹⁵N values were overestimated as were the last 6 MB-¹⁴C values.

All predicted values for organic ¹⁵N in the soil bags were close to the 95 % confidence intervals and sometimes closer with assumption 1 or assumption 2 but without any significant differences between the two assumptions.

In all cases, the model predicted a relatively large storage of ¹⁵N in the labile microbial metabolites (HL compartment), which was about 80 % of the organic forms of ¹⁵N at the end of experiment. NC ¹⁵N was predicted as being almost exhausted and the rest of the ¹⁵N was divided between microbial biomass (MB) and stable humus (HS) with a ¹⁵N-MB : ¹⁵N-HS ratio which increased with altitude (Figs. 2–7).

4 Discussion

10

4.1 Relationship between N and C cycles

These modeling results provide a positive answer to the first two questions raised in the introduction:

²⁰ 1 Can it be considered that microbial enzymatic assimilation rates are the same for C and N? Yes, we can argue that because all the parameters that had been defined for the MOMOS-C model, and their dependence on climate, quality of dead plant materials and soil texture, were retained for modeling the N cycle. This study shows that there is a strong link between the transfer processes of C and N from natural organic ²⁵ compounds.



2 Can it be considered that C transfers by microbial respiration and mortality cause simultaneous transfers of N into labile humus and inorganic forms to balance the MB C : N ratio? Yes, this modelling study demonstrates the simultaneous MB-N transfer into labile humus by microbial mortality and into inorganic N by ammonification. Moreover,
 ⁵ can the assimilation of inorganic N be modelled to sustain microbial activity in the case of an N deficit during conversion of organic forms? Yes, an assimilation of inorganic N is modelled to sustain microbial activity in case of N deficit during the transformation processes.

This study also provided an answer to question 3: can it be considered that the microbial biomass is homeostatic or does it have a C: N ratio that varies through incubation periods and is different in ecosystems at different altitudes? The model predictions using the assumption of a constant MB C: N ratio over the incubation period gave MB C: N values (Table 4) ranging from 13.6 at the tropical savannah site A(165) to 22.8 at the highest site A(3968). These values were close to the measured total C: N ratios
 ¹⁵ (Table 1) for the six sites, suggesting that the quality of living and dead organic materials converge to similar values after long fallow periods. They could appear high since the C: N ratio is generally allowed to vary within a restricted range between about 5 and 15 (Manzoni and Porporato, 2009). Other results could suggest higher values, e.g. Bottner et al. (2006) measured MB ¹⁴C : ¹⁵N ratios from 7.9 ± 1.3 for a substrate ¹⁴C : ¹⁵N

²⁰ ratio of 26.8 to 33.9 ± 7.5 for a substrate ¹⁴C : ¹⁵N ratio of 130 (2 × 10 measurements in two sites). As a first approximation for future applications of MOMOS, constant values could be used over the incubation period, especially for the low altitude sites.

Especially for the high altitude sites in this study, the predicted values were significantly more accurate when the MB C : N ratio was reduced linearly with the incubation

time. With excess C at the start of incubation, the MB C : N ratio was the highest, which encouraged microbial immobilization of inorganic N. The C : N value was reduced linearly with time to its minimum value associated with a reduction in C mineralization, with a lower slope at higher altitudes. The minimum values found for MB C : N ratios (Table 4) were in the commonly accepted range, except at the high altitude site A(3968)



where CO_2 -C respiration was reduced by lower temperatures. Bottner et al. (2006) showed that total respired CO_2 -¹⁴C was lower for nitrogen-poor straw than for nitrogen-rich straw, which was partly explained by an increase in microbial mortality (k_{MB} rate) which could increase the HL reserve which is richer in N than stable NC and can sustain MB and its conversions to inorganic N.

Except for A(165)and A(3968), the slopes of the MB C:N ratio vs. time decreased with increasing altitude (t_c in Table 4). This appears to be consistent with a decrease in metabolic rates with temperature (Eqs. 1 and 2).

Allowing negative values for inorganic ¹⁵N (immobilization of inorganic ¹⁴N) only significantly improved the predicted values for ¹⁵N mineralization at the two high altitude sites, especially at the highest. For these sites, the model predicted microbial immobilization of N not only at the start of incubation but also later on during incubation. This strange behavior needs to be investigated by other experiments.

