Development of a plant carbon-nitrogen interface coupling framework in a coupled biophysical-ecosystem-biogeochemical model (SSiB5/Triffid/DayCent-SOM v1.0): Its parameterization, implementation, and evaluation

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15 Abstract. Plant and microbial nitrogen (N) dynamics and N availability regulate the photosynthetic capacity and capture, allocation, and turnover of carbon (C) in terrestrial ecosystems. Studies have shown that a wide divergence in representations of N dynamics in land surface models leads to large <u>uncertaintyuncertainties</u> in the biogeochemical cycle of the terrestrial ecosystems and then in climate simulations as well as the projections of future trajectories. In this study, a plant C-N interface coupling framework is developed and implemented in a coupled biophysical-ecosystem-biogeochemical model

- 20 (SSiB5/TRIFFID/-DayCent-SOM v1.0). The main concept and structure of this plant C-N framework and its coupling strategy are presented. in this study. This framework takes more plant N-related metabolism processes into account. For instance, plant resistance and self adjustment is represented by a<u>A</u> dynamic C/N ratio for each plant functional type (PFT). is introduced to consider plant resistance and adaptation to N availability to better evaluate the plant response to N limitation. Furthermore, when available N is less than plant N demand, plant growth is restricted by a lower maximum carboxylation capacity of
- 25 Rubisco (Vmax) level,), reducing gross primary productivity (GPP). In addition, a module for plant respiration rates is introduced by adjusting the respiration with different rates at different plant components for<u>at</u> the same N concentration. Since insufficient N can potentially give rise to lags <u>onin</u> plant phenology, <u>phenologythe phenological</u> scheme is also adjusted with <u>a lag factor relatedin response</u> to N processes availability. All these considerations ensure a more comprehensive incorporation of N regulations to plant growth and C cycling. This new approach has been tested systematically to assess the effects of this
- 30 coupling framework and N limitation on the terrestrial carbon cycle. Long-<u>term</u> measurements from both flux tower sites with

different PFTs and global satellite-derived products are employed as references to assess these effects. The results show a general improvement with the new plant C-N coupling framework, with more consistent emergent properties, such as GPP and leaf area index (LAI), compared to <u>the</u> observations. The main improvements occur in tropical Africa and boreal regions, accompanied by a decrease <u>ofin</u> the bias in global GPP and LAI by 16.3% and 27.1%, respectively.

35 1 Introduction

Land surface processes substantially affect climate (Foley et al., 1998; Ma et al., 2013; Sellers et al., 1986; Xue et al., 2004, 2010, 2022) and are influenced by climate in turn (Bonan, 2008; Liu et al., 2019, 2020; Zhang et al., 2015), forming complex feedback loops to climate change (Friedlingstein et al., 2006; Gregory et al., 2009). To study these processes, the land surface components of Earth System Models (ESMs) have evolved from those that represent only physical processes (i.e., hydrology

40 and the energy cycle) to those that include the terrestrial carbon (C) cycle, vegetation dynamics, and nutrient processes (Cox, 2001; Dan et al., 2020; Foley et al., 1998; Oleson et al., 2013; Sellers et al., 1996; Sitch et al., 2003; Wang et al., 2010; Zhan et al., 2003).

Current land surface models have large uncertainties in predicting historical and recent C exchanges (Beer et al., 2010; Kou-Giesbrecht et al., 2023; Richardson et al., 2012) (Beer et al., 2010; Richardson et al., 2012; Zaehle et al., 2015). The

- 45 parameterization of some processes has been criticized for being oversimplified from an ecological point of view (Ali et al., 2015; Lawrence et al., 2019; Reich et al., 2006), The inclusion or exclusion of nutrient limitations on productivity is one of the critical factors., and the dynamic vegetation models tend to overestimate terrestrial C sequestration (Anav et al., 2015; Heikkinen et al., 2021; Murray-Tortarolo et al., 2013; Oliveira et al., 2021). The uncertainty/errors in predictions using land models have been attributed to many factors. The parameterization of some processes has been criticized for being
- 50 oversimplified from an ecological point of view (Ali et al., 2015; Lawrence et al., 2019; Reich et al., 2006). The inclusion or exclusion of nutrient limitations on productivity is one of the critical factors. Those The C-only models ignore significant nitrogen (N) impacts and therefore overestimate C sequestration by terrestrial ecosystems under climate change_(Peñuelas et al., 2013; Zaehle et al., 2015). Ecosystem N cycling processes are among the dominant drivers of terrestrial C-climate interactions through their impacts, mainly N limitation, on vegetation growth and productivity (Reich et al., 2006), especially
- in N-poor younger soils at high latitudes (LeBauer & Treseder, 2008; Vitousek and Howarth, 1991), and on microbial decomposition of organic matter (Hu et al., 2001)
 In the latest Coupled Model Intercomparison Project Phase 6 (CMIP6, Eyring et al., 2016. As such, the N cycle and its effect

on C uptake in the terrestrial biosphere have been incorporated into land surface models (LSMs) of ESMs (Davies-Barnard et al., 2020)(Davies-Barnard et al., 2020; Kou-Giesbrecht et al., 2023) with various representations of N processes (Ali et al., 2015; Asaadi et al., 2021; Best et al., 2011; Clark et al., 2011; Davies-Barnard et al., 2020; Ghimire et al., 2016; Goll et al., 2011; Clark et al., 2011; Davies-Barnard et al., 2020; Ghimire et al., 2016; Goll et al., 2011; Clark et al., 2011; Davies-Barnard et al., 2020; Ghimire et al., 2016; Goll et al., 2011; Clark et al., 2011; Davies-Barnard et al., 2020; Ghimire et al., 2016; Goll et al., 2011; Clark et al., 2011; Davies-Barnard et al., 2020; Ghimire et al., 2016; Goll et al., 2011; Clark et

2017; Krinner et al., 2005; Lawrence et al., 2019; Matson et al., 2002; Oleson et al., 2013; Smith et al., 2014; Thum et al., 2019; Wang et al., 2010; Wiltshire et al., 2020; Yu et al., 2020; Zhu et al., 2019).(Ali et al., 2015; Asaadi et al., 2021; Ghimire

et al., 2016; Goll et al., 2017; Lawrence et al., 2019; Oleson et al., 2013; Smith et al., 2014; Thum et al., 2019; Wiltshire et al., 2020).

- 65 <u>Adequate C-N coupling in plant N processes, however,</u>), although there were 112 different coupled models with various land surface models from 33 research teams, only about 10 models incorporated an N cycle module (Arora et al., 2020). The coupling of N processes is still an area of model development. Among these models including N processes, most of them pay more attention to microbial N dynamics in soil. The adequate C-N coupling in plant N processes has been indicated as an area that still needs intensive investigation (Thum et al., 2019; Ghimire et al., 2016; Goll et al., 2017; Yu et al., 2020; Zaehle et al.,
- 70 2015; Zhu et al., 2019). Some key plant N processes, such as N limitation on GPP, The fundamental aspects of N cycling for terrestrial biosphere models, such as N limitation of vegetation growth, strategies in which vegetation invests C to increase the N supply under N-limited conditions, and N limitation of decomposition, have been identified as important challenges for representing N cycling in terrestrial biosphere models (Meyerholt et al., 2020; Peng et al., 2020; Zaehle et al., 2015). Some key plant N processes, such as N limitation on gross primary productivity (GPP), the effect of biomass N content on autotrophic
- 75 respiration, plant N uptake, ecosystem N loss, and biological N fixation, have been introduced into LSMs with various complexity to present the N limitation effects in current land models. They include, for instance, using N to scale down the photosynthesis parameter V_{c,max} (Ghimire et al., 2016; Zaehle et al., 2015)complexities to determine the effects of N limitation in current land models. These methods include, for instance, using N to scale down the photosynthesis parameter V(c, max) (Ghimire et al., 2016; Zaehle et al., 2015) or potential GPP to reflect N availability (Gerber et al., 2010; Oleson et al., 2013;
- 80 Wang et al., 2010); defining a (Gerber et al., 2010; Oleson et al., 2013; Wang et al., 2010), defining the C cost of N uptake (Fisher et al., 2010)(Fisher et al., 2010a) and optimizing N allocation for leaf processes (Ali et al., 2015). The wide variety of assumptions and formulations of N cycling processes and C-N coupling reflects knowledge gaps and divergent theories, and further investigation is imperative (Kou-Giesbrecht et al., 2023). The coupling of N processes is still an area of model development. In the latest Coupled Model Intercomparison Project Phase 6 (CMIP6, Eyring et al., 2016), although there were
- 85 <u>112 different coupled models with various land surface models from 33 research teams, only about 10 models incorporated an N cycle module (Arora et al., 2020).(Ali et al., 2015). In many of these approaches, N limitation is represented as instantaneous down regulation of potential photosynthesis rates based on soil mineral N availability.</u>

This paper presents a recently developed process-based approach, which mainly focuses on the N limitation effects and plant respiration, and plant phenology. The dynamic plant C/N ratio is a key concept in

- 90 representing plant resistance, self-adjustment, and C/N interactions. Due to their relative immobility, plants often face significant challenges in obtaining an adequate supply of nutrients to meet the demands of basic cellular processes. A deficiency of any type of nutrient may result in decreased plant productivity and/or fertility. This paper presents a recently developed process-based plant C-N coupling framework with a consistent coupling strategy between biophysical and biogeochemical processes. The framework mainly focuses on the effects of N limitation on plant photosynthesis (Section)
- 95 <u>2.2.3</u>), plant respiration (Section 2.2.4), and plant phenology (Section 2.2.5) with a dynamic C/N ratio (CNR, section 2.2.2). The dynamic plant CNR is a more realistic representation than the fixed plant CNR in assessing the effect of N limitation on

plant C processes and interactions between plant C and N processes. (MeDowell et al., 2008; Morgan and Connolly, 2013; Stenberg and Muola, 2017).-Evidence has shown that plant C/N ratios have to change over the plant's lifecycle with nutrient availability (Chen and Chen, 2021; McGroddy et al., 2004; Meyer-Grünefeldt et al., 2015; Sardans et al., 2012; Smith, 1991;

- 100 Yang et al., 2021; Zhang et al., 2011) through plant self-adjustment. Plant cells C/N ratios are influenced by the accumulation of C polymers, such as carbohydrates and lipids, and are greater when cells are nutrient starved, or exposed to high light (Aber et al., 2003; MacDonald et al., 2002; Talmy et al., 2014). However, many land models specify fixed plant C/N ratios for each plant functional type (PFT) (e.g., Best et al., 2011; Clark et al., 2011; Krinner et al., 2005; Oleson et al., 2013; Wang et al., 2010). In this paper, we present a new plant C-N coupling framework with flexible C/N ratios (Section 2.2.2), in which N
- 105 regulates photosynthesis (Section 2.2.3), respiration (Section 2.2.4), and plant phenology (Section 2.2.5), as well as produces a consistent coupling between biophysical and biogeochemical processes. Allometric relations and empirical data sets are used to constrain the range of possible C/N ratios. This dynamic C/N ratio depends on the degree to which the N demands of different plant organs (e.g., leaf, root, and wood) are satisfied over the past several days. This plant C N framework takes some plant N metabolism processes into account and prevents unrealistic instantaneous down regulation of potential photosynthesis

110 rates.

We implement this plant C-N framework by coupling a soil organic matter and nutrient cycling model (DayCent-SOM; Del Grosso et al., 2000; Parton et al., 1998, 2010)with a biophysical/dynamic vegetation model (SSiB5/TRIFFID, the Simplified Simple Biosphere Model version 5/Top-down Representation of Interactive Foliage and Flora Including Dynamics Model, Cox, 2001; Harper et al., 2016; Liu et al., 2019; Xue et al., 1991; Zhan et al., 2003; Zhang et al., 2015). The SSiB and TRIFFID

- 115 have been extensively used for the land-atmosphere interaction studies (Harper et al., 2016; Xue et al., 2004, 2010, 2022, 2023). DayCent-SOM, which includes only the soil organic matter (SOM) cycling and trace gas subroutines from the DayCent ecosystem model (Parton et al., 1998, 2010)-, represents SOM transformations, below-ground N cycling, soil N limitation to microbial processes and plant growth, and nitrification/denitrification processes. In the coupled model, the potential N uptake depends on plant N demand from a biophysical and dynamic vegetation model, SSiB5/TRIFFID. The actual plant N uptake is
- 120 limited based on soil N availability, as predicted by DayCent-SOM (Del Grosso et al., 2000; Parton et al., 1998, 2010). Although this plant C-N coupling framework is developed based on SSiB5, TRIFFID, and DayCent-SOM, the methodology and approach in this study could be applied to other process-based land models with similar physical, biological, and ecological principles. The coupled model is verified at thirteen flux tower sites (Lund et al., 2012; Pastorello et al., 2020) with different plant functional types (PFTs) and is used to conduct several sets of global 2-D offline simulations from 1948 to 2007 to assess
- 125 the effects of the coupling process. Model predictions of global GPP and LAI have been evaluated against satellite-derived observational data(Jung et al., 2009; Sheffield et al., 2006; Zhu et al., 2013). The results demonstrate the relative importance of different plant N processes in this C-N framework.

The model used in this paper is presented in section 2.1. The development and implementation of this plant C-N framework isare presented in section 2.2. The model forcing and validation data used in this paper are presented in section 2.3. In section

130 3, the experimental design is described. In section 4, the measurements from the flux tower sites with different PFTs and the

global satellite-derived observations from <u>19481982</u>-2007 are used as references to assess the effect of the C-N coupling process on the long-term mean vegetation distribution and terrestrial C cycling using the offline SSiB5/TRIFFID/DayCent-SOM. Some issues and conclusions are presented in section 5.

2 Methods

135 2.1 Model description

2.1.1 SSiB4/TRIFFID model

The Simplified Simple Biosphere Model (SSiB, Xue et al., 1991; Sun and Xue, 2001; Zhan et al., 2003) is a biophysical model that simulates fluxes of surface radiation, momentum, sensible/<u>heat, and</u> latent heat, <u>as well as</u> runoff, soil moisture-and temperature, <u>surface temperatures</u>, and vegetation GPP_a based on energy and water balance and photosynthesis processes. The SSiB was coupled with a dynamic vegetation model, the Top-down Representation of Interactive Foliage and Flora Including Dynamics Model (TRIFFID), to calculate <u>NPP, LAI, net primary productivity (NPP), leaf area index (LAI),</u> canopy height, and PFT fractional coverage according to the C balance (Cox, 2001; Harper et al., 2016; Liu et al., 2019; Zhang et al., 2015). Moreover, the surface albedo and aerodynamic resistances are also updated based on the vegetation conditions.LAI, vegetation cover, vegetation height, and greenness. Previous work has improved the PFT competition strategy and plant physiology processes to make the SSiB4/TRIFFID suitable for seasonal, interannual, and decadal studies (Zhang et al., 2015). SSiB4/TRIFFID includes seven PFTs: (1) broadleaf evergreen trees (BET), (2) needleleaf evergreen trees (NET), (3) broadleaf

deciduous trees (BDT), (4) C3 grasses, (5) C4 plants, (6) shrubs, and (7) tundra. PFT coverage is determined by net-C availability<u>NPP</u>, competition between species, and disturbance, which includes mortality due to fires, pests, and windthrow. A detailed description and validation of SSiB4/TRIFFID can be found in Zhang et al. (2015), Liu et al. (2019), and Huang et al. (2020). In this study, The DayCent-SOM (see the next section) is introduced and coupled with the SSiB5/TRIFFID using the C-N interface coupling framework introduced in this study, which will be discussed in Sectionsection 2.2.

