

## RESEARCH ARTICLE

# Soil enzymes as indicators of soil function: A step toward greater realism in microbial ecological modeling

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## Abstract

Soil carbon (C) and nitrogen (N) cycles and their complex responses to environmental changes have received increasing attention. However, large uncertainties in model predictions remain, partially due to the lack of explicit representation and parameterization of microbial processes. One great challenge is to effectively integrate rich microbial functional traits into ecosystem modeling for better predictions. Here, using soil enzymes as indicators of soil function, we developed a competitive dynamic enzyme allocation scheme and detailed enzyme-mediated soil inorganic N processes in the Microbial-ENzyme Decomposition (MEND) model. We conducted a rigorous calibration and validation of MEND with diverse soil C-N fluxes, microbial C:N ratios, and functional gene abundances from a 12-year CO<sub>2</sub> × N grassland experiment (BioCON) in Minnesota, USA. In addition to accurately simulating soil CO<sub>2</sub> fluxes and multiple N variables, the model correctly predicted microbial C:N ratios and their negative response to enriched N supply. Model validation further showed that, compared to the changes in simulated enzyme concentrations and decomposition rates, the changes in simulated activities of eight C-N-associated enzymes were better explained by the measured gene abundances in responses to elevated atmospheric CO<sub>2</sub> concentration. Our results demonstrated that using enzymes as indicators of soil function and validating model predictions with functional gene abundances in ecosystem modeling can provide a basis for testing hypotheses about microbially mediated biogeochemical processes in response to environmental changes. Further development and applications of the modeling framework presented here will enable microbial ecologists to address ecosystem-level questions beyond empirical observations, toward more predictive understanding, an ultimate goal of microbial ecology.

## KEYWORDS

elevated CO<sub>2</sub>, functional traits, metagenomics, microbial ecological modeling, microbial functional genes, nitrogen enrichment, predictive ecology, soil enzymes

## 1 | INTRODUCTION

Projecting future carbon cycling and climate change scenarios is a grand challenge in ecology and for society (Cavicchioli et al., 2019). Microorganisms, acting as detritivores, plant symbionts, or pathogens, are critical in mediating ecosystem carbon (C) and nutrient cycling and consequently climate change (Bardgett et al., 2008; Cavicchioli et al., 2019). However, traditional biogeochemical and Earth system models (ESMs) do not explicitly consider the roles of microbial communities by assuming that microbes are in equilibrium with their environment (Schimel, 2013). Such classical models appear to work well for large-scale patterns of bulk soil organic matter pools, but they may have reached their limits, particularly when depicting transient dynamics in the face of environmental changes (Wieder et al., 2015). In the last decade, a considerable amount of effort has been devoted to explicitly integrating microbial communities and functions into microbial ecological models (e.g., Allison et al., 2010; Davidson et al., 2012; Manzoni et al., 2016; Schimel & Weintraub, 2003; Sulman et al., 2018; Tang & Riley, 2019; Wang et al., 2013; Wieder et al., 2015). Studies have shown that microbial-explicit models could more accurately represent the impacts of global change drivers, such as warming and priming effects (Wieder et al., 2015). This calls for more mechanistic microbial ecological models to advance our understanding of soil microbial and biogeochemical responses to environmental changes.

Ecosystem models with carbon–nitrogen (C–N) coupled processes have elucidated substantial impacts on the carbon–climate feedbacks that are lacking from the C-only models, for instance, smaller sensitivity of land C uptake to temperature variation or increasing atmospheric CO<sub>2</sub> concentration (Thornton et al., 2007). N availability is known to strongly influence microbial growth and C cycling (Cavicchioli et al., 2019; Treseder, 2008); hence, multiple microbial-explicit models have accounted for C–N interactions (e.g., Abramoff et al., 2017; Drake et al., 2013; Gao et al., 2020; Kyker-Snowman et al., 2020; Schimel & Weintraub, 2003). However, limited attention has been paid to the explicit representation and parameterization of multiple differential microbial groups, particularly related to the inorganic N cycle (e.g., N mineralization and immobilization, nitrification and denitrification) (Sulman et al., 2018; Wang et al., 2019). This impedes a comprehensive validation of complex C–N processes and their interactions as have been done for classical terrestrial C–N coupled models. Therefore, the introduction of mechanistic inorganic N cycling into microbial ecological models may provide new opportunities to pose and validate further hypotheses about coupled C–N cycling in response to environmental perturbations, especially elevated atmospheric CO<sub>2</sub> concentration (eCO<sub>2</sub>) and enhanced N deposition (Abramoff et al., 2017; Wieder et al., 2015).

The absence of microbial communities in ecosystem models is primarily due to the extremely high diversity and complexity of microbial communities and the lack of appropriate strategies and frameworks for using microbial information in ecological modeling (Bailey et al., 2018; Bardgett et al., 2008; Gao et al., 2020; Todd-Brown et al., 2012; Wieder et al., 2015). Because microbial communities under natural settings are extremely diverse and complex, functional traits-based approaches are very attractive and promising for explicitly accounting for the role of microbes in regulating biogeochemical cycles in ecosystem models (Falkowski et al., 2008; Klausmeier et al., 2020). However, one big challenge is how to extract and scale functional information to inform ecosystem modeling (Torsvik & Øvreås, 2002). This challenge has also become an important motivation to develop microbially explicit models (Bailey et al., 2018). Despite increasing interest in incorporating microbial functional traits into ecosystem models, it remains a major challenge to directly link genomes to global processes (Bailey et al., 2018). However, it is viable to link genomes and processes at intermediate scales with integrated applications of powerful analytical and modeling techniques (Song et al., 2017; Trivedi et al., 2013). While representing a massive number of microbial taxa in models is impractical and unnecessary, owing to functional redundancy (Bailey et al., 2018), grouping microbes and enzymes into simplified functional guilds is feasible and enables the parameterization of microbial ecological models (Chen & Sinsabaugh, 2021; Song et al., 2017).

However, it remains challenging to develop microbially explicit N transformation processes. First, the multi-step inorganic N reactions are regulated by intracellular enzymes that are located at cell membrane, cytoplasm, or periplasm (Fiencke & Bock, 2006; Schlesier et al., 2016; Song et al., 2017). These intracellular N enzymes differ from extracellular enzymes (e.g., ligninases and cellulases) and have little capability of acting on their own, leading to the concern in representing them in microbial ecological models. Second, an effective microbial or enzyme allocation scheme is warranted to handle diverse microbial communities associated with the multiple inorganic N processes. For instance, we have recently used GeoChip-based gene abundances (Shi et al., 2019) to constrain the Microbial-ENzyme Decomposition (MEND) model, where we only considered extracellular C-degrading enzymes owing to the lack of detailed representation of microbially mediated inorganic N reactions (Gao et al., 2020; Guo et al., 2020). In short, modeling efforts have not kept pace with the rapid advances in the microbial ecology of N relevant microorganisms and genes (Hu et al., 2015).

Model parameterization through calibration and validation with field observations is arduous due to the limited available long-term experimental data and large uncertainties in measurements of the state variables, fluxes, and microbial community structure and

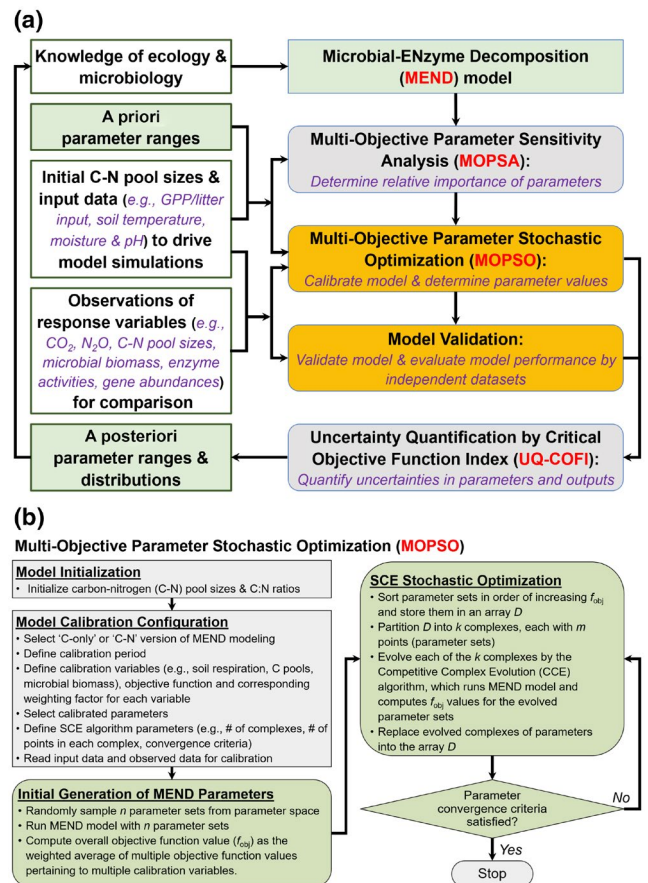
functions, as well as uncertainties in model structure and simulations (Bradford et al., 2016; Sulman et al., 2018). Consistent with these large uncertainties in observations and model simulations, recent comparison of five soil C models with different representation of microbial and mineral processes revealed that existing traditional measurements (e.g., CO<sub>2</sub> fluxes and soil C contents) were insufficient to constrain or validate ecosystem models (Sulman et al., 2018). To demonstrate the capability of microbially explicit models, development of benchmarking with multiple datasets with a variety of microbial and omics data, especially for inorganic N cycling, is needed.