4.2 Ecological consistency and parsimony

5

10

- ¹⁵ This study established that there is a strong link between the C and N assimilation, the only difference between the model for C and the model for N being the modeling of the microbial conversions to and from inorganic compounds. Carbon is removed from the system as the CO₂ from microbial respiration (Eqs. 5 and 6), while inorganic N recycles rapidly in the soil in equilibrium with microbial N. This confirms that MB acts as a very active, short-term reserve, temporarily storing C and N, releasing C by respiration,
- producing C,N labile humus compounds (HL in Fig. 1), by mortality and exudation, recycling the major part of this HL and converting to and from mineral forms of N. The MOMOS model encompasses the principle of parallel C and N assimilation (see PAR models in the introduction) as it includes simultaneous, direct microbial assimilation of
- ²⁵ plant and humus compounds, ammonium production by microorganisms and, possibly, microbial assimilation of the ammonium produced. In terms of number and definition of its parameters, MOMOS is parsimonious (Ockham's razor), this model should fill



a gap shown above in introduction for the modeling of "direct microbial control over decomposition".

References

5

15

20

Barraclough, D.: The direct or MIT route for nitrogen immobilization: a ¹⁵N mirror image study with leucine and glycine, Soil Biol. Biochem., 29, 101–108, 1997.

- Bottner, P., Pansu, M., Sarmiento, L., Hervé, D., Callisaya-Bautista, R., and Metselaar, K.: Factors controlling decomposition of soil organic matter in fallow systems of the high tropical Andes: a field simulation approach using 14C and ¹⁵N labelled plant material, Soil Biol. Biochem., 38, 2162–2177, doi:10.1016/j.soilbio.2006.01.029, 2006.
- Bradbury, N. J., Witmore, A. P., Hart, P. B. S., and Jenkinson, D. S.: Modelling the fate of nitrogen in crop and soil in the years following application of ¹⁵N-labelled fertilizer to winter wheat, J. Agr. Sci., 121, 363–379, 1993.
 - Brookes, P. C., Landman, A., Pruden, G., and Jenkinson, D. S.: Chloroform fumigation and the release of soil nitrogen : a rapid direct extraction method to measure microbial biomass nitrogen in soil, Soil Biol. Biochem., 17, 837–842, 1985.
- Carter, M. R., Parton, W. J., Rowland, I. C., Schultz, J. E., and Steed, G. R.: Simulation of soil organic carbon and nitrogen changes in cereal and pasture systems of southern Australia, Aust. J. Soil Res., 31, 481–491, 1993.

Dou, Z. and Fox, R. H.: Using NCSWAP to simulate seasonal nitrogen dynamics in soil and corn, Plant Soil, 177, 235–247, 1995.

Franko, U.: Simulation of carbon and nitrogen dynamics in rural areas, Landbauforsch. Volk., 46, 114–120, 1996.

Garnier, P., Néel, C., Mary, B., and Lafolie, F.: Evaluation of a nitrogen transport and transformation model in a bare soil, Eur. J. Soil Sci., 52, 253–268, 2001.

²⁵ Joergensen, R. G. and Mueller, T.: The fumigation-extraction method to estimate soil microbial biomass: Calibration of the k(EN) value, Soil Biol. Biochem., 28, 33–37, 1996.

Lin, B.-L., Sakoda, A., Shibasaki, R., Goto, N., and Suzuki, M.: Modelling a global biogeochemical nitrogen cycle in terrestrial ecosystems, Ecol. Model., 135, 89–110, 2000.

Manzoni, S. and Porporato, A.: Soil carbon and nitrogen mineralization: theory and models across scales, Soil Biol. Biochem., 41, 1355–1379, 2009.



30

- Mueller, T., Magid, J., Jensen, L. S., Svendsen, H., and Nielsen, N. E.: Soil C and N turnover after incorporation of chopped maize, barley straw and blue grass in the field: evaluation of the DAISY soil-organic-matter submodel, Ecol. Model., 111, 1–15, 1998.
- Neill, C. and Gignoux, J.: Soil organic matter decomposition driven by microbial growth: a simple model for a complex network of interactions, Soil Biol. Biochem., 38, 803–811, 2006.
- ⁵ model for a complex network of interactions, Soil Biol. Biochem., 38, 803–811, 2006. Nicolardot, B., Recous, S., and Mary, B.: Simulation of C and N mineralisation during crop residue decomposition: a simple dynamic model based on the C:N ration of the residues, Plant Soil, 83, 83–103, 2001.

Pansu, M. and Gautheyrou, J.: Handbook of Soil Analysis – Mineralogical, Organic and Inorganic Methods, Springer, Berlin, Heidelberg, New York, 993 pp., 2006.

10

30

Pansu, M., Thuriès, L., Larré-Larrouy, M. C., and Bottner, P.: Predicting N transformations from organic inputs in soil in relation to incubation time and biochemical composition, Soil Biol. Biochem., 35, 353–363, 2003.

