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A Bayesian framework for model calibration, comparison and analysis: application to four models for the biogeochemistry of a Norway spruce forest

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Abstract

Four different parameter-rich process-based models of forest biogeochemistry were analysed in a Bayesian framework consisting of three operations: (1) Model calibration, (2) Model
comparison, (3) Analysis of model-data mismatch.

Data were available for four output variables common to the models: soil water content and emissions of N$_2$O, NO and CO$_2$. All datasets consisted of time series of daily measurements. Monthly averages and quantiles of the annual frequency distributions of daily emission rates were calculated for comparison with equivalent model outputs. This use of the data at model-appropriate temporal scale, together with the choice of heavy-tailed likelihood functions that accounted for data uncertainty through random and systematic errors, helped prevent asymptotic collapse of the parameter distributions in the calibration.

Model behaviour and how it was affected by calibration was analysed by quantifying the normalised RMSE and $r^2$ for the different output variables, and by decomposition of the MSE into contributions from bias, phase shift and variance error. The simplest model, BASFOR, seemed to underestimate the temporal variance of nitrogenous emissions even after calibration. The model of intermediate complexity, DAYCENT, simulated the time series well but with large phase shift. COUP and MoBiLE-DNDC were able to remove most bias through calibration.

The Bayesian framework was shown to be effective in improving the parameterisation of the models, quantifying the uncertainties in parameters and outputs, and evaluating the different models. The analysis showed that there remain patterns in the data - in particular infrequent events of very high nitrogenous emission rate – that are unexplained by any of the selected forest models and that this is unlikely to be due to incorrect model parameterisation.

**Keywords:** Bayesian calibration; carbon cycle; likelihood of data; nitrogen cycle; NO; N$_2$O; prior and posterior probability distributions for parameters; uncertainty analysis; water cycle
1. Introduction

1.1 Rationale

Various recent reviews have assessed the evidence for impacts of environmental change on European forests (Hyvönen et al., 2007; Kahle et al., 2008; Luyssaert et al., 2010). Most studies have focused on changes in growth and carbon balance, but the importance of the interaction with the nitrogen cycle is increasingly recognised (de Vries et al., 2009; Sutton et al., 2008; Van Oijen et al., 2008a; Van Oijen et al., 2004). Research programmes to measure and model emissions of nitrogenous greenhouse gases from European forests and other ecosystems have been set up (Sutton et al., 2007).

The measurement of nitrous oxide (N₂O) and nitric oxide (NO) emissions from forest soils is hampered by the large spatial and temporal heterogeneity in the fluxes, and modelling these processes is still limited by availability of data (Kesik et al., 2005). Moreover, the relevant underlying mechanisms have not yet been clarified fully, and large uncertainties are present in both data and models. Available data sets not only suffer from random measurement error, but also from systematic errors associated with the positioning of measurement chambers in the field and their functioning (Butterbach-Bahl et al., 2002; Kroon et al., 2010). When modelling the systems, there is uncertainty about how to represent processes, i.e. model structural uncertainty (de Bruijn et al., 2009). Furthermore, there is uncertainty about environmental drivers and parameter values.

To improve the applicability of models to the analysis of the greenhouse gas balance of forests, these uncertainties need to be quantified and reduced. Probabilistic methods of model-data fusion or data-assimilation have come to the fore in recent years, and offer the
prospect of improved data use and uncertainty quantification (Fox et al., 2009; Wang et al., 2009). Because these methods are applications of probability theory, they require all uncertainties – in data, model inputs and model structure - to be expressed in the form of probability distributions. Bayes’ Theorem can then be employed to update the distributions when new information becomes available.

In biogeochemical modelling, most Bayesian applications have focused on parameterisation of individual models, with little attention for systematic errors in data and model structure. Wang et al. (2009) thus concluded, in a recent review on model-data fusion studies for terrestrial ecosystems, that there is a need for “developing an integrated Bayesian framework to study both model and measurement errors systematically”. The work presented here is intended to contribute to that goal.

1.2 Towards a Bayesian framework for dynamic modelling in forest biogeochemistry

We propose a framework which requires that multiple models are used in any given study, and which consists of three operations: (1) Bayesian calibration, (2) Bayesian model comparison, (3) Analysis of model-data mismatch.

The overarching objective of this paper is to demonstrate that this three-stage framework is an effective tool for the analysis of models in forest biogeochemistry. For that purpose, we used four different published models and one rich data set from the Norway spruce forest in Höglwald, Germany (Kreutzer et al., 2009). Most of the data were on the nitrogen cycle, with long time series of measurements of emissions of N₂O and NO, but we also used time series of the carbon and water cycles in the form of soil respiration and soil water content.

Bayesian calibration, i.e. the first operation in the framework, consists of defining a prior probability distribution for a model’s parameters and updating that distribution using the
data. The method has not often been applied to parameter-rich nonlinear process-based ecosystem models (Luo et al., 2009). One reason is the high computational demand associated with the technique, which is exacerbated by the long running time of the models. A second issue is the difficulty of quantifying uncertainties about random and systematic measurement errors. We show in this paper how both types of error can be accommodated in a Markov Chain Monte Carlo algorithm for Bayesian calibration.

Bayesian model comparison, the second operation used in the framework, aims to determine the extent to which the data support the different models. This is done by providing a probability distribution over models rather than parameter values. The attempt in this paper to assess whether Bayesian model comparison as a method can be useful for model selection purposes is, as far as we are aware, new for parameter-rich process-based ecosystem models.

Detailed analysis of model-data mismatch, the third operation in our framework, is not a common step in Bayesian model studies, which tend to focus on the probabilistic aspects of model behaviour rather than the internal structure of the models (Gelman and Shalizi, 2010). Bayesian calibration and model comparison effectively treat models as black boxes that convert parameter values into outputs, so this further analysis is needed to facilitate model improvement.

In summary, this paper aims to show the strengths and weaknesses of this three-operation Bayesian framework using a case-study with four models simulating the biogeochemistry of a Central European spruce forest.

2. Materials and Methods

2.1 Data

All data were taken from the Norway spruce (Picea abies L.) site at Höglwald, Germany,
latitude 48°30'N, longitude 11°10'E, altitude 540 m (Papen and Butterbach-Bahl, 1999). Trees were planted in 1907. Soil C and N were around 90,000 and 5000 kg ha⁻¹ (Kreutzer et al., 2009; Rothe, 1997). For the years 1985-1995, mean annual temperature was 7.9 °C, precipitation 888 mm, and atmospheric N-deposition as measured in the throughfall 39.4 kg N ha⁻¹ (Rothe, 1997). For 1975-1990, average global radiation was 11.3 MJ m⁻² d⁻¹ and wind speed 2.8 m s⁻¹ (data from JRC-Ispria, as cited by (Van Oijen et al., 2008b)).