In this study, building on past work (Gao et al., 2020), we attempted to improve the MEND model by developing a new microbially mediated inorganic N module that uses relevant enzymes as indicators of soil function, with the proposition of a competitive dynamic enzyme allocation scheme. The new inorganic N module accounts for the important roles of intracellular enzymes in regulating several critical inorganic N transformations, including N fixation, nitrification, and the sequential denitrification reactions from nitrate (NO<sub>3</sub><sup>-</sup>) to dinitrogen (N<sub>2</sub>) (Xue et al., 2016; Zhou et al., 2012). In addition to several important observations (soil respiration, soil concentrations of ammonium and nitrate, and abundances of two functional gene groups targeting soil organic matter [SOM] decomposition) used in Gao et al. (2020), the new MEND model was further calibrated and validated with a variety of new data from that 12-year field experiment, called BioCON (Gao et al., 2020), including SOC content, multiple inorganic N transformation rates, and the abundances of six functional gene groups important to inorganic N processes. We directly compared model outputs to the relative changes of the measured gene abundances in response to eCO<sub>2</sub>. Our results indicated that explicitly using enzymes as soil function indicators in ecosystem models and validating model predictions with gene abundance data can provide a basis for better understanding and testing hypotheses about microbially mediated biogeochemical processes under environmental changes.

## 2 | MATERIALS AND METHODS

### 2.1 | Overview of the MEND modeling framework

We developed an integrated microbial ecological modeling framework, consisting of several key components such as model development, sensitivity analysis, model calibration, validation, and uncertainty quantification (Figure 1a). The new MEND model explicitly represents distinct microbial and enzyme groups responsible for C-N transformation processes. The Multi-Objective Parameter Sensitivity Analysis (MOPSA) was used to determine the relative importance of parameters in terms of multiple response objectives (i.e., variables). The sensitivity analysis forms the cornerstone of the Multi-Objective Parameter Stochastic Optimization (MOPSO) and validation procedure. The MOPSO approach aims to determine the values of those “free” parameters by calibrating the model against observations (Figure 1b), where a stochastic optimization algorithm,



**FIGURE 1** Framework for developing the Microbial-ENzyme Decomposition (MEND) model. (a) MEND modeling framework. (b) Procedure of the Multi-Objective Parameter Stochastic Optimization (MOPSO), which directly incorporates the Shuffled Complex Evolution (SCE) algorithm into MEND model calibration

the Shuffled Complex Evolution (SCE) (Duan et al., 1992), is modularized and incorporated into the MEND model for automatically calibrating parameters. The SCE algorithm combines the strengths of several optimization strategies such as controlled random search, complex shuffling, and competitive evolution, which ensure that the parameter space is efficiently and thoroughly exploited (Duan et al., 1992; Wang et al., 2015). The MOPSO enables to fit multiple observational variables (soil respiration, C pools, microbial biomass, etc.) by minimizing the overall objective function as the weighted average of multiple objectives pertaining to these variables (Figure 1b). We further validated the model and evaluated model performance against datasets not used for model calibration. The Uncertainty Quantification by Critical Objective Function Index (UQ-COFI) approach was developed to filter the parameter sets generated by the MOPSO procedure. These filtered parameter sets by UQ-COFI represented the posterior parameter space, which were used to drive multiple model runs to quantify uncertainties in response variables due to parametric uncertainty. We employed this integrated modeling framework to guide reliable model development and application.

A detailed description of model sensitivity analysis (MOPSA) and uncertainty quantification (UQ-COFI) are presented in Supporting

Information Sections 3.3 and 3.6, respectively. In the following, we overview the new MEND model and its calibration and validation against multiple datasets.

## 2.2 | New MEND model with a competitive dynamic enzyme allocation scheme

We incorporated a new N-associated module into the old MEND (MEND-old) model (Gao et al., 2020; Wang et al., 2021) (Figure S1b; for comparison, the new MEND model [MEND-new] is shown in Figure S1a, a copy of Figure 2) by explicitly representing (a) multiple key intracellular enzymes as indicators that catalyze nitrification, sequential denitrification, and nitrogen fixation processes; (b) plant-microbial competition for inorganic N ( $\text{NH}_4^+$  and  $\text{NO}_3^-$ ); (c) ammonium ( $\text{NH}_4^+$ ) sorption; (d) nitrate ( $\text{NO}_3^-$ ) and nitrite ( $\text{NO}_2^-$ ) leaching; and (e) N gases ( $\text{NO}$ ,  $\text{N}_2\text{O}$ , and  $\text{N}_2$ ) exchange between soil and the atmosphere. A reaction rate in the model may be modified by soil pH, soil temperature, and moisture conditions (Figures S2–S4). Details on MEND-new model and its state variables, governing equations, component fluxes and parameters are described in Supporting Information Sections 1–3 and Tables S1–S6.

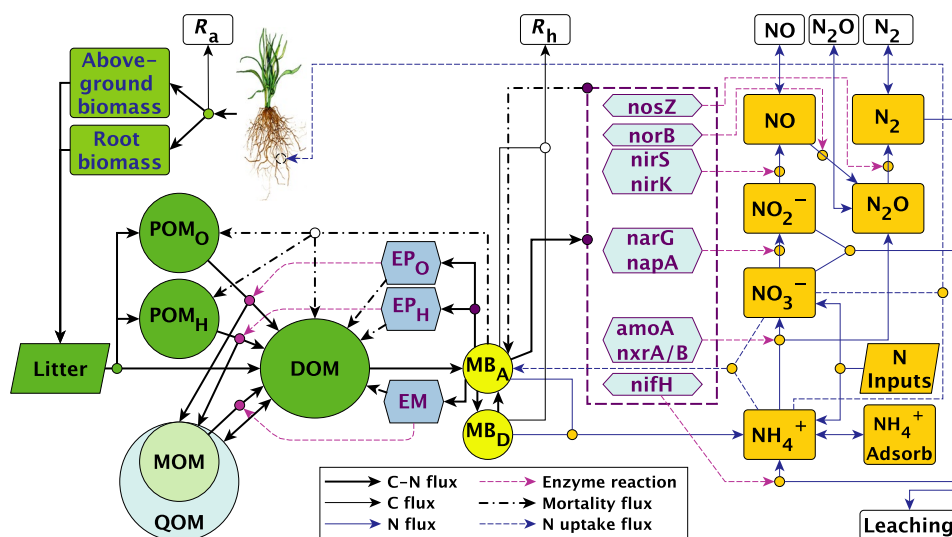
We used flexible stoichiometry (i.e., time-variant C:N ratio) for SOM and microbial biomass pools to represent microbial adaptation in response to the stoichiometric imbalance of available resources (Du et al., 2018; Fanin et al., 2017; Mooshammer et al., 2014a, 2014b; Zechmeister-Boltenstern et al., 2015). In addition to the three SOM-degrading enzyme functional groups, that is,  $\text{EP}_\text{O}$ ,  $\text{EP}_\text{H}$ , and EM, we

incorporated six new enzyme systems as indicators controlling inorganic N transformations (Figure 2), that is, nitrogenases (corresponding to functional genes of *nifH*), ammonia oxidases (*amoA*), nitrate reductases (*narG/napA*), nitrite reductases (*nirS/nirK*), nitric oxide reductases (*norB*), and nitrous oxide reductases (*nosZ*) (Xue et al., 2016; Zhou et al., 2012).

We proposed a competitive dynamic enzyme allocation scheme to deal with the synthesis of multiple enzyme groups (see Supporting Information Section 1.1.7). The enzyme allocation approach developed here is based on the synthetic results that enzyme activities are dependent on microbial biomass (Jian et al., 2016) and substrate availability (Sinsabaugh et al., 2014). A competitive allocation scheme is applied to the production of enzymes for each inorganic N transformation process, where the competitive allocation coefficient is the saturation level of an inorganic N substrate (i.e., the ratio of the substrate concentration to the corresponding half-saturation constant) relative to the sum of the saturation levels of all inorganic N substrates (Equation S40).