2.1.2 DayCent-SOM model

DayCent-SOM, a subset of DayCent that excludes plant growth, soil hydrology, and soil temperature subroutines, consists of soil mineral N pools (ammonium and nitrate) and six types of organic C and N pools consisting of two non-woody plant litter

155 pools (metabolic and structural), three coarse woody debris pools (from <u>the</u> death of large wood, fine branches, and coarse roots), and three kinetically defined organic matter pools, (active, slow, and passive); all types of organic pools except the passive pool have both <u>above-groundaboveground</u> and <u>below-groundbelowground</u> counterparts. (Table 1). Non-woody plant litter is partitioned into structural (lignin + cellulose) and metabolic (labile) litter based on the lignin: N ratio of <u>the</u> plant material (Parton et al., 1994)). The coarse woody debris pools decay in the same way that the structural pool decomposes,

160 with lignin and cellulose going to the slow soil organic matter pool and the labile fraction going to the active soil organic

matter pool. Each type of organic pool has its own intrinsic rate of decomposition, <u>which is modified by temperature and</u> moisture <u>effects</u> (Parton et al., 1994). Additionally, the decomposition rates of the structural material and coarse woody debris pools are functions of their respective lignin fractions. DayCent's litter decay model has been validated using extensive data from <u>the LIDET litter decay experiments from all over the world (Bonan et al., 2013).</u>

165 Table 1. The Nitrogen Pools in DayCent-SOM

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-	-	Aboveground	Belowground
<u>Mineral N</u> pool	-	-	Soil mineral N pools
	non-woody litter pools	<u>Surface structural N</u> Surface metabolic N	<u>Soil structural N</u> <u>Soil metabolic N</u>
<u>Organic N</u>	woody debris pool	Surface dead N	
<u>pool</u>	kinetically defined organic matter pools	<u>Surface active N</u> Surface slow organic N	<u>Soil active organic N</u> Soil slow organic N Soil passive organic N

2.2 Development of a plant Carbon-Nitrogen carbon-nitrogen (C--N) Interfaceinterface coupling framework

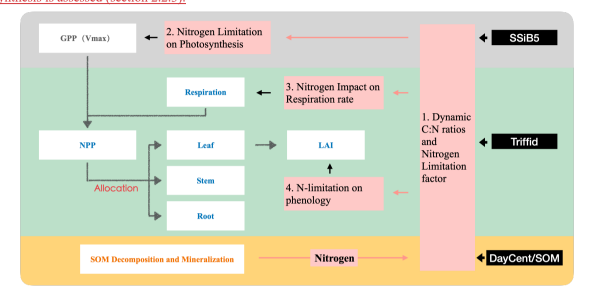
175 2.2.1 Conceptual considerations and coupling strategy

To represent C/ and N interactions, we have developed develop a plant C-N interface framework to take into account both couple biophysical and biochemical C/N-biogeochemical processes in plant life activities. the terrestrial C and N cycles. In this study, we applied the coupling framework to SSiB5/TRIFFID/-DayCent-SOM. However, this approach could be applied to any other process-based models with similar physical/biological principles. The conceptual considerations in developing this framework 180 are presented in this section. For a process-based model, introducing a consistent coupling philosophy between biophysical and biogeochemical processes is necessary. The surface water, radiation, carbon fluxes, and plant litter are calculated by SSiB5/TRIFFID. The soil N dynamics model (DayCent-SOM) is directly driven by soil temperature, soil moisture as well as, net radiation and plant C/ and N litter inputs into soil. Because the surface water, radiation, and carbon fluxes and plant litter are calculated by SSiB5, we force DayCent SOM withsoil organic pool, which are provided by the SSiB5-produced soil 185 temperature, soil moisture, and SSiB5/TRIFFID produced plant litter./TRIFFID. DayCent-SOM then computes daily changes ofin all organic matter and mineral soil pools, estimates losses of N from nitrate leaching and N₂O, NO_x, and N₂ emissions, predicts the amount of inorganic N available to plants, and updates inorganic N pools after accounting for plant N uptake by SSiB5/TRIFFID. Following plant N uptake from DayCent-SOM, our plant C-N interface framework describes Nthe effects on plant physiology from of N on photosynthesis, plant autotrophic respiration, and plant phenology plus(Fig. 1). All these 190 effects are associated with a dynamic C/N ratio (Fig. 1). Following such CNR. In the original land surface model development philosophy, our framework not only considers N limitation on the general plant (SSiB4/TRIFFID), with assumed unlimited N

availability and fixed CNR based on PFT, the assimilated C determined the N contents of leaves, stems, and roots, which

influenced photosynthesis process but also emphasizes the N limitation effect during the growth season, which can represent the physiological processes of C-N cycling and help us obtain, autotrophic respiration, NPP, and LAI. However, more information to understandevidence indicates that the attribution of N processes on CNR is not fixed in plant life, which will be further discussed in section 2.2.2. With the C cycle. dynamic CNR, the effect of N limitation on Rubisco capacity and photosynthesis is assessed (section 2.2.3).

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- 200 Figure 1. Schematic diagram of plant biogeochemistry and nitrogen impacts in SSiB5/TRIFFID/DayCent-SOM. Notes: (1) Different background <u>color representscolors represent</u> three different modules: SSiB, TRIFFID, and DayCent/SOM; (2) White boxes indicate the main processes <u>involved</u> in C-N coupling in different modules; (3) Vermeil boxes indicate how nitrogen influences plant biogeochemistry through the C-N framework.
- 205 Moreover, nitrogen is not the only dominant regulator of photosynthesis and vegetation dynamics. Reich et al., (2008) demonstrated strong relationships between respiration and N limitation based on observational data from various species. In the common N concentration range, respiration rates are consistently lower on average in leaves than in stems or roots. Therefore, this framework introduces two parameters for stems and roots based on PFT and available N, respectively, to adjust the respiration rate (section 2.2.4). Furthermore, N also affects plant phenology and can be remobilized to supply spring bud
- 210 break or vegetative shoot extension (Cox, 2001; Kolb and Evans, 2002; Marmann et al., 1997; Millard, 1994; Neilsen et al., 1997). The framework includes the impact of N on plant phenology by introducing an N limitation parameter, which will be discussed in section 2.2.5. With consideration of the effect on phenology, the N limitation effect during the growth season is emphasized. All these considerations in the framework should help to understand the effects of N processes to the C cycle more comprehensively

215 2.2.2 Dynamic C/N ratio based on plant growth and soil nitrogen storage

Plants often face significant challenges in obtaining an adequate supply of nutrients to meet the demands of basic cellular processes. Nutrient deficiency may result in decreased plant productivity and/or plant fertility(McDowell et al., 2008; Morgan and Connolly, 2013; Stenberg and Muola, 2017). Evidence has shown that plant CNR can change with nutrient availability (Chen and Chen, 2021; McGroddy et al., 2004; Meyer-Grünefeldt et al., 2015; Sardans et al., 2012; Smith, 1991). Plant cell

- 220 <u>CNRs are influenced by the accumulation of C polymers, such as carbohydrates, and are greater when cells are nutrient starved or exposed to high levels of photosynthetically active radiation (PAR) (Aber et al., 2003; MacDonald et al., 2002; Talmy et <u>al., 2014</u>). In the original land surface model (SSiB4/TRIFFID), with assumed unlimited N availability and fixed C/N ratios based on PFT, the assimilated C determined the N contents of leaf, stem, and root, which influenced autotrophic respiration, followed by GPP, LAI, and NPP. However, more evidence indicates the C/N ratio is not fixed in plant life. The studies of</u>
- 225 Ecological Stoichiometry The studies of ecological stoichiometry (Sterner and Elser, 2002), which investigates how the availability of multiple elements, including carbon, nitrogen, and phosphorus, constrainconstrains ecological interactions, revealshave revealed that plants can adjust their resource requirements. Changes inrespond and adapt to lower N resource availability will result in changes to plant C allocation and partitioning. Studies show that plants resorb only about 50% of leaf N on average (Aerts, 1996) to conserve nutrients (Clarkson and Hanson, 1980) and to increase nutrient use efficiency (Herbert
- 230 & Fownes, 1999; Vitousek, 1982). These processes cause a major internal nutrient flux and changes of C/N ratios(Herbert and Fownes, 1999; Vitousek, 1982). These processes cause changes in the CNR to reduce the impact of N limitation (Talhelm et al., 2011; Vicca et al., 2012). In addition, plant responses, such as plant resistance and self adjustment, will be limited under fixed C/N ratios, which affect plant productivity and litter N content, thus driving changes in the underground biogeochemistry and ultimately C and N uptake and storage (Drewniak and Gonzalez-Meler, 2017). Some studies show that the increase in
- 235 foliar N under increased soil N availability would improve plant responses because it allows adaptations in the stoichiometry of C and N (Bai et al., 2021; Chang et al., 2022; Ding et al., 2022; Kaiser et al., 2014; Lin et al., 2022; Zhang et al., 2022). The main impact will be to decrease C/N ratio in leaves when the available N increases, driving increases in productivity and changes in soil and litter N content (Drewniak and Gonzalez Meler, 2017). A dynamic C/N ratio is employed in our framework to obtain N states more realistically and properly represent the effect of N processes (See section 2.2.2 for more details).
- 240 A commonly used parameterization of photosynthetic C assimilation by the terrestrial biosphere in ESMsFor the response of vegetation to N limitation, i.e., the strategies in which vegetation invests C to increase N supply under N-limited conditions, some models represent flexible C/N stoichiometry, while others represent time-invariant C/N stoichiometry (Kou-Giesbrecht et al., 2023). Importantly, flexible vs. time-invariant C/N stoichiometry determines terrestrial C storage per unit N, followed by plant C allocation and partitioning. Plant responses are limited under a fixed CNR, which affects plant productivity and
- 245 <u>litter N content, thus affecting underground biogeochemistry and ultimately C and N uptake and storage. Comparing field measurements, it was found that allowing adaptations in the stoichiometry of C and N helped the land model improve the terrestrial surface C and N cycle simulation(Drewniak and Gonzalez-Meler, 2017; Medlyn et al., 2015).</u>

In this study, dynamic CNRs are introduced into SSiB5/TRIFFID. This dynamic CNR can enable vegetation to increase N uptake under N-limited conditions, reduce N limitation, and sustain terrestrial C sequestration. Plant resistance and adaptation

- 250 to N availability (N_{avail}) are represented by dynamic CNRs in SSiB5. The N availability (N_{avail}) for new growth limits the C assimilation rate through the CNRs, i.e., the model-simulated NPP should be no more than the N_{avail} is represented by the Farquhar, von Caemmerer, and Berry (FvCB) model of photosynthesis (Collatz et al., 1991; Farquhar et al., 1980). Plants require N as essential components of photosynthetic proteins involved in light capture, electron transport, and carboxylation (Evans, 1989). Nitrogen is an important constituent of the Rubisco enzyme and mitochondrial enzymes that regulate respiration
- 255 and adenosine triphosphate (ATP) generation (Makino and Osmond, 1991). One of the most important photosynthetic model parameters, the maximum carboxylation rate by the Rubisco enzyme (V_{c,max}) is a key parameter in the FvCB model (Farquhar et al., 1980) and has an extensive range across the models depending on the plant N content (Rogers, 2014). Since N is an important component of the Rubisco enzyme, leaf N content will affect V_{c,max} and thus GPP. The original FvCB model did not explicitly consider the N effect on photosynthesis. In a number of LSMs, an empirical relationship is applied to relate
- 260 $V_{e,max}$ to leaf N content N_{teaf} to generate the effect of N on photosynthesis, e.g., $V_{e,max} = i_{\psi} + s_{\psi} \times N_{teaf}$, where the intercept (i_{ψ}) and slope (s_{ψ}) are derived for each PFT based on observations (Kattge et al., 2009; Raddatz et al., 2007). Moreover, in some coupling approaches, only the relationship between the root N uptake and GPP/NPP is considered to represent the N limitation on C cycles (Ali et al., 2015; Fisher et al., 2010; Ghimire et al., 2016). However, because NPP is the difference between GPP and autotrophic respiration, adjusting NPP or GPP alone may cause the ratio between NPP and respiration to
- 265 deviate from reality. Using the process based N limitation factor produced from DayCent SOM to modify V_{c,max} (See section 2.2.3) is a more realistic way to produce the N effect on the photosynthesis process. Moreover, when calculating the N limit to Rubisco capacity, an aforementioned dynamic C/N ratio is introduced in our approach to influence the N limit effect on photosynthesis.

Nitrogen is not only a dominant regulator of vegetation dynamics, GPP, NPP, and terrestrial C cycles; Reich et al., (2008)

- 270 demonstrate strong relationships between respiration and N limitation based on observational data from various species. At any normal N concentration, respiration rates are consistently lower on average in leaves than in stems or roots. Therefore, we introduce two parameters for stems and roots, respectively, based on PFT to adjust the respiration rate in section 2.2.4. Nitrogen also affects plant phenology and can be remobilized to supply spring bud break or vegetative shoot extension (Kolb and Evans, 2002; Marmann et al., 1997; Millard, 1994; Neilsen et al., 1997). Nitrogen resorption is found during leaf
- 275 senescence and growth in evergreens (May and Killingbeck, 1992). Because plants need time to turnover, the plant N processes also have a lag effect on plant phenology (Thomas et al., 2015). Phenology in SSiB4/TRIFFID modulates LAI evolution, including leaf mortality, but it is not directly linked to N. Since different N states and supplements will lead to different lags on phenology, we add N impact on plant phenology by introducing a N limitation parameter, which will be discussed in section 2.2.5.