The primary data from the site were in the form of time series of daily measurements of soil water content and soil emissions of N₂O, NO and carbon dioxide (CO₂) (see e.g. Wu et al., 2010). Measurements at the Höglwald Forest are continuous throughout the year with fluxes being available in sub-daily resolution. However, daily mean values were used here for various years between 1994 and 2003 (1994-1996 and 2001-2003). For use in the calibration, the data were aggregated to monthly averages (Fig. 1). For N₂O and NO, we also calculated intra-annual quantiles of the frequency distribution of daily emission magnitudes (10, 50 and 90%). Monthly averages and annual statistics were only calculated for months and calendar years with more than 75% coverage, no gap-filling being applied. The data transformations led to ten different time series of data being available for use in the Bayesian analysis, four with monthly averages and six with annual quantiles, and with a combined number of data points of $n = 225$ (Table 1).

[Fig. 1 HERE]

2.2 Models

Four different deterministic process-based models of forest biogeochemistry were used in this study: BASFOR, COUP, DAYCENT and MoBiLE-DNDC (Table 2).

BASFOR is the simplest model in the group. It was designed to simulate the
interactive effects of changes in N-deposition, atmospheric [CO$_2$] and climate on the carbon balance of forests (Van Oijen et al., 2010a; Van Oijen and Thomson, 2010). The model has been subjected to Bayesian calibration before, using data from the United Kingdom (Van Oijen et al., 2005).

COUP and MoBiLE-DNDC are the two most complex models in the group. Both models were originally designed with special focus on soil processes, but recent versions of the models simulate the whole ecosystem. MoBiLE-DNDC calculates soil microclimate and hydrology, plant growth and plant-soil interactions, biogeochemical processes of the C and N cycle in soils, microbial growth and subsequent trace gas emissions. The core functionality follows the concepts developed in the DNDC suite of models (Li et al., 2000; Li et al., 1992; Werner et al., 2007). COUP was subjected to uncertainty quantification by Klemedtsson et al. (2008) and Svensson et al. (2008). The version of the model used in this paper, referred to as CoupModel, includes an N-flux submodel taken from the PnET-N-DNDC model (Norman et al., 2008). A preliminary uncertainty assessment was also carried out for MoBiLE-DNDC (de Bruijn et al., 2009).

DAYCENT, a model of intermediate complexity, traces its origins to the grassland soil model CENTURY (Parton et al., 1993), but like the previous two models it has developed into a full model for various ecosystems (Del Grosso et al., 2001), of which a small part was subjected to Bayesian calibration before (Yeluripati et al., 2009). The model version used in this paper, referred to as DailyDAYCENT, uses a daily time step for all processes.

2.3 Parameter screening
In the case of the simplest model, BASFOR, no parameter screening was applied, so all its 48 model parameters and initial constants for state variables were included in the Bayesian
calibration. Initial constants were not included in the calibrations of the other three models.

The COUP model was subjected to informal screening, based on previous work involving the same model and experimental site (Norman et al., 2008). The fraction of COUP’s output variability that was caused by uncertainty regarding the selected parameters was not quantified.

Calibration parameters for DAYCENT were selected using Morris screening (Morris, 1991) on average model outputs for soil water content and emissions of N$_2$O, NO and CO$_2$. DAYCENT has over 300 parameters but only 214 were subjected to Morris screening because the majority of the about hundred parameters of the soil water dynamics module were known to contribute about equally to the overall uncertainty, precluding identification of a subset of essential parameters. Morris screening is a global parameter sensitivity analysis, i.e. it explores combinations of parameter values across parameter space rather than just in the neighbourhood of a default parameterisation. Compared to other global sensitivity analyses, the method requires relatively few model runs (proportional to the number of parameters) which permits its use even for models with long runtimes. A subset of 17 DAYCENT parameters was selected in this way. The $r^2$ of the relations between model output variability (from runs where all parameters were varied) and the selected parameters was 0.88-0.98 for the three emission rates and 0.20 for water content, the latter value reflecting the difficulty in selecting key parameters for soil water dynamics in this model.

In the case of MoBiLE-DNDC, uncertainty in the 67 parameters of the soil chemistry submodel was considered. These were mainly parameters used to adapt or scale physical or chemical processes observed by lab studies to real world conditions, semi-empirical ratios, and k-values (decay rates for the various litter and microbial pools of varying decomposability). A substantial number of those parameters also describe Michaelis-Menten kinetics for the microbial turnover processes. The Morris screening method (Morris, 1991)
was applied to MoBiLE-DNDC using the same selection criteria as for DAYCENT, and 26
parameters were selected which together accounted for more than 60% (NO-emission) and
>90% (N₂O- and CO₂-emission, soil water content) of model behaviour.

2.4 Bayesian calibration

Bayesian calibration is the application of probability theory to parameter estimation (Jaynes,
2003; Sivia, 2006). The method finds increasing use in ecological modelling (Ogle, 2009;
Ogle and Barber, 2008; Van Oijen et al., 2005). Uncertainty about parameters is represented
as a joint probability distribution for the possible parameter values. Bayes’ Theorem is used
to determine how this distribution changes in the light of new data:

\[ P(\theta|D) \propto P(\theta) P(D|\theta) \]  (1)

Where P(\theta) and P(\theta|D) are the prior and posterior distributions for the parameters \( \theta \), i.e.
before and after conditioning on the data. The factor that modifies the prior, P(D|\theta), is the
likelihood function, which is the probability of the data for a given \( \theta \). A formal likelihood
function, integrating to unity in data space, needs to be used to be consistent with the
probability calculus, allowing Bayes’ Theorem to be applied (Rougier, (in press)).

The prior probability distribution for the parameters of a model, P(\theta), reflects a
modeller’s uncertainty about parameter values before using the data. This uncertainty is
subjective, and there was no effort in this study to impose any harmonisation on the priors for
the four different models (except for data-scaling factors, discussed later). All modellers
assigned prior distributions that could be written as the product of independent marginal
distributions for individual parameters, but different types of marginal distribution were used:
beta for BASFOR and uniform for the other three models.
The likelihood function $P(D | \theta)$ is the probability of the data $D$ (the 225 data points) given model output generated by parameter vector $\theta$. It accounts for possible measurement error. The same likelihood function was used for all models to allow formal Bayesian model comparison. As described above, the calibration data were in the form of time series of ten different variables, six for annual quantiles and four for monthly averages of daily measurements (Table 2). Estimates for the uncertainty of these variables, both random and systematic, were elicited from the data-providers: co-authors Butterbach-Bahl, Kiese and Werner. Uncertainty about random measurement error is often represented by the use of independent Gaussian distributions for the data points. However, the squared exponential in the Gaussian tends to cause asymptotic collapse of the parameter distribution even with moderate amounts of data (Clark, 2005), and may represent an overestimate of their information content. We therefore used the more heavy-tailed function proposed by Sivia (2006):

$$P(D | \theta) = \prod_{i=1}^{225} \frac{1}{\sigma_i \sqrt{2\pi}} \frac{1 - \exp(-R_i^2 / 2)}{R_i^2}$$  \hspace{1cm} (2)$$