Pansu, M., Bottner, P., Sarmiento, L., and Metselaar, K.: Comparison of five soil organic mat-

ter decomposition models using data from a ¹⁴C and ¹⁵N labeling field experiment, Global Biogeochem. Cy., 18, GB4022, doi:10.1029/2004GB002230, 2004.

Pansu, M., Martineau, Y., and Saugier, B.: A modelling method to quantify in situ the input of carbon from roots and the resulting C turnover in soil, Plant Soil, 317, 103–120, doi:10.1007/s11104-008-9791-1, 2009.

Pansu, M., Sarmiento, L., Rujano, M. A., Ablan, M., Acevedo, D., and Bottner, P.: Modeling organic transformations by micro-organisms of soils in six contrasting ecosystems: validation of the MOMOS model, Global Biogeochem. Cy., 24, GB1008, doi:10.1029/2009GB003527, 2010.

Penning de Vries, F. W. T., Jansen, D. M., ten Berge, H. F. M., and Bakema, A.: Simulation of

- Ecophysiological Processes of Growth in Several Annual Crops, Pudoc, Wageningen, 271 pp., 1989.
 - Quemada, M. and Cabrera, M. L.: CERES-N model predictions of nitrogen mineralized from cover crop residues, Soil Sci. Soc. Am. J., 59, 1059–1065, 1995.

Richter, J. and Benbi, D. K.: Modeling of nitrogen transformations and translocations, Plant Soil, 181, 109–121, 1996.

Sterner, R. W. and Elser, J. J.: Ecological stoichimometry, in: The Biology of Elements from Molecules to Biosphere, edited by: Press, P. U., Princeton and Oxford, 2002.



- Thuriès, L., Pansu, M., Larré-Larrouy, M. C., and Feller, C.: Biochemical composition and mineralization kinetics of organic inputs in a sandy soil, Soil Biol. Biochem., 34, 239–250, 2002.
 Todd-Brown, K. E. O., Hopkins, F. M., Kivlin, S. N., Jennifer M. Talbot, J. M., and Allison, S. D.: A framework for representing microbial decomposition in coupled climate models, Biogeochemistry, 109, 19–33, doi:10.1007/s10533-011-9635-6, 2012.
- Van Veen, J. A., Ladd, J. N., and Amato, M.: Turnover of carbon and nitrogen through the microbial biomass in a sandy loam and a clay soil incubated with [¹⁴C(U]glucose and [¹⁵N](NH₄)₂SO₄ under different moisture regimes, Soil Biol. Biochem., 17, 747–756, 1985.

5



Table 1. Site characteristics.

Site	El Vigia	Barinas	Tovar	Merida	Gavidia	El Banco			
Number	A(65)	A(165)	A(780)	A(1800)	A(3400)	A(3968)			
Site characteristics									
Altitude m Latitude N Longit. W Typical ecosystem Actual vegetation Temperature ^a Precipitation ^b	65 8°37'33″ 71°40'6″ Tropical rainforest Managed grassland 27.4 1825 1711	165 8°36′55″ 70°12′15″ Natural savanna Natural savanna 26.4 1565 1297	780 8°20'32″ 71°43'39″ Seasonal forest Managed grassland 23.0 1112 1054	1800 8°37'39'' 71°9'17'' Cloud forest Managed grassland 17.4 1992 785	3450 8°40'04" 70°54'58" Andean páramo 20 yr fallow 8.9 1338 557	3940 8°48'52'' 70°55'30'' High páramo Natural páramo 5.5 790 515			
Soil characteristics									
WRB type ^c	Inceptisol	Alfisol	Mollisol	Inceptisol	Inceptisol	Entisol			
C g kg ⁻¹	36.77	13.67	48.23	102.27	100.57	61.53			
N g kg ⁻¹	2.80	0.90	3.70	6.10	5.27	2.77			
D:N	13.1	15.2	13.0	16.8	19.1	22.2			
nH	5.1	5.7	6 1	5.2	4.6	4 7			
CEC ^d	13.9	5.2	13.1	26.5	24.8	12.1			
Sand (%DW)	67.3	77.0	62.0	69.3	40.0	62.0			
Silt (%DW)	24.0	14.0	31.3	25.3	42.0	30.0			
Clay (%DW)	8.7	9.0	6.7	5.3	18.0	8.0			
WHC ^e	37.82	14.69	21.38	37.71	35.67	21.42			
WCWP ^e	16.29	2.33	4.78	29.68	18.09	7.10			
WCI	27	8.5	13	33	26	14			

^a Long-term annual mean temperature in °C.