## 2.3 | Model calibration and validation

We implemented the MOPSO approach, based on the SCE algorithm (Duan et al., 1992; Wang et al., 2015), to calibrate selected model parameters according to the sensitivity analysis (Figure 1b). We aimed to determine parameter values and their uncertainties by achieving high goodness-of-fits of model simulations against experimental observations, such as soil respiration ( $R_s$ ), microbial heterotrophic respiration ( $R_h$ ), microbial biomass carbon (MBC), as well as soil C and N pools



**FIGURE 2** Diagram of the Microbial-ENzyme Decomposition (MEND) model.  $R_a$  and  $R_h$  are autotrophic and heterotrophic respiration, respectively.  $\text{POM}_\text{O}$  and  $\text{POM}_\text{H}$  are particulate organic matter (POM) decomposed by oxidative ( $\text{EP}_\text{O}$ ) and hydrolytic enzymes ( $\text{EP}_\text{H}$ ), respectively. MOM is mineral-associated OM, which is decomposed by a mixed enzyme group EM. Dissolved OM (DOM) interacts with the active layer of MOM (QOM) through sorption and desorption. Litter enters  $\text{POM}_\text{O}$ ,  $\text{POM}_\text{H}$ , and DOM. Microbes consist of active ( $\text{MB}_\text{A}$ ) and dormant microbes ( $\text{MB}_\text{D}$ ). DOM can be assimilated by  $\text{MB}_\text{A}$ . Inorganic N deposition and fertilization enter  $\text{NH}_4^+$  and  $\text{NO}_3^-$  that can be immobilized by microbes and taken up by plant roots.  $\text{NH}_4^+$  adsorption is also considered. N fixation, nitrification, and denitrification are mediated by nitrogenases (*nifH*), ammonia oxidases (*amoA*, *nxrA/B*), and N-reductases (*narG/napA*, *nirS/nirK*, *norB*, and *nosZ*), respectively. Inorganic N loss pathways include leaching ( $\text{NO}_3^-$  and  $\text{NO}_2^-$ ) and gas emission ( $\text{NO}$ ,  $\text{N}_2\text{O}$ , and  $\text{N}_2$ ) from the soil to the atmosphere

and fluxes. Each objective evaluates the goodness-of-fit of a specific observed variable. The parameter optimization attempts to minimize the overall objective function ( $J$ ) that is computed as the weighted average of multiple single-objectives (see Equation 67 in Supporting Information Section 3.4). Generally, equal weights are assigned to these objectives. However, a higher weight is recommended for a variable that is frequently measured or is vital to the research topic.

Different objective functions were used to quantify the goodness-of-fit for different variables (Supporting Information Section 3.4), depending on the measurement method and frequency. As per model validation (Refsgaard, 1997), we used datasets that were not involved in model calibration to evaluate model performance, where the same calibrated parameter values were used in model validation.

## 2.4 | BioCON datasets for model calibration and validation

Since there is no gold standard for validating model performance, it is a common practice to use published datasets in ecosystem and bioinformatic studies, which have advantages for model calibration and validation (Luo et al., 2012; Ning et al., 2020). Thus, we used experimental data (Table 1) from a well-designed, long-term multifactor free-air CO<sub>2</sub> enrichment experiment, BioCON (Biodiversity, CO<sub>2</sub>, and N deposition) (45.4010° N, 93.2010° W) in Minnesota, USA (Reich & Hobbie, 2013). The BioCON experiment aims, among other goals, to elucidate how microbe-mediated feedbacks to soil respiration are affected by N addition (+4 g N m<sup>-2</sup> yr<sup>-1</sup>) and elevated atmospheric CO<sub>2</sub> (eCO<sub>2</sub>, +180 ppm) (Adair et al., 2009, 2011). The BioCON soil is an Entisol, more specifically, a mixed, frigid Lamellic Udipsamments as per the USDA soil taxonomy (O'Geen et al., 2017; Soil Survey Staff, 1999). This excessively drained soil, derived from glacial outwash with a coarse structure, has very poor development and a sandy texture (92–94% sand and 2–3% clay in the top 114 cm) (Kazanski et al., 2021; O'Geen et al., 2017). In summary, there were four CO<sub>2</sub> × N treatments among 296 plots: ambient atmospheric CO<sub>2</sub> & ambient N supply (aCO<sub>2</sub>-aN), eCO<sub>2</sub>-aN, aCO<sub>2</sub> & enriched N supply (aCO<sub>2</sub>-eN), and eCO<sub>2</sub>-eN with each treatment having 74 plots (biological replicates). To examine the effects of plant diversity on ecosystem N cycling, the BioCON experiment also has (at each level of CO<sub>2</sub> and N) treatment plots planted with either 1, 4, 9, or 16 grassland species (Dijkstra et al., 2007).

Estimates of daily GPP (gross primary production) values were obtained from a corrected 8-day GPP product based on the MODIS GPP (MOD17A2/MOD17A2H) (Gao et al., 2020; Zhu et al., 2018) and used to drive model simulations under the control treatment (aCO<sub>2</sub>-aN). The GPP for the other three treatments was rescaled according to the general linear relationship between NPP (net primary production) and GPP (Gao et al., 2020). Meanwhile, environmental datasets measured in each CO<sub>2</sub> × N treatment were also used for model simulations for each treatment, including monthly soil pH, daily soil temperature, and moisture.

Soil samples for microbial community analysis were collected from the 296 plots in August 2009. Each sample was a composite of five

soil cores from each plot at a depth of 0–15 cm. Microbial DNA was extracted, hybridized with GeoChip arrays, and analyzed as described previously (Guo et al., 2020; Tu et al., 2014). The eCO<sub>2</sub> effect on the abundance of each functional gene (total abundance of all probes of this gene) was examined by the response ratio (Luo et al., 2006):

$$RR = \ln(x_T/x_C) \quad (1)$$

where RR is the response ratio (effect size) that quantifies the log-proportional change between the gene abundances of eCO<sub>2</sub> ( $x_T$ ) and aCO<sub>2</sub> ( $x_C$ ) samples.

The observed response ratios (RRs) between the gene abundances ( $GA_{obs}$ ) of eCO<sub>2</sub> and aCO<sub>2</sub> were used as additional data to evaluate model-simulated enzyme concentrations ( $EC_{sim}$ ), enzyme activities ( $EA_{sim}$ ), or equivalent first-order reaction rates ( $FR_{sim}$ ). As the Michaelis–Menten kinetics is used in the MEND model, the relationships among  $EC_{sim}$ ,  $EA_{sim}$ , and  $FR_{sim}$  are described as follows:

$$EA_{sim} = Vd \cdot EC_{sim} \quad (2)$$

$$FR_{sim} = (Vd \cdot EC_{sim}) / (K + S) = EA_{sim} / (K + S) \quad (3)$$

where  $EC_{sim}$  (mg C cm<sup>-3</sup>),  $EA_{sim}$  (mg C cm<sup>-3</sup> h<sup>-1</sup>), and  $FR_{sim}$  (h<sup>-1</sup>) are simulated enzyme concentration, enzyme activity, and the equivalent first-order reaction rate, respectively;  $S$  denotes the substrate (e.g., SOC) concentration; and the parameters  $Vd$  and  $K$  represent the specific enzyme activity (mg C mg<sup>-1</sup> C h<sup>-1</sup>) and the half-saturation constant (mg C cm<sup>-3</sup>), respectively.

In summary, nine C-N response variables were involved in the calibration of MEND-new (Table 1): soil CO<sub>2</sub> flux ( $R_g$ ), microbial biomass C (MBC), soil organic C (SOC), ammonium (NH<sub>4</sub><sup>+</sup>), nitrate + nitrite (NO<sub>3</sub><sup>-</sup> + NO<sub>2</sub><sup>-</sup>), as well as the reference rates of net N mineralization (FN<sub>mn-im</sub>), nitrification (FN<sub>nit</sub>), biological N fixation (FN<sub>fix</sub>), and plant N uptake (FN<sub>im-VG</sub>). Among these variables, the literature-reported biological N fixation rates (including both symbiotic and non-symbiotic N fixation) (Cleveland et al., 1999, 2013) and plant N uptake rates (Bessler et al., 2012; Harty et al., 2017; Reyes et al., 2015) in grasslands were used as reference for model calibration. To examine the predictive power of the model, we only calibrated the model against the data under the control treatment (aCO<sub>2</sub>-aN) and then applied the calibrated parameters to the other three treatments for model validation. To further investigate the model's capability in representing microbial and enzyme functional traits, we directly validated the model against literature-reported microbial C:N ratios (Xu et al., 2013) and the measured response ratios of gene abundances ( $GA_{obs}$ ).