Where $\sigma_i$ is a measure of the uncertainty about random error of the $i$-th data point, and $R_i$ is the difference between model output and $i$-th data point, divided by $\sigma_i$. The values of the $\sigma_i$ were considered to be specific to the type and magnitude of the data points, with relative values of 0.2 for the medians (Q50), 0.3 for the tail-quantiles (Q10 and Q90), and 1.0 for the monthly averages. Besides random measurement error, the data-points were considered to be subject to possible systematic error, which could result from unrepresentative positioning of the soil measurement chambers or errors in instrument calibration. This was implemented by means of four multiplicative data-scaling factors $\gamma_i$, one each for N$_2$O, NO, CO$_2$ and water.
As in other recent studies (Raupach et al., 2005), we treated the $\gamma_i$ as additional parameters to be calibrated. We considered that errors larger than a factor 2 would be very unlikely, but otherwise no assumptions about the systematic data-scaling factors were made. We therefore used an uninformative Jeffrey’s prior (Jaynes, 2003; Sivia, 2006) for the marginal prior distribution for each of the four factors:

$$P(\gamma_i) \propto 1 / \gamma_i$$ (3)

The Jeffrey’s priors for the uncertainty about multiplicative error thus are log-uniform distributions on the interval $[\frac{1}{2}, 2]$. Keats et al. (2009) also used multiplicative errors, for air pollution data in Bayesian calibration of an atmospheric transport model, but used log-normal distributions instead. As Keats et al. (2009) argued, multiplicative error is often the natural choice for measurements of non-negative quantities.

For each of the four models, the Bayesian calibration was carried out by means of Markov Chain Monte Carlo sampling (MCMC), using the Metropolis algorithm (Metropolis et al., 1953; Van Oijen et al., 2005), but model-specific choices were made of proposal distribution and method of testing chain convergence.

Before and after each model’s calibration, a preliminary parameter sensitivity analysis was carried out by calculating the partial correlation coefficients (PCC) for the relationships between individual parameters and the average simulated values of $N_2O$, NO, CO$_2$ and water. In contrast to the ordinary correlation coefficient ($r$), the PCC calculates the association between parameter and output after correcting for the linear effects of the other parameters.

2.5 Bayesian model comparison

The formal Bayesian model comparison consisted of quantifying the relative probabilities of
correctness of the four models, under the assumption that at least one of them was a correct model for the data. The comparison was, like the calibration described above, based on application of Bayes’ Theorem (Kass and Raftery, 1995):

\[ P(M|D) \propto P(M) \frac{P(D|M)}{P(D)} = P(M) \int P(D|M,\theta) P(\theta) \, d\theta \]  

(4)

Where \(P(M)\) and \(P(M|D)\) are the prior and posterior distributions for the models \(M\). In contrast to the parameter distributions, these are discrete distributions, over the four models in our comparison. As shown in the right-hand term, the factor that modifies the prior, \(P(D|M)\), is the integral of the likelihood function over the space spanned by the prior parameter distribution of each model. We refer to this integral as the model’”s „integrated likelihood”. Another common name for this quantity is „marginal likelihood” which expresses the fact that it is found by marginalising out the parameters \(\theta\).

**We a priori** assigned equal probabilities to the different models of being correct, so \(P(M)\) is uniform and the integrated likelihoods represented the relative probability for each model of being correct given the information in the data (Kass and Raftery, 1995). We quantified the integrated likelihoods as follows. For each model, 1000 parameter vectors were drawn from its prior parameter distribution. Comparison of the model outputs for these parameter vectors with the data yielded a sample of 1000 values of the likelihood, and the sample mean was taken as the estimate for the integrated likelihood of the model. Because any sampling-based method is subject to sampling error (McCulloch and Rossi, 1992), we additionally calculated the integrated likelihoods using the method suggested by Kass and Raftery (1995), as the harmonic mean of the sample generated by the MCMC.

### 2.6 Analysis of model-data mismatch
Besides the calculation of the likelihood function, the mismatch between model outputs and measurements was also quantified using more classical means. This was done separately for each of the ten output variables (Table 2) by calculating the Normalised Root Mean Square Error (NRMSE = square root of the average squared difference between model output and data, divided by the average of the data) and the squared correlation coefficient ($r^2$). NRMSE and $r^2$ are distributed quantities, because they depend on the parameterisation, so they were calculated across the range of prior and posterior parameter distributions.

Additional analysis was carried out for just the modes of the prior and posterior parameter distributions. We ran each model with the two modal parameter vectors and calculated the Mean Squared Error (MSE) for each of the four time series of monthly averages. The MSE-values were then decomposed as suggested by Kobayashi and Salam (2000):

$$\text{MSE} = \text{bias}^2 + \text{variance}^2 + 2 \text{bias} \times \text{variance} \times (1 - r)$$

Where $M$ is a simulated time series consisting of monthly averages of $N_2O$, NO, CO$_2$ or water content, $D$ is the matching data, $\sigma_M$ and $\sigma_D$ are their standard deviations, and $r$ is the correlation between the two. The decomposition consists of three terms, which can be interpreted as measures for model-data mismatch due to bias, variance error and phase shift (Kobayashi and Salam, 2000).

3. **Results**

3.1 **Bayesian calibration**
All four models were calibrated using the same MCMC-algorithm, i.e. Metropolis sampling. Burn-in and convergence were determined visually, by each modelling group separately, but an additional analysis of the Markov chains was carried out to confirm that parameter distributions had properly stabilised. The analysis was based on the fact that, after a chain reaches convergence, subsequent distinct and sufficiently long sub-chains should have similar sample means and variances. We compared the first and second halves of the chains after burn-in. The results showed that convergence was adequate for BASFOR and DAYCENT, with all parameters having similar means and variances in the two halves. However, for COUP and MoBiLe-DNDC, some parameters had not stabilised to the same extent, so the posterior parameter distributions for these two models were likely less accurate.

The calibration modified the means and reduced the variances of most marginal distributions. The average variance reduction for process-parameters was small in BASFOR and DAYCENT (3%, 4%), but larger in COUP and MoBiLE (26%, 29%). The data-scaling factors $\gamma_j$ showed greater variance reductions except for soil water content (Fig. 2).

Parameter uncertainty induced output uncertainty. The degree of prior output uncertainty was assessed by determining the quantiles of the output distributions (Table 3). For all models except DAYCENT, prior Q95 was one or two orders of magnitude larger than Q5 for the eight nitrogenous emission variables (Table 3). DAYCENT was already strongly constrained by its prior parameter distribution. For all models, soil respiration was a priori slightly more constrained than the N-emissions, whereas soil water content, which was the only state variable in the set of ten output variables, was most narrowly delimited. Overall, prior ranges were widest for BASFOR. The calibration had only little impact on the output
distributions for soil water content and almost no effect on the soil respiration distributions (Table 3). However, the posterior outputs for the nitrogenous emission variables were much more narrowly constrained than the prior distributions, with posterior Q95/Q5 ratio’s ranging from 2-5 (BASFOR, see also Fig. 1), 2-3 (COUP) and ~1.5 (MoBiLE-DNDC) (Table 3). DAYCENT was the exception with posterior ratio’s that were similar to the prior.