^b Long-term annual mean precipitation and evapo-transpiration (mm).

^c World Reference Basis.

^d Cation Exchange Capacity mmol (+) kg⁻¹. ^e Water Holding Capacity, Water Content at Wilting Point, and Initial Water Content in soil bags (% DW).



Discussion Paper	B 10, 5749– Modelin	GD 5780, 2013 g forms of						
—	SOILN	itrogen						
Discu	M. Par	M. Pansu et al.						
issi								
on F	Title	Title Page						
aper	Abstract	Introduction						
	Conclusions	References						
Disc	Tables	Figures						
ussio	14	►I						
n Pap	•							
Der	Back	Close						
—	Full Scr	een / Esc						
Discussion F	Printer-frie Interactive	Printer-friendly Version Interactive Discussion						
aper	C	O BY						

Table 2. Composition of soil in bags buried in 5–10 cm soil layer.

Site	g straw/ bag ^a	mg straw C/ bag ^b	mg straw N/ bag ^c	g soil + straw/bag ^a	mg N g ⁻¹ soil + straw ^a	% ¹⁵ N ^a	% ¹⁵ N excess ^d
A(65)	1.08	423.36	13.32	189.32	2.88	0.6094	0.2434
A(165)	1.08	423.36	13.32	225.43	0.93	0.8815	0.5155
A(780)	0.84	329.28	10.36	151.31	3.90	0.5347	0.1687
A(1800)	0.84	329.28	10.36	113.11	6.98	0.4964	0.1304
A(3400)	0.48	188.16	5.92	160.97	5.40	0.4310	0.0650
A(3968)	0.48	188.16	5.92	182.69	2.88	0.4857	0.1197

Straw C = 392 mgg^{-1} ; straw N = 12.33 mgg^{-1} ; straw C : N ratio = 31.79.

 15 N natural abundance = 0.366 %.

^a Measured values.

 b g straw × straw C. c g straw × straw N. d % 15 N – 15 N natural abundance.

Table 3. Parameter values used in the MOMOS-C model (Pansu et al., 2010) and retained in this N study.

Param-	Definition	Units	Reference	Values					
eter				A(65)	A(165)	A(780)	A(1800)	A(3400)	A(3968)
f _s	Stable C fraction of added NC	None	TAO model, Thuriès et al. (2002), Pansu and Thuriès (2002), Kaboré et al. (2010, 2011)			C	.144		
k _{VL}	Microbial assimilation rate of labile NC	day ⁻¹	MAX(0.65 – 0.0019 $\eta_{ m NC},$ 0.1) Bottner et al. (2006)			C	.590		
k _{vs}	microbial assimilation rate of stable NC	day ⁻¹	MAX(0.0037 – 0.000026 $\eta_{\rm NC}$, 0.00005) Bottner et al. (2006)	;, 0.0028					
k _{HL}	Microbial assimilation rate of labile humus	day ⁻¹	0.05 Pansu et al. (2004)				0.05		
k _{HS}	Microbial assimilation rate of stable humus	day ^{−1}	0.00005 Pansu et al. (2004)			0.0	00005		
k _{HLS}	Rate of stabilization of HL to HS	day ⁻¹	0.0003 Pansu et al. (2004)	0.0003					
k _{MB}	Mortality rate of MB	day ⁻¹	MIN(0.42 + 0.0012η _{NC} , 0.8) Bottner et al. (2006)			C	.458		
k _{resp}	Microbial respiration	day ⁻¹	–0.0008 <i>F</i> ₀₋₂₀ + 0.062 Pansu et al. (2007, 2010)	0.029	0.038	0.034	0.029	0.021	0.022
C_0^{BM}	MB-C at steady state	$g MB - {}^{14}C$ $g^{-1} added - {}^{14}C$	Pansu et al. (2010)	0.0107	0.0106	0.0192	0.0129	0.0185	0.0156

MB is microbial biomass, NC is added necromass, $\eta_{\rm NC}$ is the NC C : N ratio.