## 3 | RESULTS

Detailed results of model sensitivity analysis and uncertainty quantification are presented in Supporting Information Results 5.1 (with Figure S5) and 5.2 (with Figure S6), respectively. In the following, we



TABLE 1 BioCON data for MEND model calibration and validation

Response variable	Description	Objective Function	Number of data points
$R_s$ ( $\text{CO}_2$ )	Soil respiration = root respiration ( $R_a$ ) + microbial respiration ( $R_h$ )	$J_1 = R^2$	284
MBC	Microbial biomass carbon	$J_2 = \text{MAREt}$ , tolerance = 0.1	1
SOC	Soil organic carbon	$J_3 = \text{MAREt}$ , tolerance = 0.05	1
$\text{NH}_4^+$	Ammonium concentration	$J_4 = 0.8 \times  \text{PBIAS}  + 0.2 \times \text{MARE}$	8
$\text{NO}_3^- + \text{NO}_2^-$	Nitrate + Nitrite concentration	$J_5 = 0.8 \times  \text{PBIAS}  + 0.2 \times \text{MARE}$	8
$\text{FN}_{\text{mn-im}}$	Net N mineralization rate	$J_6 = \text{MAREt}$ , tolerance = 0.5	10
$\text{FN}_{\text{nit}}$	Nitrification flux rate	$J_7 = \text{MAREt}$ , tolerance = 0.9	10
$\text{FN}_{\text{fix}}$	N fixation flux rate	$J_8 = \text{MAREt}$ , tolerance = 0.2	1
$\text{FN}_{\text{im,VG}}$	Plant uptake rate of N	$J_9 = \text{MAREt}$ , tolerance = 0.5	1
EPO	Oxidative enzyme	For model validation only:	1
EPH	Hydrolytic enzyme	Compare simulated and observed Response Ratios (RR).	1
ENH4	Ammonium oxidase	Observed RR is the response ratio of omics-detected gene abundances between elevated $\text{CO}_2$ ( $\text{eCO}_2$ ) and ambient $\text{CO}_2$ ( $\text{aCO}_2$ ).	1
ENO3	Nitrate reductase		1
ENO2	Nitrite reductase	Simulated RR is the response ratio of MEND-modeled enzyme concentrations, activities, or reaction rates between $\text{eCO}_2$ and $\text{aCO}_2$ .	1
ENO	Nitric oxide reductase		1
EN2O	Nitrous oxide reductase		1
EN2	Nitrogenase		1

Notes:  $R^2$  denotes the coefficient of determination,  $|\text{PBIAS}|$  is the absolute value of the percent bias, MARE is the mean absolute relative error, MAREt is the MARE with a tolerance. See Supporting Information Equations (68–71) for a description of these criteria.

focus on the key results with respect to model calibration, validation, and ecological insights.

### 3.1 | Model calibration and validation of soil respiration and inorganic N processes

#### 3.1.1 | Model calibration and validation strategy in terms of the BioCON data

Based on the aforementioned sensitivity analysis and previous studies on the MEND model (Wang et al., 2013, 2015, 2019), we selected 14 important parameters (Figure S6) to conduct model calibration.

In the first step of calibration, we calibrated nine microbial physiological parameters by achieving high goodness-of-fits of model simulations against experimental observations, such as soil respiration ( $R_s$ ), microbial biomass carbon (MBC), and soil organic carbon (SOC) (Table 1). We only compared the simulated mean values of MBC and SOC to the observed reference MBC and SOC, respectively, as we only had observations at one time point for each of them. In the overall objective function (Equation 67 in Supporting Information Section 3.4), the weights of 0.50, 0.25, and 0.25 were assigned to the objectives pertaining to  $R_s$ , MBC, and SOC, respectively, owing to far more data points available for  $R_s$  (284 data points) than for MBC and SOC. The nine parameters (Table S5) included the following: (a) six parameters ( $V_g$ ,  $\alpha$ ,  $K_D$ ,  $Y_g$ ,

$kY_g$ , and  $\gamma$ ) for microbial growth, maintenance, and mortality; and (b) three parameters ( $p_{EP}$ ,  $fp_{EM}$ ,  $r_E$ ) for enzyme production, turnover, and decomposition of SOM.

As for the second step of calibration, we fixed the parameter values determined by the first step and calibrated five important inorganic N parameters (Table S5:  $\text{VN}_{\text{fix}}$ ,  $\text{VN}_{\text{nit}}$ ,  $\text{VN}_{\text{denit}}$ ,  $\text{VN}_{\text{plant}}$ , and  $Q_{\text{maxNH}_4}$ ) by fitting observed concentrations of ammonium ( $\text{NH}_4^+$ ) and nitrate + nitrite ( $\text{NO}_3^- + \text{NO}_2^-$ ), as well as the reference rates of net N mineralization ( $\text{FN}_{\text{mn-im}}$ ), nitrification ( $\text{FN}_{\text{nit}}$ ), biological N fixation ( $\text{FN}_{\text{fix}}$ ), and plant N uptake ( $\text{FN}_{\text{im,VG}}$ ) (Table 1). In the overall objective function, higher weights were used for the objectives of  $\text{NH}_4^+$  and  $\text{NO}_3^- + \text{NO}_2^-$  than the remaining N variables. As a result, there were nine individual objective functions regarding the nine C-N response variables in the model calibration: the first three objective functions were used for the calibration of microbial physiological parameters and the remaining six variables were used for the parametrization of inorganic N transformation parameters (Table 1).

The model simulation period covered the 12-year observational period (1998–2009). Model simulations for each treatment were driven by the corresponding data: GPP, soil temperature and moisture, and inorganic N ( $\text{NH}_4^+$  and  $\text{NO}_3^-$ ) input. We used the MOPSO approach to calibrate model parameters with the data from the  $\text{aCO}_2$ -aN treatment. We then validated the model using the same set of model parameters calibrated for  $\text{aCO}_2$ -aN to simulate  $R_h$  and  $R_s$ , and soil inorganic N in the other three treatments ( $\text{eCO}_2$ -aN,  $\text{aCO}_2$ -eN, and  $\text{eCO}_2$ -eN).

### 3.1.2 | Model calibration and validation results of soil respiration

Our model calibration with aCO<sub>2</sub>-aN data achieved good agreement between simulated and observed soil respiration (Figure 3a, R<sup>2</sup> = 0.60), so did the model validation of soil respiration in the other three treatments (Figure 3b, R<sup>2</sup> = 0.56–0.61). In addition, the percent bias (|PBIAS|) values of mean soil respiration were 3% for calibration and 3–11% for validation, suggesting that simulated mean soil respiration values were close to the observed ones in all four treatments. The simulated mean values of MBC and SOC were within the tolerances for MBC (10%) and SOC (5%), respectively, as expected in model simulations (Table 1).

### 3.1.3 | Model calibration and validation results of soil ammonium and nitrate

In addition, the simulated mean soil NH<sub>4</sub><sup>+</sup> and (NO<sub>3</sub><sup>-</sup>+NO<sub>2</sub><sup>-</sup>) concentrations also agreed well with the observations in both model calibration and validation (Figure 3c,d). Although model validation showed larger percent bias between simulated and observed values (|PBIAS| = 24–29% for NH<sub>4</sub><sup>+</sup> and 5–39% for NO<sub>3</sub><sup>-</sup>+NO<sub>2</sub><sup>-</sup>) than model calibration (2% for both NH<sub>4</sub><sup>+</sup> and 8% for NO<sub>3</sub><sup>-</sup>+NO<sub>2</sub><sup>-</sup>), the model validation of inorganic N concentrations could still be judged as satisfactory according to the 70% bias criterion for N modeling (Moriassi et al., 2007). Simulated variation (i.e., average standard deviation (SD) = 0.20 gN m<sup>-2</sup>) in soil NH<sub>4</sub><sup>+</sup> concentrations by the MEND-new model was also comparable to observed variation (average SD = 0.15 gN m<sup>-2</sup>), which was also true for soil NO<sub>3</sub><sup>-</sup>+NO<sub>2</sub><sup>-</sup> (average SD = 0.074 and 0.070 gN m<sup>-2</sup> for observed and simulated concentrations, respectively). For comparison, the simulated average SD values by the MEND-old model were 0.072 and 0.095 gN m<sup>-2</sup> for soil NH<sub>4</sub><sup>+</sup> and NO<sub>3</sub><sup>-</sup>+NO<sub>2</sub><sup>-</sup>, respectively.

Generally, the simulated mean NH<sub>4</sub><sup>+</sup> and NO<sub>3</sub><sup>-</sup> concentrations by MEND-new from this study showed much lower biases than MEND-old with simplified N processes as described in Gao et al. (2020), except for the NO<sub>3</sub><sup>-</sup> validation under eCO<sub>2</sub>-aN. The average |PBIAS| for NH<sub>4</sub><sup>+</sup> was reduced from 45% (MEND-old, 12–68% in range) to 21% (MEND-new, 2–28%), though the average |PBIAS| values for NO<sub>3</sub><sup>-</sup> were similar between MEND-old (11–32% with an average of 18%) and MEND-new (5–39% with an average of 18%) (Figure 3c,d). To account for the effects of the number of free (i.e., calibrated) model parameters on the model performance, we calculated the Akaike information criterion (AIC) of the two models (Goll et al., 2012). The number of free model parameters for inorganic N processes was five for MEND-old (Gao et al., 2020) and six for MEND-new (see Table S5), as most of the N-related parameters in MEND-new were determined as per literature. Compared to MEND-old, MEND-new had a slightly higher AIC under aCO<sub>2</sub>-aN (Figure 3c), but lower AIC under the other three treatments (Figure 3d).