280 2.2.2 Dynamic C/N ratio based on plant growth and soil nitrogen storage

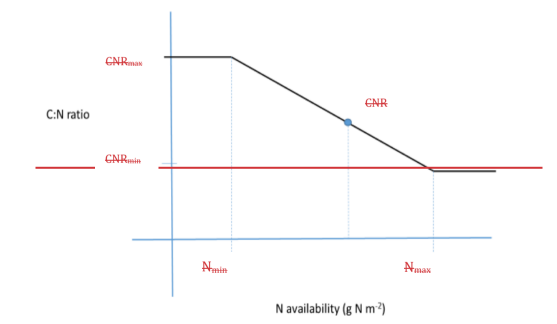
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Plant resistance and self-adjustment to N availability (N_{avail}) are represented by dynamic C/N ratios (CNRs) in SSiB5. The N availability for new growth limits the C assimilation rate through the CNRs, i.e., the model simulated NPP should be no more than N_{avail} × CNR of new plant material. In the original TRIFFID parameterization, the CNRs for different plant components (leaf, root, and woodstem) are fixed based on plant functional types (Cox, 2001); and changesthe change in CNR occurredthat occurs over the lifecycle of the plant, varied ecological process and varies with nutrient availability, and were was not considered in the original SSiB4/TRIFFID models. A linear relationship between the CNR and N_{avail} , based on the DayCent's

 $CNR = \begin{cases} CNR_{max}, & N_{avail} \le N_{min} \\ \frac{N_{avail} - N_{max}}{N_{min} - N_{max}} \times CNR_{min} + \frac{N_{avail} - N_{min}}{N_{max} - N_{min}} \times CNR_{max} & N_{min} < N_{avail} \le N_{max} \end{cases}$ (1) $avail \ge N_{max}$

parameterization, is introduced to the SSiB5/TRIFFID/DayCent-SOM for each PFT²s components, component (Fig. 2, Eq. 1).

where N_{avail} is the amount of soil mineral nitrogen that was available at the end of the previous day (g N m⁻²) calculated from 290 DayCent-SOM.



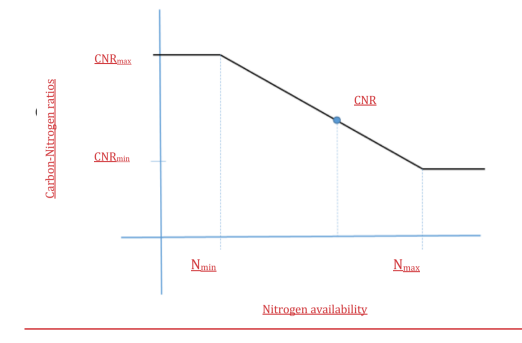


Figure 2. The relationship between soil nitrogen availability and plant carbon-nitrogen ratios.

The minimum and maximum amounts of nitrogen (N_{min}, N_{max}) necessary for the potential NPP_p (g C m⁻² day⁻¹), which is 295 first calculated from the SSiB4SSiB5/TRIFFID with unlimited N, are:

$$N_{min} = \frac{NPP_p}{CNR_{max}}$$
(2)

$$N_{max} = \frac{NPP_p}{CNR_{min}}$$
(3)

- where CNR_{min} and CNR_{max} are the minimum and the maximum C/N ratio CNRs, respectively, for each PFT's components 300 component from the DayCent (Table 1).2). Allometric relations and empirical datasets are used to constrain the range of possible CNRs. The CNR ranges CNRs of leaves, fine roots, and stems/wood are- were obtained from DayCent's user's manual and other published papers (Parton et al., 1993, 2007). Please noteNote that Eq. (2) and Eq. (3) are calculated based on the potential NPP; the CNR that is calculated based on Eqs. 1-3 ensures that when N_{avail} varies between N_{min} and $N_{max_{2}}$ the plant can adjust the CNR to support this potential NPP (as demonstrated in the schematic diagram in Figure 2). That said, the
- 305 N limitation will have no effect on the C assimilation as long as N_{avail} is largergreater than N_{min} . However, the N content of plant litter falling to the soil was determined by this dynamic C/N ratioCNR. Compared with the constant CNR, the range of possible plant carbon variation with dynamic CNR is smaller, reducing the impact of N limitation. As reviewed inat the introduction and previous beginning of this section, a number of recent studies have demonstrated that allowing adaptations in the stoichiometry of C and N would improve plant responses; for instance, an increase in available foliar N decreases the C/N 310 ratioCNR in leaves, driving an increase in productivity.

	Plant component	CNR _{min}	CNR _{max}
	leavesLeaves	20	50
Broadleaf deciduous	roots<u>Roots</u>	40	70
	woodStems	200	500
	leaves Leaves	20	40
Broadleaf Evergreen	roots <u>Roots</u>	40	70
	woodStems	150	300
	leaves Leaves	30	60
Needleleaf Evergreen	roots <u>Roots</u>	40	60
	woodStems	400	800
	leaves Leaves	20	40
C3 grass	roots <u>Roots</u>	40	50
	woodStems	40	80
	leaves Leaves	20	60
C4 grass plants	roots <u>Roots</u>	60	100
	woodStem	60	100
	leaves Leaves	20	40
shrub	roots <u>Roots</u>	40	70
	woodStems	200	400
	leavesLeaves	20	40
tundra -shrub	roots<u>Roots</u>	40	80
	woodStems	300	700

 Table 12. C:-N ranges of leaves, fine roots, and stems/wood for each plant function type (PFT).

Note: Data of The CNR_{min} and CNR_{max} data for each PFT²s components component are from DayCent's user²s manual and other publications (Parton et al., 1993, 2007)

The DayCent-SOM only provides the total available nitrogen (N_{avail}) for the plant within one grid box₅ (the soil is 3.2 m in depth), which consists of several PFTs. To apply equation 1, the nitrogen available for each PFT and its plant components in the grid box is calculated as

$$320 \quad N_{avail}(i) = N_{avail} * frac_i \tag{4}$$

$$N_{avail}(i, j) = N_{avail}(i) * \Delta C_j / \Sigma_j \Delta C_j$$

where $frac_i$ is the fraction of PFT i in one grid, and ΔCj is the fraction of carbon allocated to plant component j, which consists of <u>leaf, rootleaves, roots</u>, and <u>wood, stems</u> and is calculated in TRIFFID.

(5)

<u>Furthermore, the dynamic CNR in this framework depends on the degree to which the N demands of different plant components</u> (e.g., leaves, roots, and stems) have been satisfied over the past several days, and Eq. 1 prevents unrealistic instantaneous downregulation of potential photosynthesis rates. "Instantaneous downregulation" refers to the fact that photosynthesis rates are limited as soon as N (either in leaves or soil) is not sufficient (Reich et al., 2006; Ghimire et al., 2016), which has been applied in some N-limited models (Davies-Barnard et al., 2020). In our framework, by adjusting C/N ratios, the N limitation effect under certain conditions does not instantaneously respond to available N. The N limitation will produce effects only when available N passes some critical value (see further discussions in section 2.2.3).

2.2.3 Effect of nitrogen limitation on photosynthesis based on soil available nitrogen and the plant C-/N ratio

The widely used parameterization of photosynthetic C assimilation by the terrestrial biosphere in ESMs, including our model, is represented by the Farquhar, von Caemmerer, and Berry (FvCB) model of photosynthesis (Collatz et al., 1991; Farquhar et al., 1980). At high levels of PAR, the photosynthetic rate is limited by the amount of Rubisco in the leaf and its cycling rate.

- 335 <u>Nitrogen is an important constituent of the Rubisco enzyme and mitochondrial enzymes that regulate respiration and adenosine triphosphate (ATP) generation (Makino and Osmond, 1991). As one of the most important photosynthetic model parameters, the maximum carboxylation rate by the Rubisco enzyme (V_{c,max}) is a key parameter in the FvCB model (Farquhar et al., 1980) and has an extensive range across the models depending on the plant N content (Rogers, 2014). Therefore, leaf N content will affect V_{c,max} and thus GPP. However, the original FvCB model did not explicitly consider the effect of N on photosynthesis.</u>
- 340 While N limitation was introduced to terrestrial biosphere models, they differ in how N limitation in the plant C process is represented (Thomas et al., 2015; Fisher et al., 2010b). In a number of LSMs, an empirical relationship is applied to relate $V_{c,max}$ to leaf N content N_{leaf} to determine the effect of N on photosynthesis, e.g., $V_{c,max} = i_v + s_v \times N_{leaf}$, where the intercept (i_v) and slope (s_v) are derived for each PFT based on observations (Kattge et al., 2009; Raddatz et al., 2007). There are several different ways to represent N limitation effects in land models, including using N to scale down photosynthesis (Ghimire et
- 345 al., 2016; Goll et al., 2017; Thum et al., 2019; Yu et al., 2020; Zaehle et al., 2015; Zhu et al., 2019), scaling down potential GPP based on N availability (Gerber et al., 2010; Oleson et al., 2013; Wang et al., 2010), or defining an NPP cost of nitrogen uptake (Fisher et al., 2010). We choose the most physiological way by adjusting maximum Rubisco carboxylation rate (V_{c,max}) during the photosynthesis process, rather than adjusting NPP at the end of the photosynthesis process. -V_{c,max} regulates both C assimilation and autotrophic respiration; and the photosynthesis assimilation product, GPP, is proportional to V_{c,max}, which

We therefore introduce a downregulation of the canopy photosynthetic rate based on the available mineral N for new growth (N_{avail}) using a N-availability factor, f(N).

(6)

 $V_{c,max,Nlimit} = V_{c,max} * f(N)$

< NI

330

350

The f(N) is determined by nitrogen availability:

is proportional to nitrogen content of the Rubisco leaf reserves.

355
$$f(N) = \begin{cases} \frac{N_{avail}}{N_{min}} & N_{avail} \le N_{min} \\ 1 & otherwise \end{cases}$$

Because plants can adjust the relative allocations of C and N during N uptake via N remobilization and resorption to reduce the impact of N limitation, as discussed in the previous section for dynamic CNR, the N limit effect on photosynthesis only applies when nitrogen availability is lower than the minimum amounts-of nitrogen (N_{min}) necessary for the potential *NPP*. We take into account that plants have resistance and self adjustment through this approach to make the N limit effect neither

360 linearly nor instantaneously responsive to available N content. A linear relationship between f(N) and N_{avail} is valid only when N availability is not sufficient for the minimum N demand for new growth.
Some studies applied the same N limitation factor to NPP or GPP (Ali et al., 2015; Fisher et al., 2010; Ghimire et al., 2016).

<u>If NPP is adjusted, the same N limitation for photosynthesis is applied for plant respiration, which is not reasonable based on plant physiology (Högberg et al., 2017). Such approaches may distort the ratio of NPP and respiration. On the other hand, if
 only the GPP is adjusted for N limitation, then the N limitation for respiration is ignored.
</u>

- We chose the most physiological method by adjusting the maximum Rubisco carboxylation rate ($V_{c,max}$), which is proportional to the nitrogen content of the Rubisco leaf reserve) during photosynthesis rather than adjusting the NPP at the end of photosynthesis. $V_{c,max}$ regulates both C assimilation and autotrophic respiration, and the photosynthesis assimilation product, GPP, is proportional to $V_{c,max}$. Empirical evidence has shown that $V_{c,max}$ decreases with decreasing leaf N (Walker et al.,
- 370 <u>2014</u>). We therefore introduce a downregulation of the canopy photosynthetic rate based on the available mineral N for new growth (N_{avail}) using the N availability factor f(N).

$$V_{c,max,Nlimit} = V_{c,max} * f(N)$$

<u>The f(N) is determined by nitrogen availability:</u>

$$f(N) = \begin{cases} \frac{N_{avail}}{N_{min}} & N_{avail} \le N_{min} \\ 1 & otherwise \end{cases}$$
(7)

(6)

- 375 Because plants can adjust the relative allocations of C and N during N uptake via N remobilization and resorption to reduce the impact of N limitation, as discussed in the previous section for dynamic CNR, the N limitation effect on photosynthesis only applies when nitrogen availability is lower than the minimum amount of nitrogen (N_{min}) necessary for the potential NPP. We take into account the fact that plant responses include resistance and adaptation through this approach along with the dynamic CNR to make the N-limiting effect neither linear nor instantaneously downregulate the available N content, as
- 380 <u>discussed in the last section. A linear relationship between f(N) and N_{avail} is valid only when N availability is not sufficient for the minimum N demand for new growth.</u>

In fact, the factor, f(N) can also be applied to NPP and GPP as shown in Equations 8a b.

$$\frac{NPP_{Nlimit} = NPP * f(N)}{GPP_{Nlimit} = GPP * f(N)}$$
(8a)
(8b)

385 If NPP is adjusted (Eq. 8a), this means the same N limitation for photosynthesis is applied for plant respiration, which is not reasonable based on plant physiology (Högberg et al., 2017). Such approach may distort the ratio of NPP and respiration. If

only GPP is adjusted for N-limitation, then the N limitation for respiration is ignored. Therefore, we choose an approach to adjust *V_{c.max}*, which is related to N during the photosynthesis process and affects both C uptake and autotrophic respiration.

2.2.4 Improvement <u>in the impact</u> of nitrogen impact on <u>Nitrogen affects plant</u> respiration rates based on field 390 observations

Based on a database (Reich et al., 2008) of 2510 measurements from 287 species, the relationships between the mass-based dark respiration rate and nitrogen concentration of leaves, stems and roots were assessed. The results indicate strong respiration–nitrogen scaling relationships for all observations and for data averaged by species. At usual N concentrations, respiration rates are consistently lower on average in leaves than in stems or roots. In the original SSiB4/TRIFFID, the total

395 maintenance respiration (R_{pm}) is given by Cox (2001):

$$R_{pm} = 0.012 R_{dc} \frac{N_l + N_s + N_r}{N_l} \tag{9}$$

where R_{dc} is canopy dark respiration and is linearly dependent on $V_{c,max}$. The introduced N limitation of $V_{c,max}$ in Section 2.2.3 also influences the N effect on maintenance respiration. N_t , N_s and N_r are the N contents of leaf, stem, and root, respectively, and the factor of 0.012 is from the unit conversion. Eq. (9) assumes the respiration rates in root and stem have the same

- 400 dependence on N content as leaf. However, studies (Reich et al., 2008) have shown that the respiration rates at any common N concentration were consistently lower in leaves than in stems or roots on average. Thus, we introduce two PFT specific parameters (*ResA_s*, *ResA_r*) from field observations (Wang et al., 2006; Yang et al., 1992) to adjust root and stem respirations. Their values are listed in Table 2 The introduced N limitation of V_{c,max} in section 2.2.3 also influences the effect of N on maintenance respiration. N_l, N_s and N_r are the N contents of the leaf, stem, and root, respectively, and the factor of 0.012 is
- 405 from the unit conversion. Eq. (9) assumes that the respiration rates in roots and stems have the same dependence on the N content as that in leaves.
 Based on the information derived from field measurements for different PFTs (Reich et al., 2008; Wang et l., 2006; Yang et al., 1992), we introduce two PFT-specific parameters (*ResA_s*, *ResA_R*) to adjust root and stem respiration. Their values are listed in Table 3.