Using the samples from both the prior and posterior parameter distributions, we calculated the partial correlations between individual parameters and outputs. Posterior PCC-values tended to be higher than prior values for BASFOR and COUP, whereas the calibration decreased PCC-values for DAYCENT and MoBiLE-DNDC. However, for all models and output variables, the PCC-based ranking of the parameters changed little, so we restrict ourselves to reporting on the posterior values.

In the case of BASFOR, only one parameter was strongly correlated (|PCC| > 0.5) with N₂O- and NO-emission: the soil water content at which both emissions are equal. Soil respiration was strongly correlated with the parameters that govern decomposition rate of organic matter, and also with the light-use efficiency. Soil water content was mainly correlated with specific leaf area and leaf longevity, both of which affect the active surface area for transpiration. The results for COUP were similar. N-emissions were also mainly correlated with the N₂O-NO balance, soil respiration with decomposition rates and light-use efficiency, and water content with specific leaf area. In the case of DAYCENT, no individual parameters were a posteriori strongly correlated with model outputs, although there had been some strong correlations with the prior parameter distribution (i.e. the NO₃-N₂O conversion efficiency for N-emissions and the leaf area ratio for soil water content). In the case of MoBiLE-DNDC, no parameters were strongly correlated with N₂O-emission, but the Kₘ value for NO₂ did have a high PCC with NO-emission. Soil respiration and water content were both mainly correlated with the parameter that scales decomposition of active organic
substance as a function of soil porosity.

3.2 Bayesian model comparison

The log-transformed integrated likelihood values, calculated from samples from the prior, were as follows (between brackets the values from the alternative calculation using the harmonic mean of the sample from the posterior): BASFOR: -661.7 (-654.7), COUP: -663.5 (-651.2), DAYCENT: -738.5 (-761.2), MoBiLE-DNDC: -657.0 (-758.9). For comparison: a parameter vector whose model outputs would have gone exactly through the 225 data points, would have had a log-likelihood of -581.2. Both methods of calculating the integrated likelihoods showed that the data provided greater support for BASFOR and COUP than for DAYCENT. The two estimates of the integrated likelihood for MoBiLE-DNDC differed strongly, so it is less clear how plausible this model is.

3.3 Analysis of model-data mismatch

First, the data were compared with the ranges of model output uncertainty induced by the parameters. All time series averages of measurements were in the central intervals of the prior output distributions, between the 5% and 95% quantiles, except for soil respiration as predicted by BASFOR and MoBiLE-DNDC, and the lower quantiles of daily N$_2$O emission rates as predicted by COUP (Table 3).

Although the distributions of simulated time series averages were found to cover the data fairly well, inspection of the time series themselves revealed considerable differences between models and measurements (Fig. 3). For example, none of the models was able to reproduce the large peak in N$_2$O-emissions in early 1996 after a strong freeze-thaw event (Papen and Butterbach-Bahl, 1999), neither with the mode of the prior, not with the posterior mode (Fig. 3). This is likely to be a consequence of incomplete process representation in the
models, although a more recent version of MoBiLE-DNDC showed a possible way to account for the freeze-thaw effect (de Bruijn et al., 2009). Despite these remaining differences between model outputs and data, overall the calibration was able to remove much of the mismatch for the N-emissions and, to lesser extent, for CO$_2$-emission (compare the lower „posterior” panels of Fig. 3 to the upper „prior” ones, and see also the increased likelihoods depicted in Fig. 4). Note that some of the reduced bias in the posterior results was due to the impact of calibration on data-scaling factors (Fig. 2) rather than on model parameters. There was no apparent improvement in the simulation of soil water content, which reflected the fact that for this variable the least amount of information was available (Table 2). In one specific case, simulation of soil water by model DAYCENT using the mode of its posterior distribution, bias was increased relative to the prior mode (Fig. 3), but this result was not representative of the full posterior distribution for the water data-scaling factor ($\gamma_{\text{WATER}}$) of this model (Fig. 2). However, it does suggest that a useful – but for this model computationally demanding - additional step in the procedure would have been to determine the mode of the posterior distribution by targeted optimisation, rather than relying on the parameter sample generated by the MCMC.

[Fig. 3 HERE]

Whereas for the prior output distributions of the models most time-series averages were in the central Q5-Q95, the same did not apply to the posterior distributions. Most time-series averages were to be found in the upper tails (>Q95) of the posterior output distributions (Table 3, and compare also the examples for BASFOR in Fig. 1). There were differences, however, in how the likelihood distributions responded to the calibration, as can be seen from the posterior distributions of likelihoods (Fig. 4).
For each model and each parameter vector sampled from the prior and posterior distributions, we compared the simulated time series of the ten output variables with the corresponding data, by calculating the correlation coefficient ($r$) and the normalised root mean square error (NRMSE). Each parameter distribution thus induced ten different distributions of $r$ and NRMSE, of which we show the 5, 50 and 95% quantiles (Table 4). The posterior values of the quantiles for $r$ are often not improvements over the prior, except for Q5. So calibration tended to remove only the parameter vectors with the poorest output-data correlation. In contrast, NRMSE was improved for almost every quantile of every variable in each model (Table 4).

The MSE-decompositions for time series with monthly averages, both for the prior and posterior parameter modes, are shown in Fig. 5. Phase shift, variance error and bias were reduced to different extent for the different models.

4. Discussion

4.1 Bayesian calibration: methodological issues

Bayesian calibration uses data to update the joint probability distribution for a model’s parameters. The Bayesian approach allows for non-Gaussian distributions for both parameter uncertainty and measurement error. Our calibration was therefore based on sampling by means of MCMC rather than on matrix inversion methods. This in turn allowed us to include
systematic data error in the calibration, rather than having to estimate error terms in a first
separate step, as was done for example by Michalak et al. (2005), using maximum-likelihood
estimation.