### 3.1.4 | Model calibration and validation results of inorganic N fluxes

Biological N fixation and plant N uptake rates during model calibration and validation were generally in accordance with literature-reported data (Figure 4). The simulated biological N fixation rates in all four treatments were comparable to the ranges for grasslands reported in the literature (Cleveland et al., 1999, 2013). The N fixation rates were significantly higher under the two eN treatments compared to those under the aCO<sub>2</sub>-aN treatment (Figure 4a). However, we did not observe statistically significant eCO<sub>2</sub> effects on the N fixation rates. Our simulated plant N uptake rates were generally between 15 and 30 g N m<sup>-2</sup> yr<sup>-1</sup>, which were within the range (10–40 g N m<sup>-2</sup> yr<sup>-1</sup>) observed in grasslands (Bessler et al., 2012; Harty et al., 2017; Reyes et al., 2015). The plant N uptake rates were significantly lower under aCO<sub>2</sub>-aN than those under the other three treatments, with the highest under eCO<sub>2</sub>-eN and no significant difference between eCO<sub>2</sub>-aN and aCO<sub>2</sub>-eN or eCO<sub>2</sub>-eN (Figure 4b).

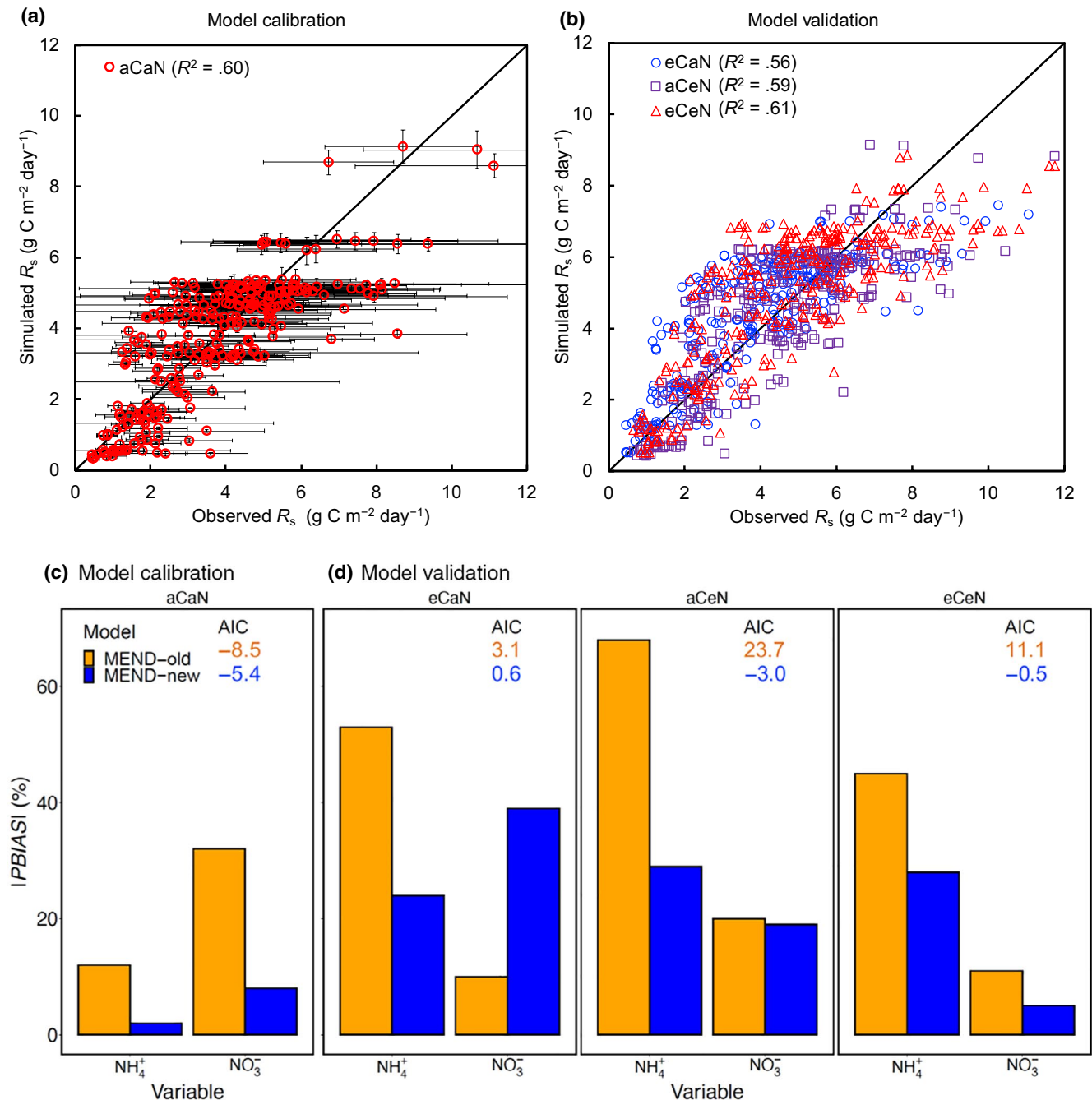
The simulated net N mineralization and nitrification rates were within the observed ranges in both model calibration and validation (Figure S7). As mentioned in the methods, we did not expect simulated values to match the measured nitrification rates and net N mineralization rates as they represented reference rates or rough estimates. Our simulated net N mineralization rates were 57–85% (with a mean of 68%) of the reference rates, with the lowest simulated actual N mineralization rate under aCO<sub>2</sub>-aN and the highest under the two eCO<sub>2</sub> treatments (Figure S7a). The simulated nitrification rates accounted for 39–54% (with a mean of 47%) of the reference values, with the lowest under the two ambient N treatments and the highest under the two enriched N treatments (Figure S7b).

## 3.2 | Model validation of microbial C:N ratios

Independent model validation showed that the microbial C:N ratios simulated by MEND-new conformed to the literature-reported mean value and the 95% confidence interval for grassland soils (Xu et al., 2013), whereas MEND-old predicted much higher microbial C:N ratios (Figure 5a). Though both models predicted lower microbial C:N under eN than aN (Figure 5b,c), only the MEND-new model revealed a statistically significant decrease in the microbial C:N as a result of N addition (Figure 5c). However, neither model demonstrated significant eCO<sub>2</sub> effect on the microbial C:N ratios (Figure S8).

## 3.3 | Model validation with functional gene abundances

We first compared the eCO<sub>2</sub> effects on enzymes simulated by the two models, that is, MEND-old and MEND-new. To make the results comparable between the two models, gene abundances were not included in the calibration of MEND-old, matching what we did for MEND-new in this study. We only examined the oxidative enzymes

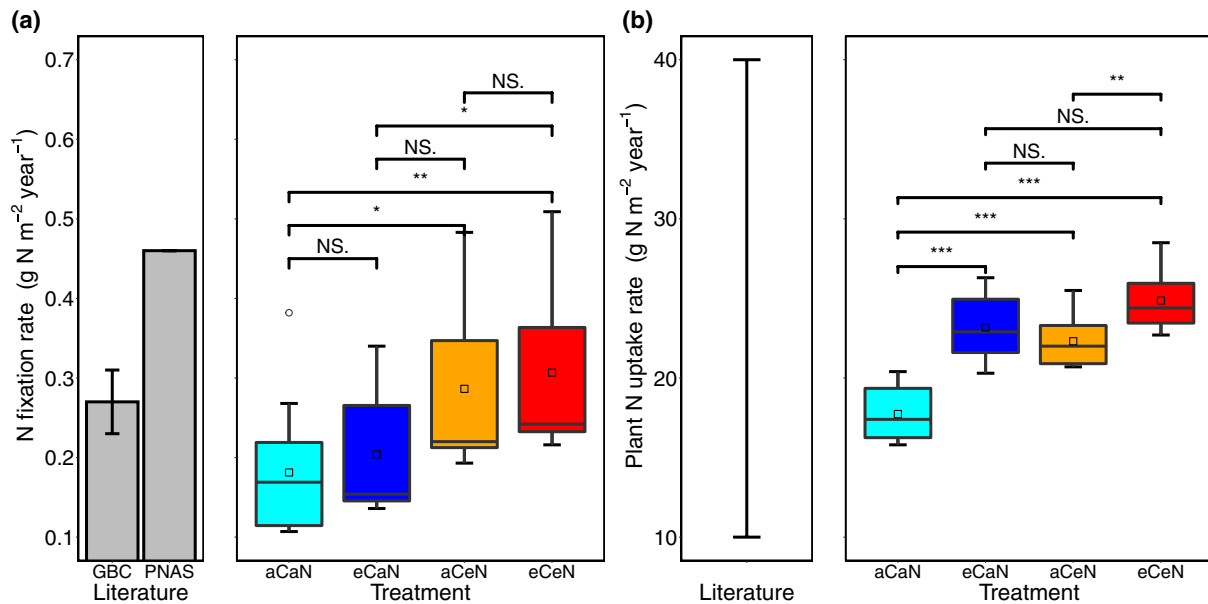


**FIGURE 3** MEND model calibration and validation. (a) Soil respiration ( $R_s$ ) calibration with ambient  $\text{CO}_2$ -ambient N (aCaN) data, (b)  $R_s$  validation with data from the other three treatments: elevated  $\text{CO}_2$ -aN (eCaN), aC-enriched N (aCeN), and eCeN. (c) Absolute value of percent bias ( $|PBIAS|$ , %) between simulated and observed mean for the calibration of ammonium ( $\text{NH}_4^+$ ) and nitrate ( $\text{NO}_3^-$ , including both  $\text{NO}_3^-$  and  $\text{NO}_2^-$ ) from aCaN. (d)  $|PBIAS|$  for the validation of  $\text{NH}_4^+$  and  $\text{NO}_3^-$  from the other three treatments. Error bars in A represent the standard deviations.  $R^2$  values in A and B denote the coefficient of determination. In (c) and (d), the two models of MEND-old and MEND-new denote the old version of MEND model as described in Gao et al. (2020) and the new MEND model in this study, respectively. The two numbers in each facet of (c) and (d) denote the Akaike information criterion (AIC, lower is better) for the two models, respectively