410
$$R_{pm,Nlimit} = 0.012 R_{dc} \frac{N_l + ResA_S * N_s + ResA_R * N_r}{N_l}$$
 (10)

PFT		Broadleaf Evergreen	Needleleaf Evergreen	C3 grass	C4 grass plants	shrub	tundra shrub
ResA _s	1.36	1.36	1.44	1.0	1.0	1.25	1.25
$ResA_R$	1.72	1.72	1.95	1.3	1.3	1.40	1.40

Table 23. The values of $ResA_S$ and $ResA_R$ for each plant function type (PFT).

Since $ResA_s$ and $ResA_R$ are generally larger than 1, new R_{pm} is larger than the original one, and the increased respiration due to the nitrogen limitation will decrease the NPP.

415

2.2.5 Effect of N limitation on the LAI based on plant phenology

Studies (Aerts and Berendse, 1988; Thomas et al., 2015) show that leaf turnover and aboveground productivity are related to nutrient availability and that plant N processes can potentially give rise to lags on phenology. In TRIFFID, a leaf phenology parameter, *p*, (Cox, 2001) is introduced to represent the vegetation's phenological status, to calculate the leaf drop rate, and to

420 adjust the model-simulated maximum possible LAI, which is based on carbon balance, (*LAI_{balance}*), to actual LAI and produce realistic phenology.

Nutrient availability affects vegetation activity and thus plant phenology (May and Killingbeck, 1992; Millard, 1994; Neilsen et al., 1997; Piao et al., 2019; Thomas et al., 2015; Vitasse et al., 2021; Zhou et al., 2022). Studies have demonstrated that variations in nitrogen availability could change the spring and fall phenology, such as spring bud break or vegetative shoot

425 extension (Yang et al., 2016; Yin et al., 2017; Fu et al., 2019), as well as the length of the growing season (Wang and Tang, 2019; Zhou et al., 2022). Increased soil nitrogen availability could supplement nutrient deficiencies and thus stimulate plant growth under low temperatures in early fall (Luke McCormack et al., 2014; Delpierre et al., 2016; Yin et al., 2017) and delay the end of the growing season (Wingler et al., 2006).

In TRIFFID, the leaf mortality rate and a leaf phenology parameter, p, (Cox, 2001), are introduced to represent the vegetation's

430 phenological status (Eqs. 11 and 12) and to adjust the model-simulated seasonal maximum possible leaf area index (LAI_{balance}), which is based on surface carbon balance (Cox, 2001; Enquist et al, 1998), to determine the actual LAI and produce realistic phenology.

$$LAI = p \times LAI_{balance} \tag{11}$$

and

$$435 \quad \frac{dp}{dt} = \begin{cases} -\gamma_p & \gamma_{lm} > 2\gamma_0 \\ \gamma_p (1-p) & \gamma_{lm} \le 2\gamma_0 \end{cases}$$
(12)

where <u>the</u> leaf constant absolute drop rate γ_p = 20 yr⁻¹, the leaf mortality rate γ_{lm} is a function of temperature T (Cox, 2001), and the minimum leaf turnover rate γ₀ = 0.25 (Cox, 2001). This phenology parameter, p, This phenology in SSiB4/TRIFFID modulates LAI seasonal evolution, which considers leaf mortality and the temperature threshold for leaf drop, but it is not directly linked to N. The phenology parameter p indicates that "full leaf" is approached asymptotically during the growing season, and p is reduced at a constant absolute rate when the mortality rate is largergreater than a certain threshold value. Otherwise, p increases, but the rate of increase is reduceddecreases as the growing season evolves. To reflect progresses.

Since different N states and supplies affect phenology, as discussed above, this framework includes the impact of N on plant phenology by introducing N limitation in SSiB5/TRIFFID/DayCent-SOM, we to take into account the effects of N on <u>phenology. We</u> assume that p is limited by N availability, with the new nitrogen limited limitation $p_{N \ limit}$ determined by

(13)

445 $p_{N \ Limit} = f(N) \times p$

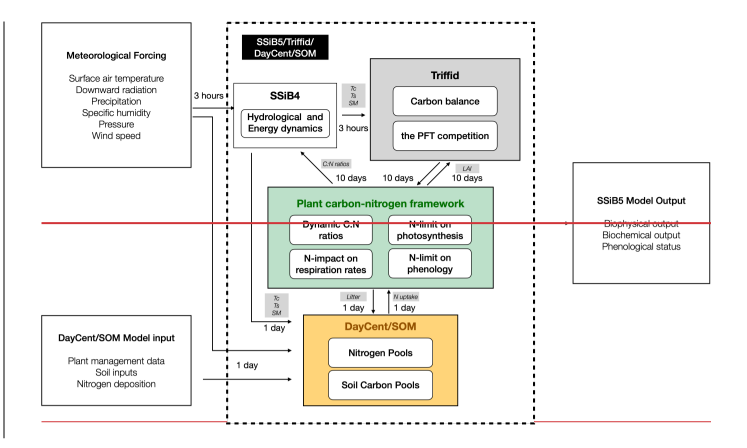
where f(N) is calculated as described in section 2.2.3.

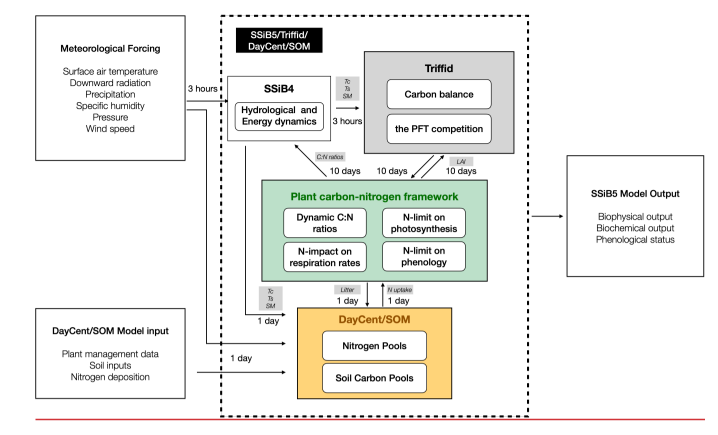
significant during the growth season.

2.2.6 The computational flow of SSiB5/TRIFFID/DayCent-SOM

In SSiB5/TRIFFID/DayCent-SOM, SSiB5 provides GPP, autotrophic respiration, and other physical variables, such as canopy and soil temperatures and soil moisture, every 3 hours for TRIFFID (Fig. 3). The TRIFFID accumulates the GPP from SSiB5 450 and produces biotic C, PFT fractional coverage, vegetation height, and LAI every ten days, which are used to update surface properties in SSiB5, such as albedo, surface roughness length, and aerodynamic/ and canopy resistances. The plant C-N framework uses the meteorological forcings (i.e., air temperature, humidity, wind, radiation, and precipitation) and physical variables (i.e., soil moisture and soil temperature) provided by SSiB5 every 3 hours and the biophysical properties (vegetation fraction and biotic C) provided by TRIFFID, which is are updated every ten days. The plant C-N interface framework calculates

455 the dynamic C/N ratiosCNR, N-limited photosynthesis, and N-impacted respiration rate every 3 hours. The Closs and potential N uptake are accumulated accumulate within one day in the C-N Interface Framework interface framework, and plant C and N litter fall litter fall are transferred to DayCent-SOM at the end of the day. DayCent-SOM calculates the amount of inorganic N available for plant N uptake (Navail) and the N losses from nitrate leaching and N-trace gas emissions each day. The TRIFFID updates the vegetation dynamics based on the C balance on day 10, including PFT competition. The updated vegetation dynamics are transferred to SSiB5 to calculate N-limited phenology to reflect N-the impact of N on the C cycle, which is 460





465 Figure 3. Flowchart of plant carbon-nitrogen interactions in SSiB5/TRIFFID/DayCent-SOM; <u>the main variables are listed</u> between <u>the two modules are listed</u>.

Notes: Tc: canopy temperature; Ts: land surface temperature; SM: soil moisture; GPP: gross primary productivity; Res: autotrophic respiration.

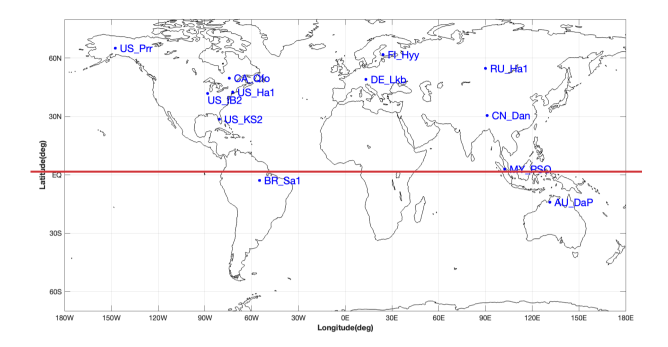
470 2.3 Model forcing and validation data

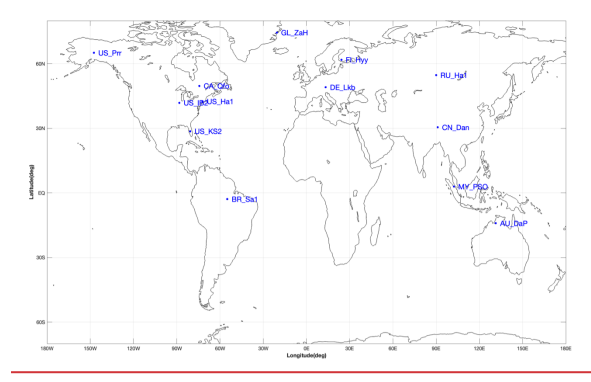
Long-_term measurements from flux tower sites with different PFTs and global satellite-derived products are employed as references to systematically assess the effects of this coupling framework and N limitation on the terrestrial carbon cycle. Flux tower <u>sitessite</u> data are presented in section 2.3.1. The global meteorological forcing and validation data are listed in sections 2.3.2 and 2.3.3 <u>separately, respectively</u>.

475 2.3.1 Ground measurement data

To validate the coupled model, twelvethirteen sites with representative biome types and climate zones were selected to evaluate the simulations of the seasonal patterns of GPP, sensible heat flux, and latent heat flux. The<u>All</u> driving data were a half-hourly dataset, including air temperature, specific humidity, wind velocity, air pressure, precipitation, and short-shortwave and long-

wavelongwave radiation data from the FLUXNET 2015 dataset (Pastorello et al., 2020). <u>The GL-ZaH data were obtained from</u>
 a tundra heath site (Lund et al., 2012). The geographical distribution of <u>the</u> selected FLUXNET 2015 sites is displayed in Figure 4, and the detailed site information is listed in Table 3. <u>4</u>.





485 Figure 4. Geographical distribution of selected FLUXNET 2015 sites. The information of these FLUXNET sites is listed in Table 14.

 Table 3.4. The
 FLUXNET sites, latitude (LAT), longitude (LONG), plant function type (PFT), and time frame (Time) used for the

 SSiB5/TRIFFID/-DayCent-SOM model validation.

Site_ID	Site name	LAT	LONG	PFT	Time
AU_DaP	Daly River Savanna	-14.06	131.32	C4 grassplants	2007-2013
BR-Sa1	Santarem-Km67-Primary Forest	-2.86	-54.96	Broadleaf Evergreen	2002-2011
CA_Qfo	Quebec - Eastern Boreal, Mature Black Spruce	49.69	-74.34	Needleleaf Evergreen	2003-2010
CN-Dan	Dangxiong	30.50	91.07	C3 grass	2004-2005
DE_Lkb	Lackenberg	49.10	13.30	Needleleaf Evergreen	2009-2013
FI_Hyy	Hyytiala	61.85	24.29	Needleleaf Evergreen	1996-2014
MY_PSO	Pasoh Forest Reserve	2.97	102.31	Broadleaf Evergreen	2003-2009
RU_Ha1	Hakasia steppe	54.73	90.00	C3 grass	2002-2004
US_Ha1	Harvard Forest EMS Tower (HFR1)	42.54	-72.17	Broadleaf deciduous	1991-2012

US_IB2	Fermi National Accelerator	Laboratory- B	Batavia (l	Prairie site)	41.84	-88.24	C3 grass	2004-2011
US-KS2	Kennedy Space Center (scru	ıb oak)			28.61	-80.67	Shrub	2003-2006
US_Prr	Poker Flat Research Range	Black Spruce	Forest		65.12	-147.49	Needleleaf Evergreen	2010-2014
<u>GL_ZaH</u>	Zackenberg Heath	<u>74.47</u> <u>-20</u>) <u>.55</u>	<u>Tundra</u>	<u>2000-20</u>	<u>14</u>		

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2.3.2 Meteorological forcing data

The Princeton global meteorological dataset for land surface modellingmodeling (Sheffield et al., 2006) was used to drive the SSiB4/TRIFFID global simulations from 1948 to 2007 at 1° x 1° spatial resolution and a 3-hour temporal interval. This dataset, includingwhich included surface air temperature, pressure, specific humidity, wind speed, downward short waveshortwave radiation flux, downward long wavelongwave radiation flux, and precipitation, was constructed by combining a suite of global observation-based datasets with the National Center for Environmental Prediction/National Center for Atmospheric Research reanalysis data.