Although the theory is straightforward, it is easy to overestimate the information
content of any dataset, and this may lead to unsupported changes in the parameter
distributions. For example, when the common assumption is made that each new data point
adds independent new information to the calibration, the parameter distributions will
asymptotically collapse with sample size (Clark, 2005). Modellers thus need to elicit realistic
assessments of measurement uncertainty from the data-providers (Moala and O'Hagan,
2010). This issue was important in the case-study presented here because the dataset was
fairly large (n=225), covering times series of four variables. We applied four techniques to
ensure a realistic, albeit subjective assessment of the information content of the data: (1)
using the monthly temporal scale as the one at which the models were supposed to be
applicable, together with the frequency distribution of daily emission events, (2) allowing for
random errors in the data, (3) allowing for systematic errors in the data by the use of the four
scaling factors, (4) using a heavy-tailed likelihood function (Sivia, 2006). The adjustment of
temporal scale is a common technique in atmospheric physics, applied whenever models
produce more smooth results than measurements and thereby induce apparent correlations
between measurement errors (Prinn, 2000). We considered the implementation of these four
techniques to be partly the responsibility of the data-providers (in the case of random and
systematic errors) and partly that of the modellers and data-providers together (temporal scale
and likelihood function). Using this approach, parameter uncertainties were reduced
markedly but the distributions did not collapse. The techniques applied are generic and may
be widely applicable to calibration of complex dynamic models using long time series.
4.2 Bayesian calibration: impact on parameter uncertainty of the forest models

Parameter uncertainties were reduced strongly compared to the prior, and the likelihood distributions were shifted toward higher values for all four models (Fig. 4). The degree of uncertainty reduction varied between the models, as did the balance between changing data-scaling parameters and process parameters. Although our dataset included measurements of the three major biogeochemical cycles (nitrogen, carbon and water), there was still some lack of balance because about 75% of the data points were for emissions of N₂O and NO (Table 1). Therefore, most of the improvement of model behaviour (reduction of likelihood and NRMSE, increase in $r$) was for these variables. For all models, the parameters that were changed the most were related to the soil nitrogen dynamics.

The results of our preliminary parameter sensitivity analysis, consisting of calculating partial correlations with the different outputs, need to be interpreted with care. A high value of the PCC for a specific parameter-output combination suggests that the parameter – within its range of uncertainty - strongly affects the output. Therefore knowledge about the process governed by that parameter is key to understanding variability in the output. The opposite may not be true: a strong but non-linear effect may yield a low PCC. Also, the importance of a parameter is not an intrinsic property: it depends on the distribution of that parameter. Whenever Bayesian calibration reduces the variance of a parameter, the contribution of that parameter to output variability is expected to decrease. However, PCC-analysis is not a variance-decomposition method (Saltelli et al., 2000), so it may not be able to show that effect, and indeed, for two out of the four models posterior values of PCC were generally larger than the prior values.

With these caveats, the results of the PCC-analysis did reveal commonalities between the models. Across the models, N-emissions were mainly correlated with N₂O-NO partitioning, soil respiration mainly with decomposition but also tree productivity, and soil
water content with leaf area dynamics. These agreements between the models suggest that
some of the differences between complex process-based models may not overly affect their
behaviour. The models compared here all represent, albeit in very different ways, the linkages
between the ecosystem C-, N- and H₂O-cycles. Therefore there are inevitable similarities in
the overall feedback structure of the models, imposed by constraints of stoichiometry and
mass-balance of the three biogeochemical cycles. These similarities may outweigh details of
process representation and parameterisation (Van Oijen et al., 2004; Van Oijen et al., 2010b).

The prior PCC-values were, as expected, indicative of which parameters were most
informed by the data in the subsequent calibration. For all four models, the relative decrease
of marginal variance tended to be greatest for those parameters that had a strong a priori
correlation with N₂O-emission, the output variable for which the largest number of data
points were available. The parameter variance reduction accounted for by the prior PCC-
values ranged from 25% (COUP, MoBiLE) to 29% (BASFOR) and even 35% (DAYCENT).
PCC-analysis might thus play a useful role in parameter screening before calibration.

4.3 Bayesian model comparison

Comprehensive model comparison requires taking into account parameter uncertainty. A
complex model might, in principle, be able to predict complex biogeochemical time series
more closely than a simple model. But if it is unclear what the parameterisation of the
complex model for good prediction should be, then its predictive capacity is reduced. A
simple model whose parameters are well-known might then perform better. Regarding model
complexity, there is a trade-off between the need to represent the intricacies of the real world
and the need to minimise parameter uncertainty. We thus need a method for comparing the
behaviour of the models not just at the modes of the parameter distributions or at the
maximum likelihood estimates, but across their whole parameter distributions. Bayesian
model comparison is such a method. This method has been used before in environmental
modelling, in the elegant study by Tuomi et al. (2008) who compared different functions
describing the impact of temperature on soil respiration, but to our knowledge this is the first
application to complex process-based ecosystem models. Formally, the relative magnitudes
of the integrated likelihoods equate to relative probabilities for the individual models of being
correct, conditional on a correct model being present in the comparison. However, in
environmental modelling all models are incorrect in some way, so we prefer to use the
integrated likelihoods as a guide towards plausible model structures rather than as
probabilities of correctness (Gelman and Shalizi, 2010; Kass and Raftery, 1995).

Bayesian model comparison requires that a common likelihood function is used with
all models – because the criterion for comparison is the integrated likelihood of the models,
where the integration is over the prior parameter distribution. Therefore only those data can
appear in the likelihood function, which are part of the output set of each model. For
example, in the current study, we could not include vertical profiles of soil temperature in the
likelihood function because the simplest model BASFOR has only one soil compartment.
Three of the models had remarkably similar values of the integrated likelihood, the exception
being the low value for DAYCENT, which was thus identified as the least plausible model.
There was only a slight preference for MoBiLE-DNDC. We analysed these prior integrated
likelihood values further by considering the underlying distributions of likelihoods associated
with the four different categories of output variable ($\text{N}_2\text{O}$, NO, respiration, water) (Fig. 4).
The lower value of the integrated likelihood for DAYCENT can be seen to be mainly due to
poorer performance for $\text{N}_2\text{O}$. The similarity between the integrated likelihoods for the other
models was seen to extend to the underlying distributions of category-wise likelihoods (Fig.
4).
4.4 Analysis of model-data mismatch

The NRMSE and $r$ statistics provided useful additional information beyond the formal Bayesian model comparison. When going from prior to posterior, $r$ did not improve (apart from Q5) in any of the models, but NRMSE improved throughout (Table 4). The calibration thus was more successful in reducing the average magnitude of the differences between simulations and measurements, than in aligning the distribution of the outputs over time. There clearly remain difficulties for all models in simulating the large interannual variation in nitrogenous emission characteristics. The models were also similar in that the posterior NRMSE was lowest for soil water content and highest for the monthly values of $\text{N}_2\text{O}$-emission (Table 4). It was easier to simulate water than nitrogen dynamics.

The decomposition of the MSE for the modes of the prior and posterior parameter distributions gave further information. The MSE-decomposition is only possible for long time series, i.e. the monthly data (Table 1). There were not enough annual quantiles available to allow the reliable estimation of the variance and phase shift terms. The calibration reduced the MSE for the parameter modes of all four models, and all four categories of output variables (Fig. 5), confirming the effectiveness of the calibration. However, the analysis revealed large differences between the models. The simplest model, BASFOR, had the highest variance error for $\text{N}_2\text{O}$ and NO. This suggests that a simple model may not be able to respond quickly enough to changes in the environment that affect nitrogenous emissions. The low integrated likelihood of DAYCENT has already been attributed to poor simulation of $\text{N}_2\text{O}$-emission (Fig. 4), and the MSE-decomposition showed that this was mainly due to a large phase shift for $\text{N}_2\text{O}$ emission (Fig. 5). In fact, DAYCENT had very low bias and variance error for $\text{N}_2\text{O}$, so it was able to capture general characteristics (mean, „peakiness“) of the time series of emission very well, but not the timing of emission events.