(Figure 5d) and hydrolytic enzymes (Figure 5e) that are associated with the C cycle because only these two groups are included in both models. The response ratios (RRs) of simulated enzyme concentrations ( $\text{EC}_{\text{sim}}$ ), enzyme activities ( $\text{EA}_{\text{sim}}$ ), and the first-order reaction rates ( $\text{FR}_{\text{sim}}$ ) by MEND-old were significantly higher than the

response ratios of observed gene abundances ( $\text{GA}_{\text{obs}}$ ). The simulated response ratios by MEND-new were generally lower than those by MEND-old, except for the  $\text{FR}_{\text{sim}}$  of hydrolytic enzymes under eN and the  $\text{FR}_{\text{sim}}$  of oxidative enzymes. In short, compared to MEND-old, the simulated response ratios by MEND-new were generally closer





**FIGURE 4** Comparison between simulated rates and literature-reported nitrogen flux rates. (a) Biological N fixation rate; the “Literature” data were from Cleveland et al. (1999, GBC) and Cleveland et al. (2013, PNAS), where the bars show the mean values and the error bar shows the value range. (b) Plant N uptake rate; the “Literature” data were from Bessler et al. (2012) and Reyes et al. (2015), where the error bar denotes the value range. The difference in simulated rates between paired treatments was tested by the Wilcoxon signed-rank test. “\*”, “\*\*\*”, and “\*\*\*\*” denote significant difference with  $p$ -value  $< .05$ ,  $p$ -value  $< .01$ , and  $p$ -value  $< .001$ , respectively. “NS.” means not significant

to the measured values. Particularly, only the  $EA_{sim}$  by MEND-new correctly reflected the negative response in the oxidative enzymes observed under eN (Figure 5d).

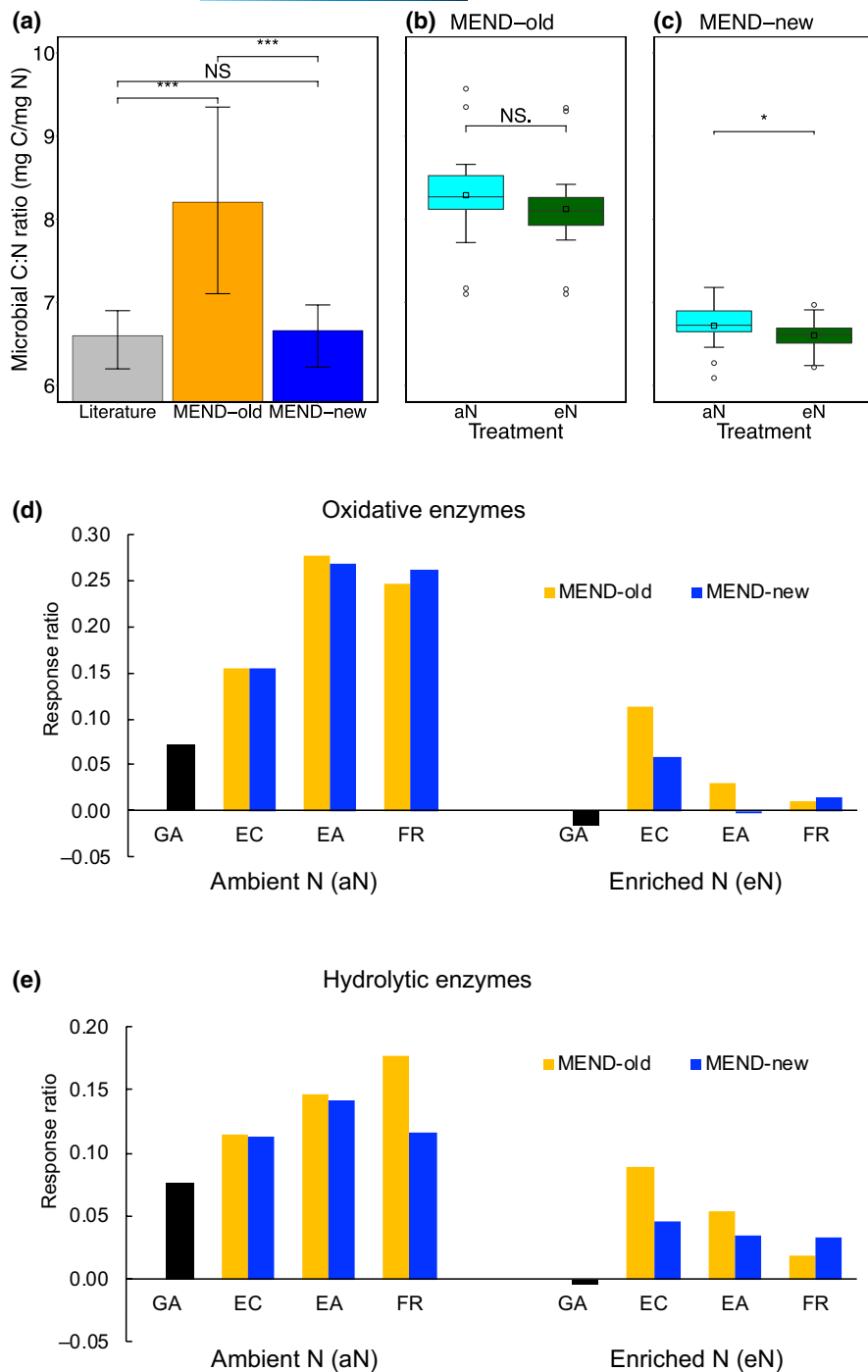
We further evaluated the similarity or dissimilarity between MEND-new simulated and observed response ratios of all eight enzymes associated with the C and N cycling by the Wilcoxon signed rank test (Conover, 1998). The simulated response ratios consist of  $EC_{sim}$  (Figure 6a),  $EA_{sim}$  (Figure 6b), or  $FR_{sim}$  (Figure 6c) for eight enzymes, whereas the observed response ratios include  $GA_{obs}$  for eight corresponding genes (Table 1 and Figure 6).

The simulated results of response ratios indicate that the  $eCO_2$  effects on the enzymes were more pronounced under aN than under eN, consistent with the responses in  $GA_{obs}$ , that is, 50% CI = 0.03–0.06 under aN versus –0.02 to –0.01 under eN (Figure 6). We also found that, among the three simulated variables ( $EC_{sim}$ ,  $EA_{sim}$ , and  $FR_{sim}$ ), only  $EA_{sim}$  responses were not significantly different from the responses of  $GA_{obs}$  under aN or eN according to the Wilcoxon signed-rank test (Figure 6b). Our results showed positive responses of  $EA_{sim}$  under aN for six out of eight enzymes, which concurred with the changes in  $GA_{obs}$ . However, the other two enzyme groups (NO and  $N_2O$  reductases) exhibited slightly negative response ratios (–0.019 and –0.003) when comparing  $eCO_2$ -aN to  $aCO_2$ -aN, which were not consistent with  $GA_{obs}$  responses (0.03 and 0.05). In addition, negative response ratios of  $EA_{sim}$  under eN were found for all enzymes except two groups (hydrolytic enzymes and  $NO_2^-$  reductases), which generally concurred with the changes in  $GA_{obs}$  under eN.