2.3.3 Global remote-sensing data

- To assess the climatological status, variation, and trends of <u>the</u> simulated LAI, two widely used global LAI products were used as references in this study: the Global Inventory Modeling and Mapping Studies (GIMMS) LAI and the Global LAnd Surface Satellite (GLASS) LAI. GIMMS-LAI is based on the third generation of <u>Normalized Difference Vegetation Indexthe</u> <u>normalized difference vegetation index</u> (NDVI3g) from the GIMMS group and an <u>Artificial Neural Networkartificial neural</u> <u>network</u> model (Zhu et al., 2013). GIMMS-LAI provides a 1/12-degree resolution, 15-day composites, and spans<u>from</u> July 1981 to December 2011. GLASS-LAI is generated from Advanced Very High Resolution Radiometer (AVHRR) (from 1982 to 1999 with 0.05-degree resolution) and Moderate Resolution Imaging Spectroradiometer (MODIS, from 2000 to 2012 with
- 1 km resolution) reflectance data using general regression neural networks (Xiao et al., 2014). <u>The GIMMS</u> and GLASS <u>LAILAIs</u> and the meteorological forcing data for the overlap period <u>offrom</u> 1982 to 2007 were remapped to <u>a</u> 1-degree spatial resolution and a monthly temporal interval.
- The Model Tree Ensemble (MTE) GPP product (Jung et al., 2009) was used as a reference to evaluate <u>the</u> simulated GPP. 510 MTE is based on a machine learning technique in which the model is trained to predict the five C fluxes at FLUXNET sites driven by observed meteorological data, land cover data, and the remotely-__sensed fraction of absorbed <u>photosyntheticphotosynthetically</u> active radiation (Jung et al., 2009). The trained model was then applied at the grid scale driven by gridded forcing data. <u>The MTE-GPP</u> data were resampled to <u>a</u> 1-degree spatial <u>resolution</u> and a monthly temporal resolution. However, the MTE data do not include CO₂ fertilization. Liu et al. (2019) <u>discussdiscussed</u> this issue and 515 <u>indicateindicated</u> that the lack of CO₂ fertilization mainly affects the trend. Since this paper focuses on climatological

 $\frac{\text{mean}_{\text{mean}}}{\text{mean}}$ as well as differences between different experiments in which the CO₂ fertilization effect would be is largely cancelled, the missinglack of CO₂ fertilization in the FLUXNET-MTE is not a factor in interpreting our results.

3 Experimental designdesigns

520

To illustrate the reliability of the schemes which<u>that</u> represent different processes of plant N in our framework, we first evaluated the model's short-term performance using in-_situ measurements (section 3.2). Then, four sets of sensitivity experiments were designed to quantify the major effects of the plant N process and the relative contributions of different plant N processes on the terrestrial ecosystem carbon cycle (section 3.3).

3.1 Initial conditions for the dynamic vegetation model

The initial condition of the dynamic vegetation SSiB4/TRIFFID needs to be obtained from a long-term equilibrium simulation
(Zhang et al., 2015). There are different ways to initialize the surface conditions for quasi equilibrium simulations. Following previous SSiB4/TRIFFID studies (Huang et al., 2020; Liu et al., 2019; Zhang et al., 2015), we set up the initial conditionconditions for the run using the SSiB vegetation map and SSiB vegetation table, which are based on ground surveys and satellite-derived information (Dorman and Sellers, 1989; Sellers et al., 1986; Xue et al., 2004; Zhang et al., 2015) with 100% occupation at each grid point for the dominant PFT and zero occupation for other PFTs. We then ran the SSiB4/TRIFFID model with the climate forcing and the atmospheric CO₂ concentration at the 1948 level for 100 years to reach equilibrium conditions. The vegetation and soil conditions from the equilibrium results were used as the initial conditions for the subsequent model runs.

Determining the initial conditions for SSiB5/TRIFFIID/DayCent-SOM was carried out as described for SSiB4/TRIFFID with one additional step in order to initialize the global soil C and N levels. We saved 60 years of daily litter C/N inputs and soil temperature and moisture conditions from the SSiB4/TRIFFID that, which were based on historical meteorological forcings (1948-2007). An offline version of DayCent-SOM was run for 2000 years for each grid cell using these 60 years of data, repeated over and overrepeatedly, to determine the quasi-equilibrium soil C & and N levels; these soil C and N values were read in by SSiB5/TRIFFIID/DayCent-SOM at the start of the global simulation in 1948. This approach was applied for both measurement sites and global 2-D simulations.

540 3.2 Site-level validation

This paper focuses on the impact of N processes on the climatology of the global carbon cycle. Most current Dynamic Global Vegetation Modelsdynamic global vegetation models (DGVMs) are mainly focused on long-term (decadal to thousands of years or even longer) simulations at the global scale; the diurnal and seasonal variations are not athe subject forof their modellingmodeling. Moreover, adequate long-term in-situ measurements are also not available for comparison. However,

since the SSiB5/TRIFFID is a process-based model, we can evaluate the model's short-term performance using in-situ 545 measurements.

Twelve Thirteen sites with representative biome types and elimatesclimate zones (Table 34 and Fig. 4) were selected to evaluate the simulations of seasonal patterns of fluxes overacross these sites. The siteSite-level simulations were conducted by SSiB4/TRIFFID (a C-only model) and SSiB5/TRIFFID/DavCent-SOM separately to validate the model's performance. The

550 model results were compared against the observed daily data obtained by the flux tower, including the GPP, sensible heat flux, and latent heat flux.

3.3 Global 2-D2D offline control runruns and sensitivity runs

In this study, the SSiB4/TRIFFID and the SSiB5/TRIFFID/DayCent-SOM were applied to conduct a series of global 2-D2D offline runs (Table 45). All these runs employed the quasi-equilibrium simulation results as the initial condition, conditions 555 and were then were driven by the historical meteorological forcing from 1948 through 2007. The run using the SSiB4/TRIFFID is referred to as the control run (Exp. SSiB4 hereafter). Using the control simulation, we first evaluated the ability of the model to produce the climatology and variability of several biotic variables by comparing the results to multiple observation-based datasets. In addition to the control run, sixfour sets of sensitivity experiments were conducted to quantify the major effects of the N process and C-N interface coupling methodology on the C cycle. These sensitivity experiments were designed as follows:

560

(1) Nitrogen limitation on photosynthesis (Exp. NIPSN): The same meteorological forcing used for the control (Exp. SSiB4) drives the model, but dynamic $\frac{C/N \text{ ratios} CNR}{CNR}$ and N limitation on $V_{c.max}$ (Eq. 6) are introduced. The difference between Exp. SSiB4 and Exp. NIPSN indicates the effect of N limitation on photosynthesis.

(2) Nitrogen impact on Respiration rate (Exp. NlResp): The model was driven by the same meteorological forcing

565 used for Exp. SSiB4, but dynamic C/N ratioCNR and N impacts on autotrophic respiration (Eq. 10) are introduced. The difference between Exp. SSiB4 and Exp. NIResp indicates the effect of N impact on the respiration rate.

(3) Nitrogen limitation on Phenology (Exp. NIPhen): The model was driven by the same meteorological forcing used for Exp. SSiB4, but dynamic C/N ratioCNR and N impacts on phenology (Eq. 13) arewere introduced. The difference between Exp. SSiB4 and Exp. NlPhen indicates the effect of nitrogen limitation on phenology.

- 570 (4) SSiB5/TRIFFID/DayCent-SOM (Exp. SSiB5): The model was driven by the same meteorological forcing used for Exp. SSiB4, but all four C-N coupling processes in the framework, i.e., dynamic C/N-ratioCNR, N impacts on photosynthesis, autotrophic respiration, and phenology, are introduced. The difference between Exp. SSiB4 and Exp. SSiB5 indicates the effect of N dynamics, especially the sensitivity of C cycle variability and trendtrends to N process coupling. Furthermore, the difference between Exp. NIPSN and Exp. SSiB5 indicates the uncertainty (or possible errors) due to missing N effects
- on autotrophic respiration and phenology in the coupling framework. 575 Although the model runs were from 1948 to 2007, we only present the results from 1982-2007 to avoid spinning up for the SSiB5/TRIFFID/DayCent-SOM after SSiB4/TRIFFID and DayCent-SOM each reached their historical equilibrium conditions.

Since the results from Exps. SSiB5 and NIPSN showed statistically significant differences from Exp. SSiB4 over many parts of the world₇₂ in the following discussion, we will mainly focus on <u>the differences between</u> these two experiments' differences with and Exp. SSiB4.

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100-year equilibrium	Initial condition	Real-forcing 1948-2007	simulation
Fixed climatology forcing		Transient for	rcing
Control experiment		SSiB4:	Control experiment
		NIPSN:	Nitrogen limitation on photosynthesis_(Vmax), Eq.6
		NIResp:	Nitrogen impact on Respiration rate, Eq.10
		NIPhen:	Nitrogen limitation on Phenology, Eq. 13
		SSiB5: _	_including all four nitrogen processes

4. Results

585 To test this framework, measurements from flux tower sites with different PFTs and global satellite-derived products from 1982–2007 are employed as references. The results from site simulation and global 2-D simulations are presented in sections 4.1 and 4.2⁻⁻⁻⁻₁₂₂ respectively. As mentioned in section 2, the framework takes some plant N metabolism processes into account. To illustrate the relative contributions of different plant N processes <u>onto</u> the terrestrial ecosystem carbon cycle, four sets of sensitivity experiments were designed (Table 4<u>5</u>). The analyses are presented in section 4.2.

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4.1 Evaluations using the measurements from flux tower sites

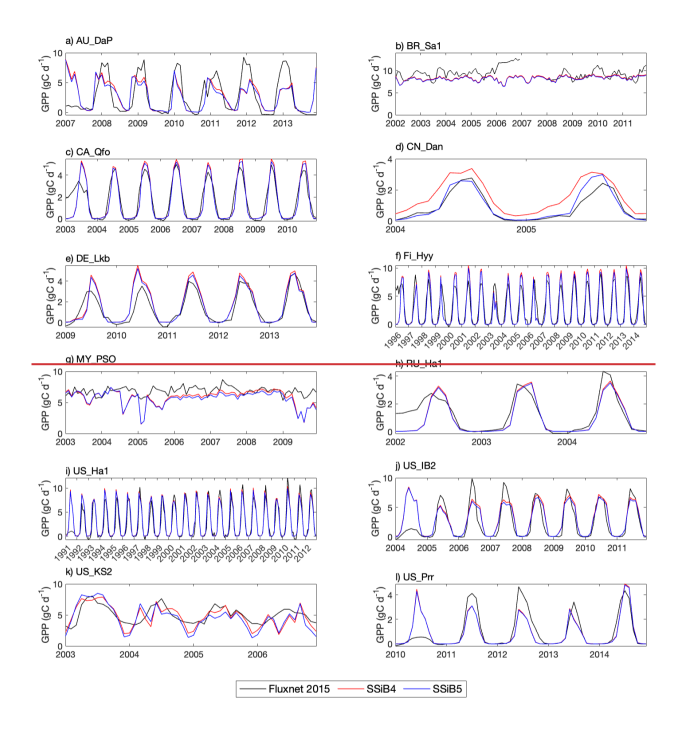
Land models with dynamic vegetation and nitrogen processes normally focus on the-long-term climate simulation with simulations at large spatial -scales. In this section, we validate the model performance for twelvethirteen sites with several years of simulation (Table 34) to ensure that, as a process-based model, the short-term SSiB5/TRIFFID in the short term simulation is still able to properly represent the surface processes at seasonal scales after introducing the introduction of DayCent-SOM through the interface coupling framework. This evaluation also provides a glance at the model's performance at several sites with various elimateclimates and PFTs (Table 34) with short-term data to gain preliminary confidence for further evaluation.

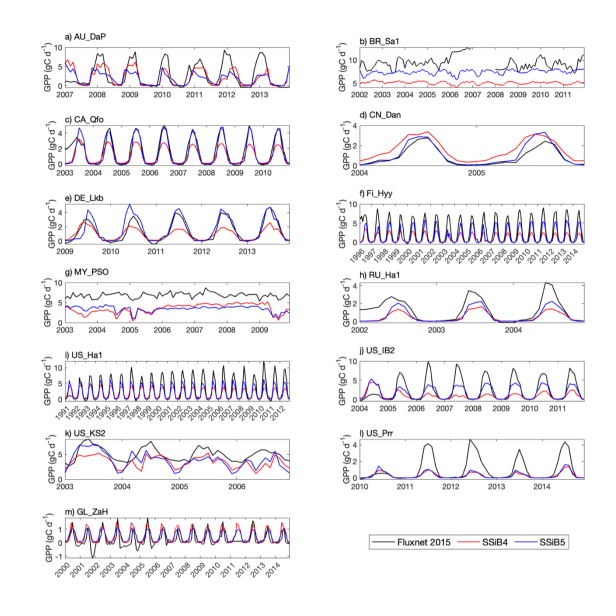
Figures 5, 6, and 7 show that both SSiB4 and SSiB5/TRIFFID/DayCent-SOM produce a reasonable seasonal cycle for GPP,

- 600 sensible heat, and latent heat fluxes, respectively, and that the results are close to observation.the observations. Table 56 summarizes the major results. We use bias, root-mean-squaredsquare error (RMSE), as well as and standard deviation to assess model performance against the in-situ site measurements. When we evaluateevaluated the 12 sites average of the 13 sites, the biases for GPP and sensible and latent heat fluxes are decreased by about approximately 7%, 1817%, and 2-%, respectively. The average RMSEs over the 1213 sites for these three variables are also decreased by about approximately 2%, consistent
- with the reduction in bias. Furthermore, the SSiB5/TRIFFID/DayCent-SOM producesproduced a closer standard deviation for GPP, sensible heat flux, and latent heat flux than did SSiB4/TRIFFID for the 1213-site averages. By and largeOverall, in these short-term simulations with specified initial vegetation conditions, both SSiB4 and SSiB5 produce reasonable GPP and surface heat fluxes compared with in-_situ measurements, but adding N processes (SSiB5) shows a slight improvement forslightly improved the 1213-site average. Although these improvements are rather marginal (except for the bias reduction for sensible
- 610 heat), the results nevertheless demonstrate that, with short-term simulation, <u>the improvement in the</u> model simulations² improvement is rather consistent.

With closer checking of the SSiB4 to SSiB5 results at each site, the results display various characteristics. For instance, <u>while</u> some sites mainly show the improvement in both bias and RMSE, while others show improvement only in one or the other. Moreover, while some sites show improvementshowed improvements in all three variables (GPP₇ and latent and sensible heat

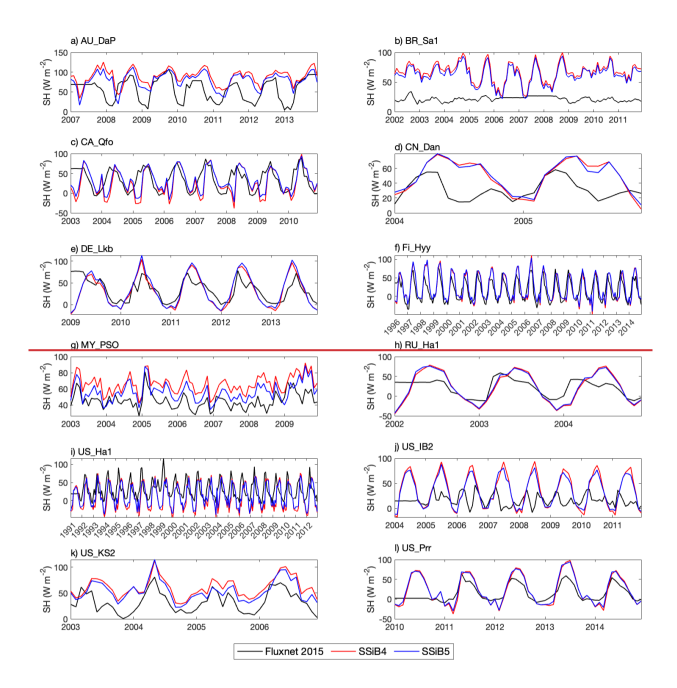
- 615 fluxes), others only show improvement forshowed improvements in one or two variables. It should be pointed outnoted that SSiB4 and SSiB5 are mainly used for global studies. For the validation of in-situ measurements, proper optimization of some site-specific soil and vegetation parameters is necessary (Xue et al., 1996, 1997). In this study, no model parameters were optimized during this validation exercise for a better fit between the simulated results and FLUXNET measurements. The discussions above leadled us to conduct long-term experiments at a global scale to comprehensively investigate the N process
- 620 effects of N processes and to help understand the mechanisms governing the global carbon cycle, which will be discussed in the following section.