Most data were in the central Q5-Q95 ranges of the prior output distributions of the
models (Table 3). However, perhaps surprisingly, most data were to be found in the upper tails (>Q95) of the posterior output distributions (Table 3). This is because of a trade-off between the different variables in the calibration. Such trade-off is inevitable with models that are imperfect and cannot capture the whole range of behaviour of all variables simultaneously. Moreover, there is also the distinct possibility of systematic error in the measurement of the different categories of output variables. Where the models were unable to reconcile all the data, the calibration tended to modify the settings of the scale parameters (Fig. 2). It is important to establish whether the likely underlying cause of the revision of the scale factors was indeed systematic measurement error or model structural error. One way of attempting this is to consider the differences between the models in their posterior estimates for the scaling factors. A posteriori, all models suggested that the measurements for CO₂-emissions were unrealistically high (Fig. 2: γ_{CO₂} mostly <1), but only the BASFOR model suggested the same for the N-emissions (γ_{N2O} and γ_{NO} <<1). There thus is some doubt about the CO₂-measurements, and likewise about the capacity of BASFOR to simulate N-dynamics.

Note that, in our approach, the data-scaling factors are intended to represent the idiosyncrasies of specific datasets, so we cannot expect the calibrated scaling values to apply elsewhere. However, if calibration of a model using data from multiple sites were shown to consistently lead to scaling factors different from unity, we would expect the error to be predominantly the model’s rather than that of the data. Here we only had data from one site available, so no strong conclusions can be drawn.

4.5 The Bayesian framework

The three-operation Bayesian framework proposed here - calibration, comparison, analysis of model-data mismatch – was shown to work well in this study. Most of the techniques employed in each of these operations are novel in their application to complex dynamic
models of forest biogeochemistry, in particular in combination with the use of data from processes that vary strongly over time (Luo et al., 2009). The application of the framework showed that model parameter uncertainty could be reduced in all models, irrespective of their level of complexity. However, it also showed that the models still suffer from structural deficiencies, even if we allow for the possibility of errors in the data, and that the deficiencies are stronger for the nitrogen cycle than for the carbon and water cycles.

There are limitations associated with this study and a caveat has to be made regarding the use of the results. We only used data from one site, Höglwald, and the general applicability of the models to European forests needs to be tested with data from other sites. When data from these new sites become available, the posterior distributions found here will become prior distributions in further calibration. Another limitation of the study was the use of parameter screening, which was considered necessary for the three models with the highest computational demand, but there may be environmental conditions under which simulated forest biogeochemistry is sensitive to the excluded parameters. Also, uncertainty concerning environmental drivers such as weather conditions was ignored as it was expected to be small compared to the structural and parameter uncertainties. This assumption needs to be verified.

Wang et al. (2009) called for the development of an integrated Bayesian framework that can account for the different sources and types of error arising in environmental modelling. The Bayesian framework proposed here is an integrated one in the sense that its three operations were linked methodologically and in that its three operations provide complementary information. Methodologically, the Bayesian calibration made use of the same likelihood function as the Bayesian model comparison. Calibration was also linked to comparison in the use of MCMC to explore parameter space, with the thus generated sample being used in the model comparison for estimating the integrated likelihood. Because this estimate, based on the harmonic mean of the sampled likelihoods, can be unstable (Chib and
Jeliazkov, 2001), we also used the method of directly sampling from the prior. Methodological links further existed between the calibration and the analysis of model-data mismatch, in that the sample generated by the calibration was used to calculate the NRMSE across the posterior parameter distribution, rather than just for one parameter vector.

More importantly than these methodological links, the three operations in the framework also complemented each other in how they help improve the modelling. Calibration reduced parameter uncertainty, model comparison reduced uncertainty about the relative plausibility of the different models, and the analysis of model-data mismatch showed which parts of the models needed most improvement, which in this case was the nitrogen dynamics.

5. Conclusions

- Bayesian calibration can be used to reduce parametric uncertainty of complex dynamic models for forest biogeochemistry.
- Bayesian calibration allows for the use of datasets that contain long time series of gas emissions with high intra- and interannual variability, and with both random and systematic error.
- Data need to be compared with models at the appropriate temporal scale. This may involve, as shown here, monthly averaging and the calculation of annual frequency distributions. These transformations, and the use of heavy-tailed likelihood functions that account for uncertainty about random and systematic measurement errors, can help prevent collapse of the parameter distributions in the calibration.
- Bayesian model comparison can be used to calculate the relative conditional probabilities of models being correct, irrespective of the type and complexity of the
considered models.

- Bayesian model comparison treats models as black boxes, so it can only identify which models are implausible, but it cannot identify any specific model deficiencies.
- Analysis of model-data mismatch can help identify model weaknesses by decomposition of the MSE and by showing how the NRMSE and the correlation coefficient $r$ vary for the different processes simulated by the models.
- Together, the three operations of Bayesian calibration, Bayesian model comparison, and analysis of model-data mismatch, constitute a promising framework for uncertainty reduction and improvement of complex dynamic models in forest biogeochemistry.

- This was confirmed by the case-study analysed here, in which four different parameter-rich process-based models of forest biogeochemistry were confronted with long time-series of biogeochemical data. Parameter uncertainties were reduced in all models and the relative model plausibilities were quantified, with MoBiLE-DNDC having a slight preference over the other models. The simplest model, BASFOR, was shown to underestimate variance of nitrogenous emissions even after calibration. The model of intermediate complexity, DAYCENT, simulated the time series well but with large phase shift. COUP and MoBiLE-DNDC were able to remove most bias through calibration.

- The calibration not only reduced parameter and model uncertainty, but also identified possible systematic error in the measurement of soil respiration, to which all models assigned data-scaling factors less than unity with high posterior probability.
- There remain patterns in the data - in particular infrequent events of very high nitrogenous emission rate - that are unexplained by any of the models, even after calibration. Given the intensive exploration of parameter space in the calibration, this
is unlikely to be due to incorrect model parameterisation.

- The analysis showed that the models still suffer from structural deficiencies, even if we allow for the possibility of errors in the data. The deficiencies are stronger for the nitrogen cycle than for the carbon and water cycles.

Acknowledgements

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References


Del Grosso, S.J. et al., 2001. Simulated interaction of carbon dynamics and nitrogen trace gas fluxes using the


Table 1. Overview of data used for calibration. Variables are annual quantiles of daily emission rates where indicated, and monthly averages otherwise. The period of measurement indicates the years of first and last measurement, and \( n \) is the total number of data points over that period. The columns marked \( \text{min} \), \( \text{mean} \) and \( \text{max} \) show the extreme values and the mean of the \( n \) values.