## 4 | DISCUSSION

### 4.1 | Ecosystem modeling with explicit enzymes as indicators of soil function

The MEND-new model developed here offers new capabilities to investigate microbial-enzyme mediated N fixation, nitrification, and denitrification, and plant-microbe competition for inorganic N, as well as inorganic N leaching and gaseous emissions processes, which adds additional features to the original MEND-old model (Gao et al., 2020; Wang et al., 2020). The oxidative and hydrolytic enzymes for depolymerizing SOM are actual molecules which are independently functional. However, the intracellular N enzymes (responsible for biological N fixation, nitrification, and denitrification) are not physical molecules and thus have little ability to function independently of a living cell (Fiencke & Bock, 2006; Schlesier et al., 2016; Song et al., 2017). Toward this end, we treat these inorganic N enzymes as simple bioindicators of likely activity of cellular-level microbial physiology. Explicit representing these intracellular N enzymes in the model is more “pseudo-mechanistic” rather than “truly mechanistic” (Hommel, 2020), but it provides a tractable way to capture complex biological dynamics of inorganic N cycling. Although enzyme-enabled representation of more detailed C-N transformation processes increases model complexity, it potentially offers important insights into microbial control over biogeochemical cycles and the interactions between multiple physical, chemical, and biological processes.

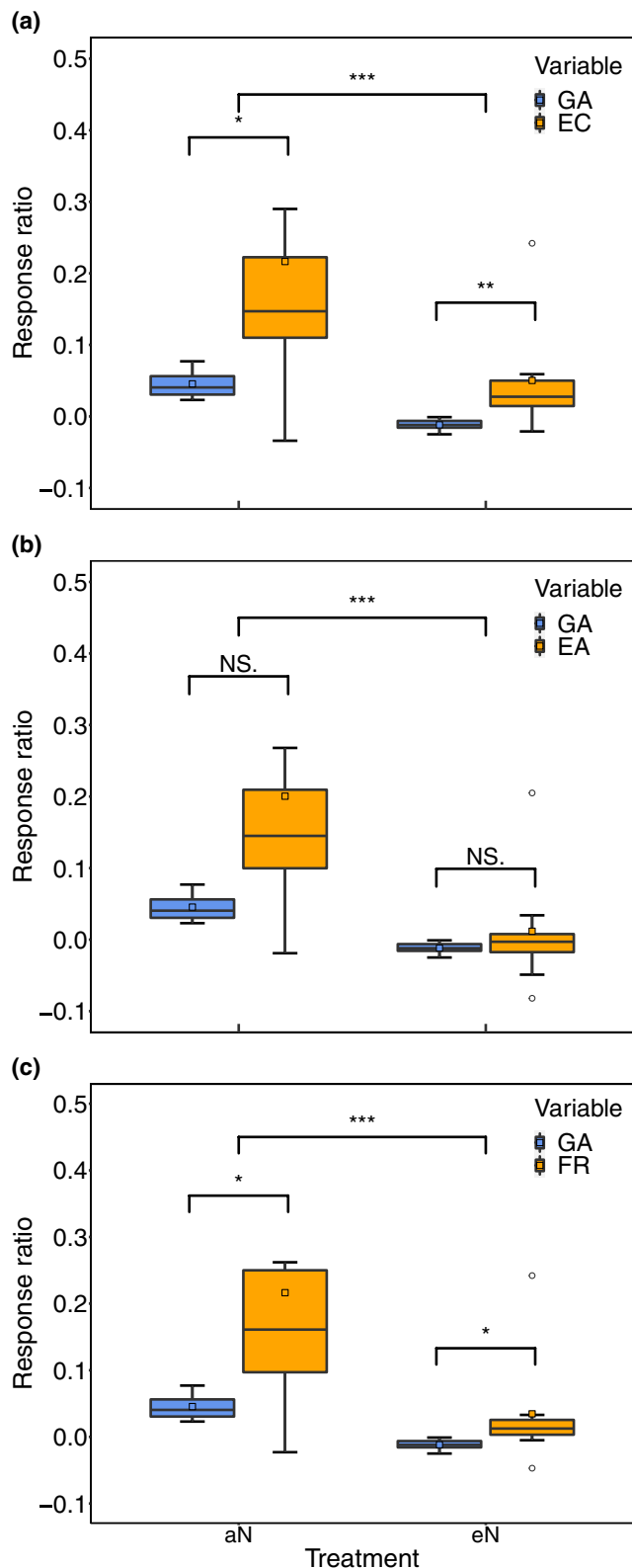


**FIGURE 5** Comparison of microbial C:N ratios and functional enzymes simulated by two models (MEND-old and MEND-new). (a) modeled versus literature-reported microbial C:N ratios (error bars denote the 95% confidence interval); (b) MEND-old modeled microbial C:N ratios under ambient N (aN) and enriched N (eN); (c) MEND-new modeled microbial C:N ratios under aN and eN; (d) elevated  $\text{CO}_2$  ( $\text{eCO}_2$ ) effect on oxidative enzymes; (e)  $\text{eCO}_2$  effect on hydrolytic enzymes. MEND-old and MEND-new denote the old version of MEND model as described in Gao et al. (2020) and the new MEND model in this study, respectively. The “Literature” data in (a) were from Xu et al. (2013). The  $\text{eCO}_2$  effects in the year of 2009 (d and e) are quantified by the response ratio (RR), which is defined as the logarithmic ratio of a variable under  $\text{eCO}_2$  to that under ambient  $\text{CO}_2$  ( $\text{aCO}_2$ ) as per ambient N (aN) or enriched N (eN) treatment. The RRs are calculated pertaining to observed gene abundances ( $\text{GA}_{\text{obs}}$ ), simulated enzyme concentrations ( $\text{EC}_{\text{sim}}$ ,  $\text{mg C cm}^{-3}$  soil), simulated enzyme activities ( $\text{EA}_{\text{sim}}$ ,  $\text{mg C cm}^{-3} \text{h}^{-1}$ ), and simulated first-order reaction rates ( $\text{FR}_{\text{sim}}$ ,  $\text{h}^{-1}$ ). The difference between paired data was tested by the Wilcoxon signed-rank test. “\*”, “\*\*”, and “\*\*\*” denote significant difference with  $p$ -value  $< .05$ ,  $p$ -value  $< .01$ , and  $p$ -value  $< .001$ , respectively. “NS.” means not significant

In contrast with enzyme-based models like MEND, the gene-centric approach was developed for examining ocean N cycling, where the gene abundances can be directly modeled to mediate chemical reactions (Reed et al., 2014). The gene-centric approach offers the advantage of direct comparison between modeled and measured gene abundances. However, currently, there is not enough information available for identifying appropriate biomarker genes for a specific metabolic pathway (Reed et al., 2014). In addition, for modeling a complex system with many processes, compared with the models characterized by enzyme groups, the number of genes may increase dramatically, resulting in difficulties and uncertainties

in estimating a vast number of parameters for these genes. In terms of ecosystem-level modeling that relies on bulk concentrations, it is currently more feasible to adopt the strategy with aggregated enzyme groups than the gene-centric approach.

We also proposed a competitive dynamic enzyme allocation scheme to assist the incorporation of multiple enzyme systems. Here, ‘dynamic’ means the allocation of each enzyme group varies with time, and ‘competitive’ implies that multiple enzyme systems compete with each other as per the relative saturation levels of the corresponding substrates. Enzyme allocation problems have been studied theoretically (Müller et al., 2014) or empirically (Sinsabaugh



**FIGURE 6** Elevated  $\text{CO}_2$  ( $\text{eCO}_2$ ) effects on functional genes/enzymes quantified by the response ratio (RR) in the year of 2009. (a) RRs of observed gene abundances ( $\text{GA}_{\text{obs}}$ ) versus simulated enzyme concentrations ( $\text{EC}_{\text{sim}}$ ,  $\text{mg C cm}^{-3}$  soil), (b) RRs of  $\text{GA}_{\text{obs}}$  versus simulated enzyme activities ( $\text{EA}_{\text{sim}}$ ,  $\text{mg C cm}^{-3} \text{h}^{-1}$ ), (c) RRs of  $\text{GA}_{\text{obs}}$  versus simulated first-order reaction rates ( $\text{FR}_{\text{sim}}$ ,  $\text{h}^{-1}$ ). The RR is defined as the logarithmic ratio of a variable under  $\text{eCO}_2$  to that under ambient  $\text{CO}_2$  ( $\text{aCO}_2$ ) as per ambient N (aN) or enriched N (eN) treatment. Each boxplot includes eight RR values from eight genes (enzymes): two groups (oxidative and hydrolytic) for the decomposition of soil organic matter, nitrogenases (nifH), ammonia oxidases (amoA), and four N-reductases (narG/napA, nirS/nirK, norB, and nosZ). The difference in RR between two variables was tested by the Wilcoxon signed-rank test. “\*”, “\*\*\*”, and “\*\*\*\*” denote significant difference with  $p$ -value  $< .05$ ,  $p$ -value  $< .01$ , and  $p$ -value  $< .001$ , respectively. “NS.” means not significant

realize that this allocation approach could not be directly evaluated as it is currently challenging to measure in situ production rates, particularly, of multiple enzyme systems. However, our model calibration and validation with a variety of inorganic N concentrations and fluxes indirectly demonstrated the applicability of this competitive dynamic enzyme allocation scheme, which was further supported by the model evaluation with measured gene abundance data.