625

Figure 5. Simulated seasonal variations of <u>in</u> GPP against observations at <u>twelve FLUXNET</u><u>thirteen</u> sites representing different SSiB5 PFTs. Note: <u>theThe</u> information about these FLUXNET sites is listed in Table <u>34</u>.



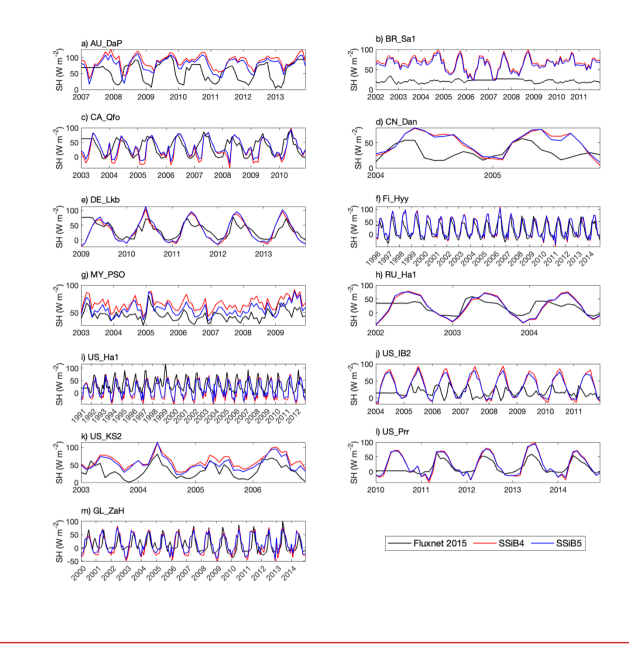
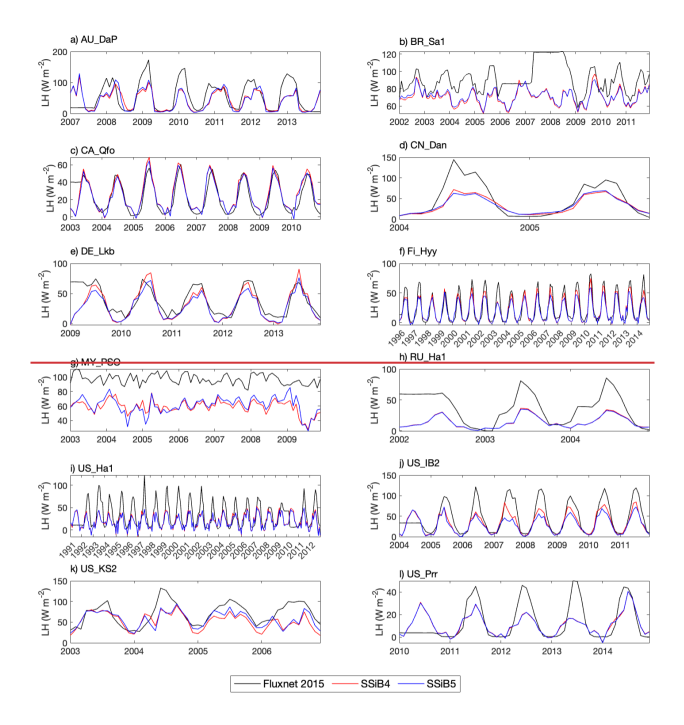


Figure 6. Same as Figure 5, but for sensible heat flux.



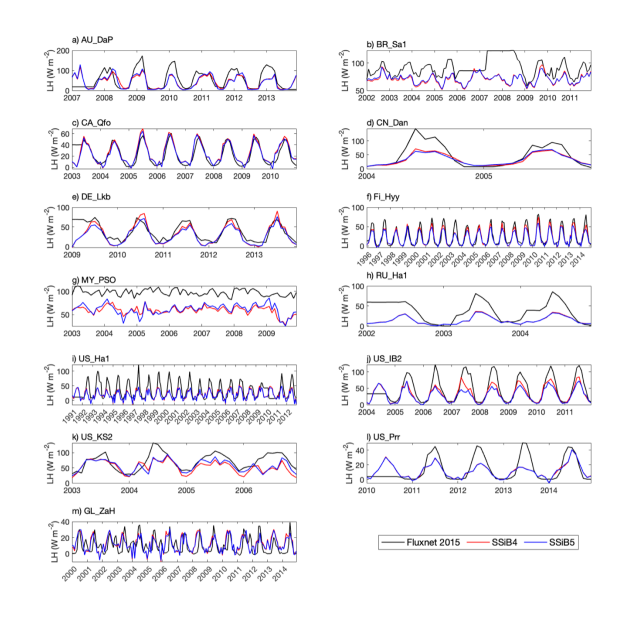


Figure 7. Same as Figure 5, but for the latent heat flux.

 635
 Table 56. The GPP, sensible heat flux, and latent heat flux intercomparisons comparisons of bias, standard deviation and RMSE between SSiB4 and SSiB5 over twelveat the thirteen sites.

 Sile_ID
 Bias
 Standard deviation
 RMSE

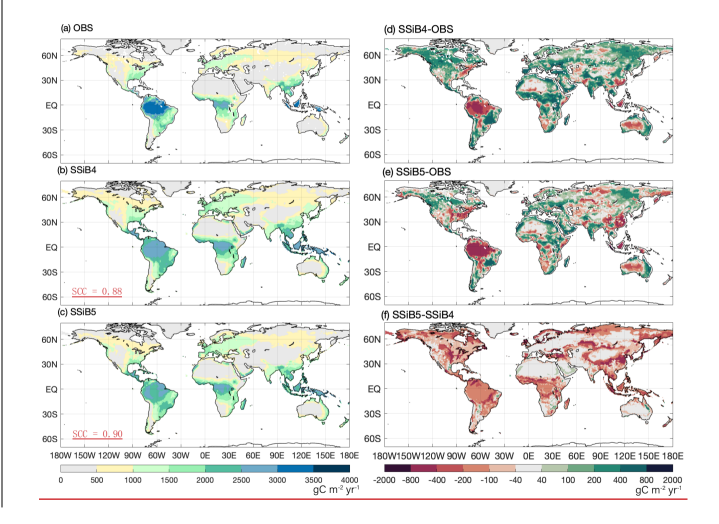
 Sile_ID
 Bias
 Standard deviation

 SSiB4
 SSiB4
 SSiB5
 SIB4
 SSiB5
 SSiB4
 SSiB5
 SSiB4
 SSiB5
 SSiB4
 SSiB5
 SSiB4
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	Site ID	Bias		Standard de	viation	RMSE		
	Site_ID -	SSiB4	SSiB5	Fluxnet	SSiB4	SSiB5	SSiB4	SSiB5
GPP	AU_DaP	0.05	-0.05	3.11	2.46	2.33	2.60	2.61
(g C d ⁻¹)	BR-Sa1	-1.07	-1.20	1.31	0.57	0.55	1.77	1.84
	CA_Qfo	-0.05	-0.11	1.71	1.99	1.92	0.78	0.75
	CN-Dan	0.70	0.08	0.92	1.08	1.03	0.80	0.33
	DE_Lkb	0.34	0.25	1.50	1.80	1.71	0.80	0.74
	FI_Hyy	-0.11	-0.22	2.93	3.47	3.32	1.51	1.44
	MY_PSO	-1.02	-1.20	0.65	1.28	1.21	1.63	1.72
	RU_Ha1	-0.24	-0.27	1.29	1.31	1.27	0.69	0.69
	US_Ha1	0.36	0.27	3.31	3.36	3.30	1.31	1.28
	US_IB2	0.56	0.42	2.91	2.70	2.57	1.80	1.79
	US-KS2	-0.28	-0.52	1.37	1.76	2.01	1.35	1.54
	US_Prr	-0.08	-0.10	1.43	1.30	1.28	0.86	0.86
	GI	<u>ZaH</u> <u>0</u> .	<u>28 0.25</u>	<u>0.50</u>	<u>0.53</u> <u>0.37</u>	<u>0.48</u>	<u>0.43</u>	
<mark>12<u>13</u>-site a</mark>	iverage	0.41 <u>40</u>	0. 38<u>37</u>	1. <mark>87<u>76</u></mark>	1. <mark>92</mark> 82	1. 88 <u>76</u>	1. 33 26	1. 30 23
Sensible	AU_DaP	32.47	23.13	28.26	19.64	21.05	36.24	36.32
Heat	BR-Sa1	45.29	40.94	4.04	16.32	15.98	25.61	25.07
Flux	CA_Qfo	-7.04	-2.34	27.77	33.18	29.37	9.54	9.20
(W m ⁻²)	CN-Dan	17.96	18.53	14.44	22.38	20.75	25.60	26.99
	DE_Lkb	-3.12	0.16	25.13	35.39	36.91	17.83	18.15
	FI_Hyy	5.53	7.20	28.17	33.57	33.63	8.99	10.91
	MY_PSO	20.49	10.86	10.03	11.30	11.98	39.22	37.99
	RU_Ha1	-0.14	0.84	21.71	39.19	38.02	29.42	29.67
	US_Ha1	-18.34	-15.80	24.40	33.71	29.42	24.33	24.66
	US_IB2	20.21	18.26	11.95	32.89	29.19	23.16	28.72
	US-KS2	27.74	20.81	21.01	19.17	20.14	27.31	24.73
	US_Prr	8.10	9.35	20.93	36.84	35.45	12.02	12.01
	<u>GL_Za</u>		<u>03</u> <u>27.08</u>	<u>36.04</u>	<u>31.48</u> <u>29.20</u>	<u>27.37</u>		
<mark>12<u>13</u>-site a</mark>	iverage	17.20<u>16.05</u>	14.02<u>13.33</u>	<u>19.82</u> 20.38	27.80 28.43	26.82 27.18	23. 27<u>73</u>	22.70 23.9
Latent	AU_DaP	-11.02	-10.83	45.72	30.03	33.93	36.24	36.32
Heat	BR-Sa1	-20.47	-19.82	16.15	9.44	8.47	25.61	25.07
Flux	CA_Qfo	2.21	0.96	18.06	18.63	17.56	9.54	9.20
(W m ⁻²)	CN-Dan	-12.63	-12.57	42.39	22.13	20.77	25.60	26.99
	DE_Lkb	-7.39	-10.00	22.81	24.57	20.79	17.83	18.15
	FI_Hyy	-3.06	-4.84	23.22	19.21	16.64	8.99	10.91
	MY_PSO	-38.18	-36.18	7.07	9.24	11.64	39.22	37.99
	RU_Ha1	-22.89	-23.10	25.68	10.43	10.08	29.42	29.67
	US_Ha1	-11.94	-13.14	27.06	15.53	14.71	24.33	24.66
	US_IB2	-12.90	-17.38	36.91	24.68	20.70	23.16	28.72
	US-KS2	-17.74	-13.41	27.63	20.28	19.65	27.31	24.73
	US_Prr	-1.90	-1.87	16.44	9.62	9.68	12.02	12.01
	<u>GL_Za</u>	<u>H 2.76 2.</u>	<u>26 10.17</u>	<u>9.67</u>	<u>9.17 11.15</u>	<u>10.48</u>		
1213-site a		13.93<u>12.80</u>	13.67 12.70	25.7624.56	17. <mark>82</mark> 19	17.95 16.45	<u>24.2722.68</u>	23.7022.3

4.2 Evaluation of GPP and LAI at global scale the Global Scale

The SSiB model is mainly used for global study, in this section, we evaluate the model's performance usingclimate studies. It
is important to adequately produce the observed global GPP and LAI-data. The model's performance in these aspects is evaluated in this section. The SSiB4/TRIFFID-simulated global PFT distribution has been extensively discussed in Zhang et al. (2015) and Liu et al. (2019). The simulation results are generally consistent with observation. The spatial distribution from the SSiB5/TRIFFID/DayCent-SOM did not show substantial difference and will not be discussed here. The simulated GPP averaged over 1982-2007 iswas compared to the FLUXNET-MTE GPP (Jung et al., 2011) to examine the impact of N processes and itstheir coupling with C and ecosystem processes. Both SSiB4/TRIFFID (Exp. SSiB4) and SSiB5/TRIFFID/DayCent-SOM (Exp. SSiB5) capture the distribution of global GPP (Fig. 8) and its latitudinal distribution (Fig. 9a).



650 Figure 8. The 1982-2007 average gross primary production comparison for (a) FLUXNET-MTE GPP (OBS), (b) SSiB4/TRIFFID (SSiB4), and (c) SSiB5/TRIFFID/DayCent/SOM (SSiB5) and the difference between (d) SSiB4-OBS, (e) SSiB5-OBS, and (f) SSiB5-SSiB4. Note: SCC indicates the spatial correlation coefficient between the model simulation and satellite-derived datasets (OBS).

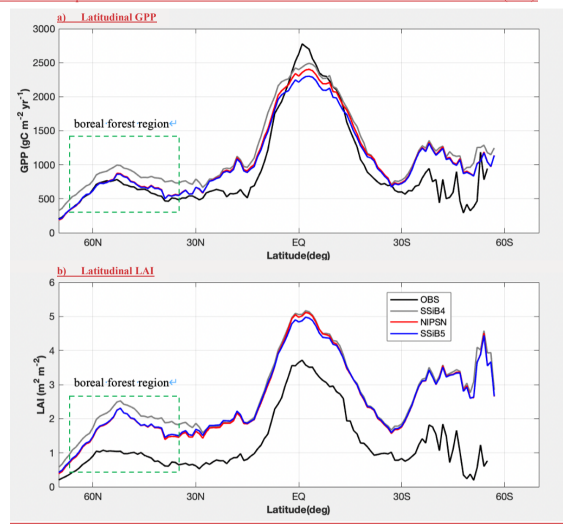


 Figure 9. Intercomparisons of latitudinal LAI and GPP among OBS, SSiB4 (control), NIPSN (N limitation on photosynthesis only), and
 SSiB5 (all N processes) over the period of 1982-2007. Note: Observed LAI is the GIMMS LAI.