<table>
<thead>
<tr>
<th>Variable</th>
<th>Unit</th>
<th>Period of measurement</th>
<th>( n )</th>
<th>( \text{min} )</th>
<th>( \text{mean} )</th>
<th>( \text{max} )</th>
</tr>
</thead>
<tbody>
<tr>
<td>( \text{N}_2\text{O} ) (Q10)</td>
<td>kg N ha(^{-1}) y(^{-1})</td>
<td>1994-2003</td>
<td>6</td>
<td>0.08</td>
<td>0.21</td>
<td>0.31</td>
</tr>
<tr>
<td>( \text{N}_2\text{O} ) (Q50)</td>
<td>kg N ha(^{-1}) y(^{-1})</td>
<td>1994-2003</td>
<td>6</td>
<td>0.27</td>
<td>0.52</td>
<td>0.85</td>
</tr>
<tr>
<td>( \text{N}_2\text{O} ) (Q90)</td>
<td>kg N ha(^{-1}) y(^{-1})</td>
<td>1994-2003</td>
<td>6</td>
<td>0.55</td>
<td>2.32</td>
<td>9.14</td>
</tr>
<tr>
<td>NO (Q10)</td>
<td>kg N ha(^{-1}) y(^{-1})</td>
<td>1994-2002</td>
<td>5</td>
<td>1.65</td>
<td>2.33</td>
<td>2.79</td>
</tr>
<tr>
<td>NO (Q50)</td>
<td>kg N ha(^{-1}) y(^{-1})</td>
<td>1994-2002</td>
<td>5</td>
<td>4.81</td>
<td>6.64</td>
<td>8.30</td>
</tr>
<tr>
<td>NO (Q90)</td>
<td>kg N ha(^{-1}) y(^{-1})</td>
<td>1994-2002</td>
<td>5</td>
<td>10.65</td>
<td>13.65</td>
<td>18.52</td>
</tr>
<tr>
<td>( \text{N}_2\text{O} )</td>
<td>kg N ha(^{-1}) y(^{-1})</td>
<td>1994-2003</td>
<td>70</td>
<td>0.03</td>
<td>0.99</td>
<td>16.55</td>
</tr>
<tr>
<td>NO</td>
<td>kg N ha(^{-1}) y(^{-1})</td>
<td>1994-2003</td>
<td>61</td>
<td>1.74</td>
<td>7.72</td>
<td>24.04</td>
</tr>
<tr>
<td>Rsoil</td>
<td>mg C m(^{-2}) h(^{-1})</td>
<td>1995-1997</td>
<td>36</td>
<td>23.9</td>
<td>113.3</td>
<td>255.2</td>
</tr>
<tr>
<td>Water</td>
<td>% (vol)</td>
<td>1994-1996</td>
<td>25</td>
<td>27.1</td>
<td>33.6</td>
<td>37.0</td>
</tr>
</tbody>
</table>
Table 2. Overview of the models.

<table>
<thead>
<tr>
<th>Property</th>
<th>BASFOR</th>
<th>COUP</th>
<th>DAYCENT</th>
<th>MoBiLE-DNDC</th>
</tr>
</thead>
<tbody>
<tr>
<td># State variables (trees, soil)</td>
<td>14 (6,8)</td>
<td>56 (8,48)</td>
<td>20 (10,10)</td>
<td>38 (10,28)</td>
</tr>
<tr>
<td># Parameters (in calibration)</td>
<td>48 (48)</td>
<td>&gt;300 (23)</td>
<td>&gt;300 (17)</td>
<td>67* (26)</td>
</tr>
<tr>
<td>Time step</td>
<td>Daily</td>
<td>Hourly</td>
<td>Daily</td>
<td>Daily (but less than 1 minute for diffusion processes)</td>
</tr>
<tr>
<td>Inputs: Environmental time series</td>
<td>Radiation, temperature, precipitation, humidity, wind speed, N-deposition, [CO₂]</td>
<td>Radiation, temperature, precipitation, humidity, wind speed, N-deposition</td>
<td>Radiation, temperature, precipitation, humidity, wind speed, N-deposition</td>
<td>Radiation, temperature, precipitation, N-deposition, [CO₂]</td>
</tr>
<tr>
<td>Inputs: Environmental constants</td>
<td>Soil water retention curve, rooting depth</td>
<td>Soil water retention curve, rooting depth</td>
<td>Soil water retention curve, rooting depth</td>
<td>Layer-specific texture, bulk density, field capacity, pH</td>
</tr>
</tbody>
</table>

* For the soil chemistry module only.
Table 3. Summary of model output, with prior and posterior uncertainties. All values shown refer to time series averages, to be compared with the last-but-one column of Table 1. For each model, the table entries are three quantiles (5, 50 and 95%) of the output distributions generated by the prior and posterior parameter distributions. In **bold**: posterior distributions for which the posterior width (Q5-Q95) is at least an order of magnitude less than for the prior.