#### 4.2 | Rigorous calibration and validation of microbial ecological models

Rigorous calibration and validation of microbial ecological models against observations is essential for assessing and refining models. However, finding appropriate datasets to validate microbial and enzymatic reactions in ecosystem models exhibits significant challenges. Treating inorganic N enzymes as indicators of soil function also allows the use of corresponding gene abundance data in ecosystem modeling, yet the relationship between enzymes and their coding genes is complicated (Bailey et al., 2018). Here, we used gene abundance data for model validation instead of calibration, because we attempted to explore the possible relationships between simulated ecosystem functioning (i.e., enzyme concentrations, enzyme activities, or reaction rates) and gene abundance. We showed that the changes in enzyme activities, rather than enzyme concentrations and the first-order reaction rates, are better explained by the responses in gene abundances. This may be due to the inclusion of more (eight in this study vs. two in Gao et al. (2020)) enzyme systems and relevant gene abundance data, which could introduce larger variation in the data resulting in differential modeling performance in terms of multiple variables. Therefore, we need more paired measurements of gene abundances and process rates under long-term field conditions in various ecosystems. DNA-based functional gene abundances have been thought to integrate longer-term (hours to days or longer) microbial potential in the physicochemical environment (Petersen et al., 2012; Rocca et al., 2015). Thus, we infer that DNA-based functional gene abundance is likely a better predictive

et al., 2002; Sinsabaugh & Moorhead, 1994) based primarily on stoichiometric information (Allison et al., 2011). These previous studies were generally focused on limited groups of enzymes (Averill, 2014; Müller et al., 2014), in contrast to the eight enzyme systems regulated by our competitive dynamic enzyme allocation scheme. We

variable for enzyme activity than for enzyme concentration or reaction rate. In the MEND model, enzyme activity contains the information of both active enzymes and their specific activity. To this end, enzyme activity represents the potential enzyme-catalyzed biogeochemical rates not limited by substrate availability (Ouyang et al., 2018; Petersen et al., 2012), whereas substrate availability is considered in the actual reaction rate (i.e.,  $FR_{sim}$  calculated by Equation 3). This interpretation supports our results on stronger relationship between  $GA_{obs}$  and  $EA_{sim}$  than between  $GA_{obs}$  and the other two variables ( $EC_{sim}$  and  $FR_{sim}$ ).

Very few studies have adopted gene abundance data in ecosystem or environmental modeling, where the model-data integration practices were often implemented for a short time period (e.g., 20 days) based on laboratory data (Gao et al., 2020; Li et al., 2017; Pagel et al., 2016; Song et al., 2017). Compared to these short-term laboratory-based modeling studies, it is likely more challenging to conduct gene-informed long-term (e.g., years to decades or longer) ecosystem modeling in the field, owing to complex spatiotemporal environmental conditions and large uncertainties in measurements, as demonstrated by the current study.

We also adopted the differential split-sample test to conduct a rigorous model calibration (for the baseline treatment  $aCO_2$ -aN) and validation (for the other three treatments under differential  $CO_2$  and N supply), which has been considered as the best possible approach for model parameterization (Refsgaard, 1997) and helped to demonstrate the predictive power of the calibrated model. During this process, we implemented advanced model-data integration by combining a wide spectrum of observations ranging from conventional measurements (e.g., soil respiration fluxes, concentrations of  $NH_4^+$  and  $NO_3^-$ ), to less frequently measured variables (e.g., all kinds of inorganic N fluxes and microbial biomass), and to rarely available gene abundance data of multiple enzyme systems that regulates SOM decomposition and inorganic N processes.

Simulation of some processes and properties were improved using our new modeling approach, while others were not. For example, we incorporated new data associated with N processes from the BioCON experiment into model calibration and validation, and compared to the BioCON results from the MEND-old model (Gao et al., 2020), the simulated  $NH_4^+$  and  $NO_3^-$  concentrations from this study were improved as indicated by much lower biases and generally lower AIC (except  $aCO_2$ -aN). By contrast, the model performance in soil respiration simulations was consistent between MEND-new ( $R^2 = 0.56$ – $0.61$ ) and MEND-old ( $R^2 = 0.53$ – $0.61$ ). Previous studies have demonstrated that the incorporation of more detailed biogeochemical processes might not necessarily improve modeling performance of soil respiration, as multi-objective model calibration aims to find a compromise between different objectives, such as various observed C-N pool sizes and process rates other than soil respiration (Bao et al., 2012; Davidson et al., 2012; Wang & Chen, 2012; Wang et al., 2019). Such model-data integration with multiple datasets on diverse system processes is crucial for examining the model's capability in representing a multitude of soil biogeochemical processes. In addition, model goals are not limited to improving

gross predictions but also gaining insights to underlying processes. Mechanistic understanding and representation of microbially mediated biogeochemical processes would help depict ecosystem responses to diverse perturbations more confidently.

Our direct validation of simulated microbial C:N ratios exemplifies the predictive power of the MEND-new model, given that microbial C:N ratios were not included in model calibration. The near congruence between observed and simulated microbial C:N ratios indicated the substantive improvement of the MEND-new model over the old version through mechanistic representation of N processes including dynamic N mineralization-immobilization and the competitive N uptake between plants and microbes. In addition, the MEND-new model predicted decreased microbial C:N ratios under enriched N supply (Xiao et al., 2018), owing to insignificant changes in microbial biomass C and significantly increased microbial biomass N.

It should be noted that the soil system studied in this study could be not well representative. The soil in the experimental site is a Typic Udipsamments that is minimally developed with no diagnostic horizons, and it could have little potential for stabilizing organic matter by mineral sorption or occlusion in aggregates (Schimel & Schaeffer, 2012; Six et al., 2002). Mineral interactions and spatial processes could play a small role in regulating the processing of plant detritus or SOM, or of N cycling processes in this soil. In addition, anaerobiosis and anerobic micro-sites are likely uncommon in the coarsely textured soils with high saturated hydraulic conductivity (O'Geen et al., 2017), which will certainly affect N (especially denitrification) dynamics, differently from other soils that are more developed with more texture structure (Kristensen et al., 2000). Nevertheless, this simple soil system is a perfect test-bed in many ways for experimental and modeling ideas. However, the parameterization of the model might not be readily applicable to other soils that have a fine texture and/or aggregate development, in which microbe-substrate-mineral interactions regulate the functioning of the biological components of the soil system. More likely, the model might just need different parameterization or perhaps more sophisticated treatment of organic-mineral interactions. Testing this would be a natural next phase in evaluating the model's applicability in diverse soils and ecosystems.

In summary, this study presents substantive methodological and ecological advances relative to previous studies, including our recent publication (Gao et al., 2020), in that (a) the MEND-new model now includes a more detailed representation of enzyme-catalyzed N transformation processes, with the addition of a competitive dynamic enzyme allocation scheme to tackle the synthesis of multiple enzyme systems; (b) the model was calibrated against a variety of observed N fluxes and validated by gene abundances for six N-associated processes, indicating that the changes in enzyme activities, rather than enzyme concentrations and reaction rates, were better explained by the measured gene abundances in responses to  $eCO_2$ ; and (c) the MEND-new model's predictions agreed well with the literature in terms of microbial C:N ratios and decreased

microbial C:N as a result of N addition, whereas the MEND-old model did not. Taken together, our results indicated that representing microbial-enzyme groups in ecosystem models is a potentially valuable step forward to develop robust predictive models that interpolate or extrapolate observed interactions among microbes and soil C-N cycling, likely bolstering confidence in the assessments and projections of carbon-climate feedbacks. Pertaining to model refinement, a comprehensive understanding of microbial communities and their roles in regulating specific C and nutrient processes is essential for successful incorporation of enzymes-based bioindicators in ecosystem modeling. The newly refined MEND model has the potential to provide a powerful avenue for understanding and testing hypotheses about microbially mediated soil biogeochemical processes under environmental changes.

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### CONFLICT OF INTEREST

The authors declare they have no conflict of interest.

### AUTHORS' CONTRIBUTIONS

All authors contributed intellectual input and assistance to this study and manuscript preparation. The original concept and modeling strategy were developed by Gangsheng Wang, Jizhong Zhou, Peter B Reich, and Sarah E Hobbie. Field experiments are maintained by Peter B Reich and Sarah E Hobbie. Model input data were compiled by Qun Gao, Yunfeng Yang, and Gangsheng Wang. The MEND modeling was developed and conducted by Gangsheng Wang. All data analysis and integration were guided by Gangsheng Wang and Jizhong Zhou. The paper was written by Gangsheng Wang and Jizhong Zhou, with help from Qun Gao, Yunfeng Yang, Peter B Reich, and Sarah E Hobbie.

### DATA AVAILABILITY STATEMENT

The model code and data that support the findings of this study are openly available in GitHub at <https://github.com/wanggangsheng/MEND.git>. The BioCON experimental data can be freely accessed at <https://www.cedarcreek.umn.edu/research/data>.

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