35

Table 7. Regional and global GPP for (a) FLUXNET-MTE GPP (observation), (b) SSiB4 (control), (c) NIPSN (N limitation on photosynthesis only) and (d) SSiB5 (N limitation on photosynthesis, autotrophic respiration, and phenology).

					GPP M	ean (gC n	$n^{-2} yr^{-1}$)		
Regions	Sub-regions	MT	E	5	SSiB4 NIPSN		SSiB5		
		mean	bias	mean	bias	mean	bias	mean	bias
	West Africa	893		1147	254(28.5%)	963	70(7.9%)	915	22(2.5%)
Arid and Semi- Arid Areas	West NA	438		549	111(25.4%)	454	16(3.5%)	431	-7(-1.6%)
	SA Savanna	1665		1860	195(11.7%)	1763	98(5.9%)	1675	10(0.6%)
	East Africa	1228		1533	306(24.9%)	1427	199(16.2%)	1356	128(10.4%)
	East Asian semi-arid	1440		1470	30(2.1%)	1199	-241(-16.7%)	1139	-301(-20.9%)
NH High-Mid	NA High-Mid Latitude	552		814	262(47.6%)	700	149(27.0%)	665	114(20.6%)
Latitude Areas	Eurasian High-Mid	844		966	122(14.5%)	871	27(3.2%)	827	16(-2.0%)
	Amazon Basin	2993		2668	-326(-10.9%)	2631	-362(-12.1%)	2500	-494(-16.5%)
Equator	Southeast Asia	2778		2540	-238(-8.6%)	2419	-359(-12.9%)	2298	-480(-17.3%)
	Equator Africa	2522		2645	123(4.9%)	2611	89(3.5%)	2481	-42(-1.7%)
Subarctic Areas	NA Subarctic	234		364	130(55.7%)	240	6(2.4%)	228	-6(-2.7%)
and Tibet	Eurasian Subarctic	331		484	153(46.2%)	328	-3(-1.0%)	311	-20(-6.0%)
	Tibet	409		561	153(37.3%)	298	-111(-27.2%)	283	126(-30.8%)
Global		863		1082	220(25.4%)	991	129(14.9%)	942	79(9.1%)

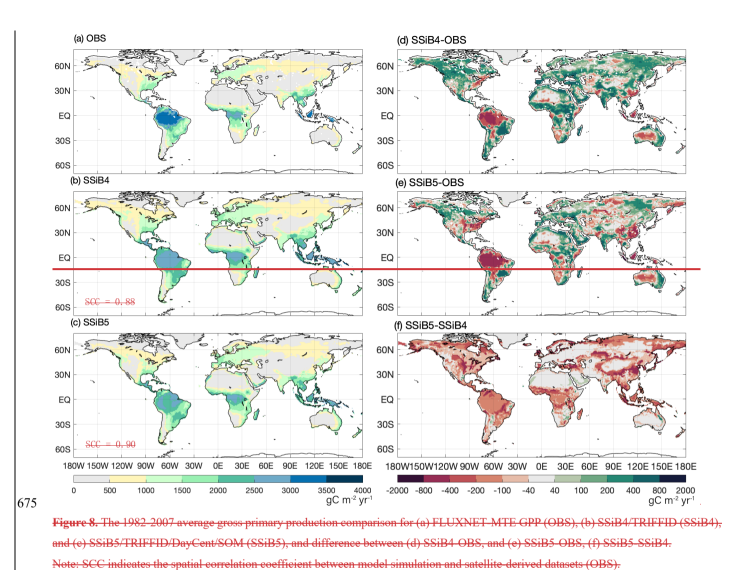
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Note: the numbers in parentheses are relative biases: (bias/MTE mean)

The highest GPP occurs in the tropical evergreen forestforests and generally decreases with the increase in latitudes in increasing latitude according to both the observations and the model simulation simulations (Figs. 8 and 9a). Exp. SSiB4-

665 simulated GPP has <u>a</u> positive bias over many parts of the world (Fig. 8d), including tropical Africa and the North American and eastern Siberian boreal regions, but <u>a</u> negative bias in some regions, mainly in the Amazon tropical forest. The simulated global GPP is 1082.36 g C m⁻² yr⁻¹ (Table 6),7), which is higher than the estimate,estimated value of 862.86 g C m⁻² yr⁻¹ in FLUXNET-MTE (Jung et al., 2011). After introducing the N limitation for the three processes, the SSiB5 reduced the positive bias in SSiB4 over many parts of the world (Figs. 8e, 8f, and 9a). Exp. SSiB5's global GPP prediction, 941.81 g C m⁻² yr⁻¹, is

670 closer to <u>the</u> observations <u>compared tothan</u> Exp. SSiB4, with a 16.3% reduction in the bias (Table 67). Furthermore, the temporal correlation coefficients between <u>the</u> observed and simulated monthly/annual mean GPPs are increased from 0.46/0.98 (Exp. SSiB4) to 0.50/0.99 (Exp. SSiB5), respectively (Fig. 10), showing improvement in <u>the</u> simulation of the seasonal cycle in SSiB5. The correlation for interannual variability in SSiB4 is already very high (0.98). SSiB5 shows no substantial improvement<u>continues the high correlation of SSiB4</u>.



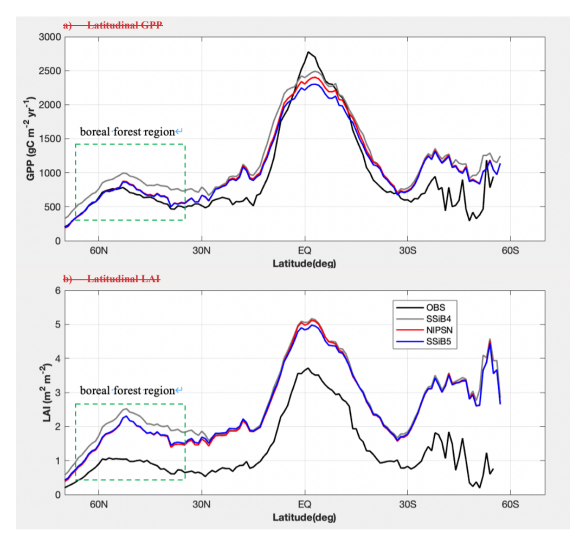


Figure 9. Intercomparisons of latitudinal LAI and GPP among OBS, SSiB4 (control), NIPSN (N limitation on photosynthesis only), and SSiB5 (all N processes) over the period 1982-2007.

685 photosynthesis only) and (d) SSiB5 (N limitation on photosynthesis, autotrophic respiration, and phenology).

	Sub-regions	GPP Mean (gC m ⁻² yr ⁻¹)								
Regions		MTE		5	SSiB4	ľ	NIPSN	SSiB5		
		mean bias		mean	bias	mean	bias	mean	bias	
Arid and Semi- Arid Areas	West Africa	893		1147	254(28.5%)	963	70(7.9%)	915	22(2.5%)	
	West NA	438		549	111(25.4%)	454	16(3.5%)	431	-7(-1.6%)	
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	East Africa	1228		1533	306(24.9%)	1427	199(16.2%)	1356	128(10.4%)	
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NH High-Mid	NA High-Mid Latitude	552		814	262(47.6%)	700	700 149(27.0%)		114(20.6%)	
Latitude Areas	Eurasian High-Mid	844		966	122(14.5%)	871	27(3.2%)	827	16(-2.0%)	
Equator	Amazon Basin	2993		2668	-326(-10.9%)	2631	-362(-12.1%)	2500	-494(-16.5%)	
	Southeast Asia	2778		2540	-238(-8.6%)	2419	-359(-12.9%)	2298	-480(-17.3%)	
	Equator Africa	2522		2645	123(4.9%)	2611	89(3.5%)	2481	-42(-1.7%)	
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	Eurasian Subarctic	331		484	153(46.2%)	328	-3(-1.0%)	311	-20(-6.0%)	
	Tibet	409		561	153(37.3%)	298	-111(-27.2%)	283	126(-30.8%	
Global		863		1082	220(25.4%)	991	129(14.9%)	942	79(9.1%)	

Note: the numbers in parentheses are relative biases: (bias/MTE mean)

690 The improvement, however, is not homogeneous over<u>across</u> the globe but displays apparent regional characteristics. The GPP biases in tropical Africa, <u>the North American boreal region</u>, <u>the South American savanna</u>, and <u>the central U.S. showshowed</u> substantial <u>reductionreductions</u> (Fig. 8f), which <u>helphelped</u> improve the spatial distribution of SSiB5. The global spatial correlation coefficient increases from 0.88 to 0.90 (Fig. 8). <u>MeanwhileMoreover</u>, the GPP simulations <u>did not improve</u> in some regions, such as in <u>the temperate East Asian mixed forest-grassland regions and in some areas <u>inof</u> Siberia (Fig. 8), <u>did not improve.8)</u>. In particular, the negative GPP bias in the Amazon <u>is-increased</u> (Fig. 8f). This phenomenon <u>has</u>-also appeared in</u>

the offline test inat the Amazon site (the BR-Sa1 Sitesite, Table 54). Du et al. (2020) indicate indicated that phosphorus (P) has morea greater effect in tropical areas. This paper mainly focuses on model development and preliminary global evaluation. More regional evaluation is necessary for further investigation, especially for the regions where the N limitation is not dominant.

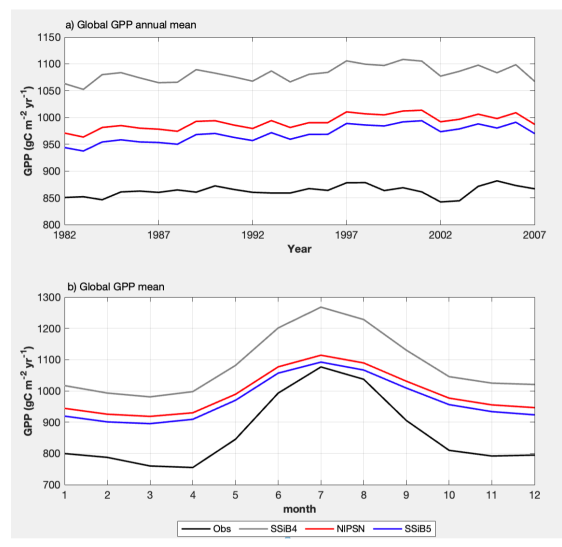


Figure 10. Intercomparisons of global monthly/annual mean GPPs among OBS, SSiB4 (control), NIPSN (N limitation on photosynthesis only), and SSiB5 (all N processes) over the period <u>of</u> 1982-2007.

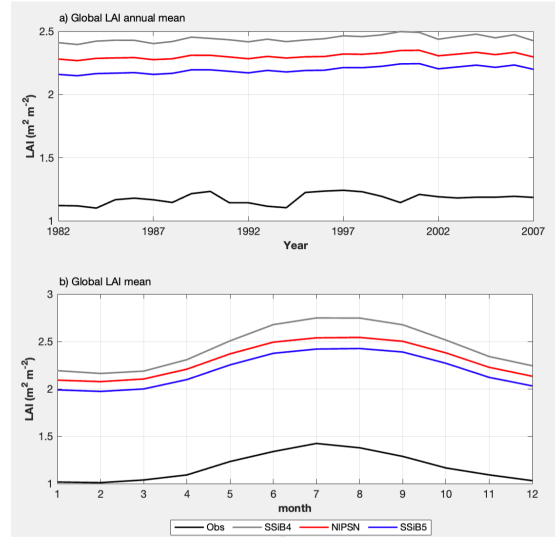
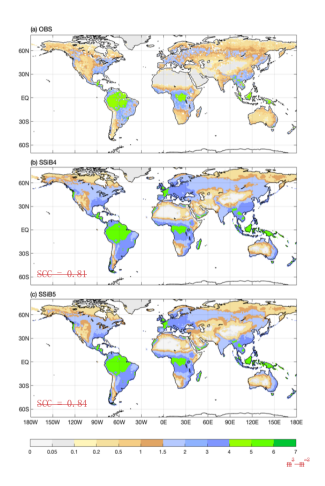
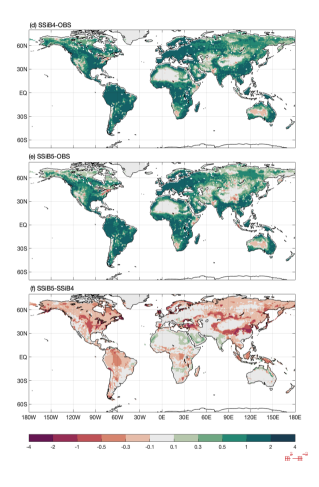


Figure 11. Same as figure Figure 10, but for the LAI. Note: Observed LAI is the GIMMS LAI.

Furthermore, the N-limitation-effect of N limitations on the LAI simulation iswas also investigated. Both-Compared with satellite-derived products, both SSiB4 and SSiB5 produceexhibit reasonable spatial distribution compared with satellitederived productsdistributions (Figs. 12a-c). The highest LAI occurs in the tropical evergreen forestforests and decreases with latitude in both the observations and the model (Fig. 9b). Compared with the control, Exp. SSiB5 also generally reducesreduced the positive bias in the simulated LAI compared to the control (Fig. 12f). The simulated LAI in Exp. SSiB4 has a global positive bias. After introducing the three N limitation processes, the positive bias is reduceddecreased over most parts of the world (Fig. 12f). Globally, Exp. SSiB5 has an LAI bias of 0.94/1.12 for GIMMS/GLASS, respectively (Table 78), which is lower than the LAI bias of 1.26/1.44 for GIMMS/GLASS, respectively, in Exp. SSiB4, with a substantial 31.1% reduction in the bias (compared to GIMMS, Table 78). However, thea substantial positive bias still exists substantially overacross the globe (Fig. 12e). In a land model intercomparison, the positive LAI prevailed in almost every dynamic vegetation model (Murray-Tortarolo et al., 2013). Our study showsshowed that imposing N limitation is an adequate step to overcome the dynamic vegetation models' systematic LAI positive bias, but thethis issue ishas still not solvedbeen resolved and requires morefurther investigation. -In addition, the correlation coefficients between the observed and simulated monthly/annual average LAIs are improved from 0.49/0.97 (Exp. SSiB4) to 0.51/0.98 (Exp. SsiB5) (Fig. 11).