<table>
<thead>
<tr>
<th>Var.</th>
<th>Unit</th>
<th>Dist.</th>
<th>BASFOR</th>
<th>COUP</th>
<th>DAYCENT</th>
<th>MoBiLE-DNDC</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td>Q5</td>
<td>Q50</td>
<td>Q95</td>
<td>Q5</td>
</tr>
<tr>
<td>$N_2O$ (Q10)</td>
<td>kg N ha$^{-1}$ y$^{-1}$</td>
<td>Prior</td>
<td>0.01</td>
<td>0.05</td>
<td>1.78</td>
<td>0.01</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Post.</td>
<td><strong>0.02</strong></td>
<td><strong>0.03</strong></td>
<td><strong>0.04</strong></td>
<td><strong>0.03</strong></td>
</tr>
<tr>
<td>$N_2O$ (Q50)</td>
<td>kg N ha$^{-1}$ y$^{-1}$</td>
<td>Prior</td>
<td>0.03</td>
<td>0.18</td>
<td>3.83</td>
<td>0.02</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Post.</td>
<td><strong>0.07</strong></td>
<td><strong>0.12</strong></td>
<td><strong>0.17</strong></td>
<td><strong>0.06</strong></td>
</tr>
<tr>
<td>$N_2O$ (Q90)</td>
<td>kg N ha$^{-1}$ y$^{-1}$</td>
<td>Prior</td>
<td>0.12</td>
<td>1.13</td>
<td>8.08</td>
<td>0.16</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Post.</td>
<td><strong>0.29</strong></td>
<td><strong>0.46</strong></td>
<td><strong>0.65</strong></td>
<td><strong>0.11</strong></td>
</tr>
<tr>
<td>NO (Q10)</td>
<td>kg N ha$^{-1}$ y$^{-1}$</td>
<td>Prior</td>
<td>0.03</td>
<td>0.10</td>
<td>8.99</td>
<td>3.42</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Post.</td>
<td><strong>0.05</strong></td>
<td><strong>0.11</strong></td>
<td><strong>0.18</strong></td>
<td><strong>1.60</strong></td>
</tr>
<tr>
<td>NO (Q50)</td>
<td>kg N ha$^{-1}$ y$^{-1}$</td>
<td>Prior</td>
<td>0.09</td>
<td>0.45</td>
<td>12.21</td>
<td>5.95</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Post.</td>
<td><strong>0.25</strong></td>
<td><strong>0.79</strong></td>
<td><strong>1.43</strong></td>
<td><strong>2.41</strong></td>
</tr>
<tr>
<td>NO (Q90)</td>
<td>kg N ha$^{-1}$ y$^{-1}$</td>
<td>Prior</td>
<td>0.31</td>
<td>2.38</td>
<td>19.99</td>
<td>10.56</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Post.</td>
<td><strong>0.99</strong></td>
<td><strong>2.69</strong></td>
<td><strong>4.57</strong></td>
<td><strong>4.41</strong></td>
</tr>
<tr>
<td>$N_2O$</td>
<td>kg N ha$^{-1}$ y$^{-1}$</td>
<td>Prior</td>
<td>0.05</td>
<td>0.39</td>
<td>4.42</td>
<td>0.07</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Post.</td>
<td><strong>0.11</strong></td>
<td><strong>0.18</strong></td>
<td><strong>0.24</strong></td>
<td><strong>0.08</strong></td>
</tr>
<tr>
<td>NO</td>
<td>kg N ha$^{-1}$ y$^{-1}$</td>
<td>Prior</td>
<td>0.15</td>
<td>0.92</td>
<td>13.25</td>
<td>6.05</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Post.</td>
<td>0.43</td>
<td>1.16</td>
<td>1.99</td>
<td>2.96</td>
</tr>
<tr>
<td>CO₂</td>
<td>mg C m$^{-2}$ h$^{-1}$</td>
<td>Prior</td>
<td>44.2</td>
<td>69.4</td>
<td>106.8</td>
<td>76.5</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Post.</td>
<td>56.2</td>
<td>76.5</td>
<td>99.5</td>
<td>67.6</td>
</tr>
<tr>
<td>Water (vol)</td>
<td>%</td>
<td>Prior</td>
<td>28.3</td>
<td>31.6</td>
<td>33.7</td>
<td>33.2</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Post.</td>
<td>28.7</td>
<td>31.0</td>
<td>33.0</td>
<td>33.3</td>
</tr>
</tbody>
</table>
Table 4. Comparison of data with model outputs: correlation coefficient ($r$) and normalised root mean square error (NRMSE). The table shows quantiles (Q5, Q50, Q95) of the distributions of $r$ and NRMSE induced by prior and posterior parameter distributions. In **bold**: posterior values that are improvements over the prior ($r$ increased, NRMSE reduced).

<table>
<thead>
<tr>
<th>Var.</th>
<th>Dist.</th>
<th>Statistic</th>
<th>BASFOR</th>
<th>COUP</th>
<th>DAYCENT</th>
<th>MoBiLE-DNDC</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td>Q5</td>
<td>Q50</td>
<td>Q95</td>
<td>Q5</td>
</tr>
<tr>
<td>N$_2$O (Q10)</td>
<td>Prior</td>
<td>$r$</td>
<td>-0.37</td>
<td>0.42</td>
<td>0.72</td>
<td>-0.83</td>
</tr>
<tr>
<td></td>
<td></td>
<td>NRMSE</td>
<td>0.44</td>
<td>1.07</td>
<td>11.95</td>
<td>0.41</td>
</tr>
<tr>
<td></td>
<td>Post.</td>
<td>$r$</td>
<td>-0.75</td>
<td>-0.19</td>
<td>0.49</td>
<td>-0.60</td>
</tr>
<tr>
<td></td>
<td></td>
<td>NRMSE</td>
<td><strong>0.41</strong></td>
<td><strong>0.47</strong></td>
<td><strong>0.53</strong></td>
<td><strong>0.31</strong></td>
</tr>
<tr>
<td>N$_2$O (Q50)</td>
<td>Prior</td>
<td>$r$</td>
<td>-0.46</td>
<td>0.65</td>
<td>0.88</td>
<td>-0.62</td>
</tr>
<tr>
<td></td>
<td></td>
<td>NRMSE</td>
<td>0.40</td>
<td>1.02</td>
<td>7.64</td>
<td>0.35</td>
</tr>
<tr>
<td></td>
<td>Post.</td>
<td>$r$</td>
<td>-0.53</td>
<td>-0.04</td>
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<td>-0.73</td>
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Fig. 1. Dots: Monthly averages of measured emissions of N$_2$O, NO, CO$_2$ and of soil water content. The lines represent output of model BASFOR. Dashed red line: output for the mode of the prior parameter distribution. Thick black line: output for the mode of the posterior. Thin black lines: 5% and 95% quantiles of the posterior output distribution.

Fig. 2. Posterior marginal distributions for the four data-scaling factors that represent systematic multiplicative error in the data according to the different models.

Fig. 3. Differences between measurements and simulations (positive values indicating underestimates by the models) for time series with monthly averages. Top 4 panels: simulations using the mode of the prior parameter distribution. Bottom 4 panels: simulations with the posterior mode.

Fig. 4. Distributions of log-likelihoods for each of the four models, for the four categories of output variables. Grey: prior, black: posterior.

Fig. 5. Decomposition of the Mean Squared Error (MSE) associated with the modes of the prior and posterior parameter distributions, for the time series with monthly data. The MSE-values for the N$_2$O and NO are in the same units (kg N ha$^{-1}$ y$^{-1}$ squared), the
MSE-values for soil respiration and soil water content are in squared mg C m\(^{-2}\) h\(^{-1}\) and squared %, respectively.
Figure 1

- N$_2$O emission (kg N ha$^{-1}$ yr$^{-1}$)
  - Obs
  - Prior Mode
  - Post. Mode
  - Post. Q95, Q5

- NO emission (kg N ha$^{-1}$ yr$^{-1}$)

- CO$_2$ emission (mg C m$^{-2}$ h$^{-1}$)

- Water in soil (% vol)
Figure 2

N\textsubscript{2}O

NO

CO\textsubscript{2}

WATER

Legend:
- BASFOR
- COUP
- DAYCENT
- MoBiLE
Figure 3

N₂O emission (kg N ha⁻¹ yr⁻¹)

CO₂ emission (mg C m⁻² h⁻¹)

Water in soil (% vol)

N₂O emission (kg N ha⁻¹ yr⁻¹)

NO emission (kg N ha⁻¹ yr⁻¹)

Water in soil (% vol)
Figure 4: BASFOR, COUP, DAYCENT, MoBiLE

N$_2$O emission

NO emission

CO$_2$ emission

Water in soil