BGD 10, 5749–5780, 2013 **Modeling forms of** soil Nitrogen M. Pansu et al. **Title Page** Abstract Introduction Conclusions References Tables Figures 14 Back Close Full Screen / Esc Printer-friendly Version Interactive Discussion $(\mathbf{\hat{u}})$ (cc)

Discussion Paper

Discussion Paper

Discussion Paper

Discussion Paper

Paramotor	Definition	Linite	Description				/aluaa		
Falametei	Demilion	Units	Description	A(65)	A(165)	A(780)	A(1800)	A(3400)	A(3968)
$\eta_{\rm MB}^{\rm lim}$	MB C : N ratio	None	Assumption 1 constant C : N ratio	18.3	13.6	19.3	18.9	19.1	22.8
F test	Eq. (13)		Assumption 1 constant C : N ratio	6.5 ^a	9.2 ^a	3.1 ^b	1.6 NS	0.6 NS	0.7 NS
$\eta_{\rm MB}^{\rm min}$	Minimum value of MB C : N ratio	None	Assumption 2 variable C : N ratio	13.7	12.9	14.2	13.0	12.2	18.7
$\eta_{\rm MB}^{\rm max}$	Maximum value of MB-C : N ratio	None	Assumption 2 variable C : N ratio	23.5	33.7	24.0	23.7	21.4	42.8
t _c	Time for linear decrease from η_{MB}^{min} to η_{MB}^{max}	days	Assumption 2 variable C : N ratio	139	5	226	325	978	66
F test	Eq. (13)		Assumption 2 variable C : N ratio	68.6 ^a	7.8 ^a	32.9 ^a	12.4 ^a	2.3 NS	6.1 ^a
$F_{y_{H12}}$ test	Eq. (14)		Assumption 2, negative values not possible	10.6 ^a	0.8 NS	10.6 ^a	7.7 ^a	3.7 ^b	3.6 ^b
$F_{y_{H12}}$ test	Eq. (14)		Assumption 2, negative values possible					4.2 ^b	9.2 ^a
k _l	Rate of transfer of mineral ¹⁵ N to plants and losses	day ⁻¹	Assumption 1 constant C : N ratio	0.65	0.14	0.24	0.25	0.17	0.48
k _l	Rate of transfer of mineral ¹⁵ N to plants and losses	day ⁻¹	Assumption 2 variable C : N ratio	0.21	0.11	0.19	0.21	0.16	0.07

Table 4. Values and significance of additional parameters for MOMOS-N (this study).

MB is the microbial biomass.

^a 1 % significance level.

^b 5%, significance level.

NS not significant.











Fig. 2. Labeled microbial biomass (left), inorganic (centre) and organic forms (right) of ¹⁵N in A(65), points are measurements with 95% confidence intervals, lines are values predicted by the model using (1) assumption 1 (dashed lines) or (2) assumption 2 and strategy (a) (solid lines), MB is microbial biomass, HL is labile humus ¹⁵N, HS is stable humus ¹⁵N, NC is necromass ¹⁵N.





Fig. 3. Labeled microbial biomass (left), inorganic (centre) and organic forms (right) of ¹⁵N in A(165), points are measurements with 95% confidence intervals, lines are values predicted by the model using (1) assumption 1 (dashed lines) or (2) assumption 2 and strategy (a) (solid lines), MB is microbial biomass, HL is labile humus ¹⁵N, HS is stable humus ¹⁵N, NC is necromass ¹⁵N.





Fig. 4. Labeled microbial biomass (left), inorganic (centre) and organic forms (right) of ¹⁵N in A(780), points are measurements data with 95% confidence intervals, lines are values predicted by the model using (1) assumption 1 (dashed lines) or (2) assumption 2 and strategy (a) (solid lines), MB is microbial biomass, HL is labile humus ¹⁵N, HS is stable humus ¹⁵N, NC is necromass ¹⁵N.





Fig. 5. Labeled microbial biomass (left), inorganic (centre) and organic forms (right) of ¹⁵N in A(1800), points are measurements with 95% confidence intervals, lines are values predicted by the model using (1) assumption 1 (dashed lines) or (2) assumption 2 and strategy (a) (solid lines), MB is microbial biomass, HL is labile humus ¹⁵N, HS is stable humus ¹⁵N, NC is necromass ¹⁵N.





Fig. 6. Labeled microbial biomass (left), mineral (centre) and organic forms (right) of ¹⁵N in A(3400), points are measurements with 95% confidence intervals, lines are values predicted by the model using (1) assumption 1 (dashed lines), (2) assumption 2 and strategy (a) (solid lines), or (3) assumption 2 and strategy (b) (dashed lines) MB is microbial biomass, HL is labile humus ¹⁵N, HS is stable humus ¹⁵N, NC is necromass ¹⁵N.





Fig. 7. Labeled microbial biomass (left), mineral (centre) and organic forms (right) of ¹⁵N in A(3968), points are measurements with 95% confidence intervals, lines are values predicted by the model using (1) assumption 1 (dashed lines), (2) assumption 2 and strategy (a) (solid lines), or (3) assumption 2 and strategy (b) (dashed lines) MB is microbial biomass, HL is labile humus ¹⁵N, HS is stable humus ¹⁵N, NC is necromass ¹⁵N.