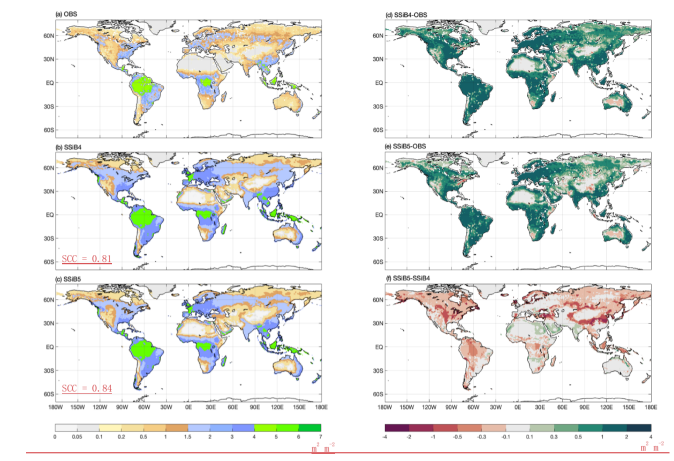


Figure 12. Same as Figure 8, but for the LAI.

Note: SCC indicates the spatial correlation coefficient between the model simulation and the GIMMS LAI (OBS).

It is interesting to note that despite the global general LAI reduction, the SsiB5SSiB5 slightly increased the LAI estimation in
North Africa and India (Fig. 12). The N impacts of N on phenology and respiration cause a slight shift in the vegetation from shrubshrubs (N. Africa) or C4 grassplants (India) to C3 grassgrasses in these areas, which contributes to the increases in GPP and LAI increase (Fig. 13). Furthermore, in areas such as the Amazon, and East Asian mixed forest-grassland regions, SSiB5 improved only improves the LAI simulation, and not the GPP simulation. We will further identify the effect of N limitation on the photosynthesis process and other processes on simulated GPP and LAI. More observational data are necessary to gain more understanding.

Table 78. Regional and Globalglobal LAI for (a) GIMMS LAI (observation), (b) GLASS LAI (second observation), (c) SSiB4 (control), (d) NIPSN (N limitation on photosynthesis only) and (ed) SSiB5 (N limitation on photosynthesis, autotrophic respiration, and phenology). The bias is relative to the GIMMS LAI.

		LAI Mean (m ² m ⁻²)										
Regions	Sub-regions	GIMMS		(GLASS		SSiB4		NIPSN		SSiB5	
		mean	bias	mean	bias	mean	bias	mean	bias	mean	bias	
Arid and Semi- Arid Areas	West Africa	1.08		1.01	-0.07(-6.5%)	2.04	0.96(88.9%)	1.89	0.81(75.0%)	1.73	0.65(60.2%)	
	West NA	0.62		0.49	-0.13(-21.0%)	1.38	0.76(122.6%)	1.18	0.56(90.3%)	1.09	0.47(75.8%)	
	SA Savanna	1.99		1.91	-0.18(-4.0%)	3.34	1.35(67.8%)	3.23	1.24(62.3%)	2.97	0.98(49.2%)	
	East Africa	1.59		1.55	-0.04(-2.5%)	3.02	1.43(89.9%)	2.89	1.30(81.8%)	2.66	1.07(67.3%)	
	East Asian semi-arid	1.60		1.36	-0.24(-15.0%)	3.35	1.75(109.4%)	2.84	1.24(77.5%)	2.61	1.01(63.1%)	
NH High-Mid	NA High-Mid Latitude	0.84		0.49	-0.35(-41.7%)	1.91	1.07(127.4%)	1.66	0.82(97.6%)	1.53	0.69(82.1%)	
Latitude Areas	Eurasian High-Mid	1.14		0.57	-0.57(-50.0%)	2.29	1.15(100.9%)	2.08	0.94(82.5%)	1.91	0.77(67.5%)	
Equator	Amazon Basin	4.19		4.08	-0.11(-2.6%)	6.01	1.82(43.4%)	5.98	1.79(42.7%)	5.50	1.31(31.3%)	
	Southeast Asia	3.93		3.88	-0.05(-1.3%)	4.68	0.75(19.1%)	4.68	0.75(19.1%)	4.31	0.38(9.7%)	
	Equator Africa	3.83		3.76	-0.07(-1.8%)	5.74	1.91(49.9%)	5.72	1.89(49.3%)	5.27	1.44(37.6%)	
Subarctic Areas and Tibet	NA Subarctic	0.32		0.14	-0.18(-56.3%)	0.71	0.39(121.9%)	0.51	0.19(59.4%)	0.47	0.15(46.9%)	
	Eurasian Subarctic	0.33		0.12	-0.21(-63.6%)	0.87	0.54(163.6%)	0.65	0.32(97.0%)	0.60	0.27(81.8%)	
	Tibet	0.64		0.54	-0.10(-15.6%)	1.36	0.72(112.5%)	0.81	0.17(26.6%)	0.75	0.11(17.2%)	
Global		1.18		1.00	-0.18(-15.3%)	2.44	1.26(110.8%)	2.31	1.13(95.8%)	2.12	0.94(79.7%)	

Note: the numbers in parentheses are relative biases

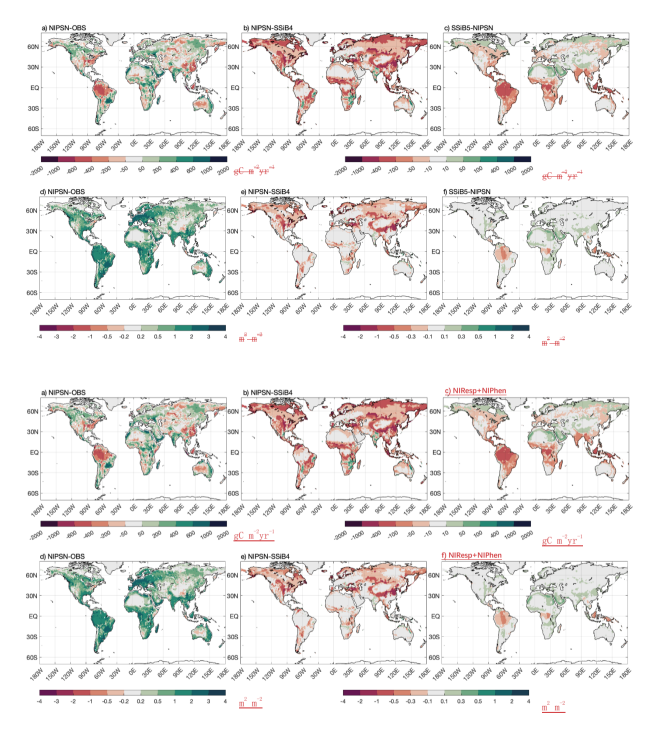


Figure 13. The 1982-2007 average gross primary production difference (a) NIPSN-OBS, (b) NIPSN-SSiB4, (c) SSiB5-NIPSNNIResp+NIPhen, and leaf area index difference (d) NIPSN-OBS, (e) NIPSN-SSiB4, (f) SSiB5-NIPSNNIResp+NIPhen

Note: NIPSN is N limitation on photosynthesis (Vc, max) only.

750 We imposed N limitation on several processes. Among them, Exp. NIPSN showshas the largest and most significant impact. The This paper mainly discusses the results from Exp. NIPSN, which apply applies Eq. (6) to scale down the V_{cmax} , are discussed here. Please note, the differences between Exp. SSiB5 and Exp. NIPSN show the effects due to another two N limitation processes (Eqs. 10 and 13), Exp. Exp. NIPSN has a lower global GPP bias (128.52 g C m⁻² vr⁻¹) compared to FLUXNET MTE estimates than does Exp. SSiB4 does (219.50 g C m⁻² yr⁻¹) (Fig. 13, Table 67), but it is larger than that of Exp. SSiB5, in which the bias is 79 g C m⁻² yr⁻¹ (Table 67). In addition, Exp. NIPSN has a global LAI bias of 1.13 (Fig. 13, 755 Table 78), which is also lower than the LAI bias in Exp. SSiB4 (1.26); but higher than that in Exp. SSiB5 (0.94). The largest reductions in the magnitude of the LAI bias are in North America, the Eurasian continents continent, and tropical Savanna regions in South America and Africa (Figs. 13b and 13e). That said, the N limitation on of photosynthesis plays a dominant role, contributing to about approximately 65%/41% of the improvement forin the GPP/LAI simulations in 760 Exp. SSiB5, respectively. Adjusting $V_{c,max}$ is the most direct and process-based approach based on physiology and yieldshas the largest greatest impact. But However, the effects of N limitation on the other two processes is are still substantial. The N limitations on respiration and phenology have the most greatest impact in tropical forest and savanna regions (Figs. 13c and 13f). For The GPP, they also reducereduced the positive bias over boreal regions and the negative bias in theover polar regions. The results from Exp. NIResp or Exp. NIPhen individually did not show a statistically significant impact. However, the sum 765 of these two N limitations still has substantial impacts on many parts of the world, as displayed in Fig. 13b, mainly in tropical rainforests and some midlatitude regions. In addition, the differences between Exp. SSiB5, which includes three limitations, and Exp. NIPSN, as displayed in Figs. 10 and 11, also delineate the characteristics of the global impacts of these two effects at seasonal and interannual scales.

770 5 Discussion and Conclusions

This study presents improvements in modellingmodeling the C cycle by introducing plant N processes into—the SSiB5/TRIFFID/DayCent-SOM, using DayCent-SOM to obtain the amount of N available to plants and plant soil N uptake. The approach presented in this study can also be applied to other models with similar physical and biological principles. The new C-N coupling framework allows us to use <u>the_dynamic C/N ratiosCNR</u> to represent plant resistance and self-adjustmentresponse, which allowallows adaptations in the stoichiometry of C and N. Since these processes can increase nutrient use efficiency and reduce the impact of N limitation through N remobilization and resorption, N-limit<u>the N limitation</u> effect woulddoes not linearly noror instantaneously respond to <u>the</u> available N content. A linear relationship between <u>the</u> N limitation factor and available N is valid only when N availability is not sufficient for the minimum N demand for new growth. This is an advantage of our approach. That said, withWith the new model structure, Nthe impacts of N on GPP are predicted

780 directly but not linearly with leaf N content, which is affected by the state of plant growth, autotrophic respiration, and plant phenology.

By comparing site-level results from SSiB4 and SSiB5 to FLUXNET GPP and surface heat fluxes from twelvethirteen sites with representative biome types and climate zones, we gained confidence in the ability of the new N processes to enhance global model performance. We also evaluated the model performance against global satellite product data sets datasets for GPP

- 785 and LAI. In general, with the new plant C-N coupling framework, SSiB5/TRIFFID/DayCent-SOM producesproduced significantly less absolute bias for GPP and LAI than did the baseline version of SSiB4/TRIFFID (without N processes), with a global decrease ofin the bias in GPP and LAI byof 16.3% and 27.1%, respectively. The main improvements are found in tropical Africa and the boreal forest. The more realistic representation of the dynamic C/N ratiosCNR and plant C-N framework leads to general improvements in SSiB5/TRIFFID/DayCent-SOM'sthe global C cycling simulations_of
- 790 <u>SSiB5/TRIFFID/DayCent-SOM</u>. From the perspective of plant physiology (Högberg et al., 2017), the downregulation of the canopy photosynthetic rate based on the available mineral N for <u>the</u> growth of plant tissues is more reasonable than the simple and direct downregulation of GPP or NPP. In fact, we have conducted a test to directly downscale GPP and NPP, and our simulation results (not shown) support this viewpoint. This coupled model can better reproduce observed state variables and their emergent properties (such as GPP, NPP, LAI, and respiration).
- 795 Despite the general improvement globally, the GPP simulation in temperate East Asian mixed forest-grassland regions seems to be worse with SSiB5 than with SSiB4. In some regions, such as the Amazon, while SSiB4 produced lower GPP than did the observations, the imposed N limitation in SSiB5 further increased the bias in these regions. This mismatch is a common issue reported in a number of publications (Anav et al., 2015; Liu et al., 2019; Piao et al., 2013). Further investigations are necessary. Recently, the important influence of phosphorus availability on terrestrial ecosystem carbon uptake has been
- 800 increasingly recognized (Du et al., 2020). Recently, initiated ecosystem-scale manipulation experiments in phosphorus-poor environments (Fleischer et al., 2019) call for the need for new phosphorus-enabled LSMs to track these actions (Goll et al., 2017; Reed et al., 2015). We plan to incorporate other plant processes, such as plant/soil phosphorus processes, to further improve the performance of the model in the future. More evaluations for regions where N limitation is not dominant are necessary.
- 805 Although the global GPP of SSiB5 was similar to that of the satellite- derived GPP, the positive bias for the LAI was still very large (Table 7). Recent review papers seem to confirm that overestimation of LAI is a common issue in current dynamic vegetation models. Murray-Tortarolo et al. (2013) and Anav et al. (2013) evaluated the performance of dynamic vegetation models in simulating LAI from a CMIP model intercomparison. The simulated LAI for almost every dynamic vegetation model is twice as large as the satellite-derived LAI. More recent studies (Zaehle et al., 2015; Mueller et al., 2019; Gristina et al. (2013) evaluated the performance of dynamic vegetation model is twice as large as the satellite-derived LAI.
- 810 al., 2020; Oliveira et al., 2021; Heikkinen et al., 2021) have confirmed this shortcoming in current dynamic vegetation models. <u>Further investigations are necessary.</u> <u>Anthropogenic N input is one of the major factors affecting C–N coupling and N limitation. The anthropogenic N inputs to</u> terrestrial ecosystems have been much greater than the vegetation N fixation in recent decades in some areas, such as eastern

China and the central USA, which can relieve N limitations (Tian et al., 2022). Due to the scope of this paper, this issue is not

815 addressed in this paper but is an important subject for further investigation to comprehensively understand the N limitation effect. Finally, this is an offline experiment in which the atmospheric forcing (such as downward radiation) is fixed. With a fixed atmospheric demand, the heat flux response due to the N limitation effect is also limited, as shown in section 4.1. A comprehensive assessment of the effect of N limitation on heat fluxes and atmospheric circulation needs to be conducted in a fully coupled atmosphere–land model.

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Data availability. The evaluation/reference data sets from model data discussed in this paper are archived at https://doi.org/10.5281/zenodo.7196869

825 Code availability. The source code of biophysical-ecosystem-biogeochemical model, SSiB version5/TRIFFID/DayCent-SOM is archived at https://doi.org/10.5281/zenodo.7297108

Author contributions. ZX, YX, MH, and YL designed the coupling strategy between SSiB4/TRIFFID and DayCent-SOM. ZX conducted the simulation with suggestions from YX, WG, and WP. ZX, YX, and MH drafted the text and ZX made the figures. All authors (ZX, YX, WG, MH, YL and WP) have contributed to the analysis and the text.

Competing interests. The authors declare that they have no conflict of interest.

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