



Evaluating atmospheric CO₂ effects on gross primary productivity and net ecosystem exchanges of terrestrial ecosystems in the conterminous United States using the AmeriFlux data and an artificial neural network approach



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ABSTRACT

Quantitative understanding of regional gross primary productivity (GPP) and net ecosystem exchanges (NEE) and their responses to environmental changes are critical to quantifying the feedbacks of ecosystems to the global climate system. Numerous studies have used the eddy flux data to upscale the eddy covariance derived carbon fluxes from stand scales to regional and global scales. However, few studies incorporated atmospheric carbon dioxide (CO₂) concentrations into those extrapolations. Here, we consider the effect of atmospheric CO₂ using an artificial neural network (ANN) approach to upscale the AmeriFlux tower of NEE and the derived GPP to the conterminous United States. Two ANN models incorporating remote sensing variables at an 8-day time step were developed. One included CO₂ as an explanatory variable and the other did not. The models were first trained, validated using eddy flux data, and then extrapolated to the region at a 0.05° × 0.05° (latitude × longitude) resolution from 2001 to 2006. We found that both models performed well in simulating site-level carbon fluxes. The spatially-averaged annual GPP with and without considering the atmospheric CO₂ were 789 and 788 g C m⁻² yr⁻¹, respectively (for NEE, the values were -112 and -109 g C m⁻² yr⁻¹, respectively). Model predictions were comparable with previous published results and MODIS GPP products. However, the difference in GPP between the two models exhibited a great spatial and seasonal variability, with an annual difference of 200 g C m⁻² yr⁻¹. Further analysis suggested that air temperature played an important role in determining the atmospheric CO₂ effects on carbon fluxes. In addition, the simulation that did not consider atmospheric CO₂ failed to detect ecosystem responses to droughts in part of the US in 2006. The study suggests that the spatially and temporally varied atmospheric CO₂ concentrations should be factored into carbon quantification when scaling eddy flux data to a region.

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1. Introduction

Net ecosystem carbon exchange (NEE) and gross primary productivity (GPP) are two major fluxes involved in the ecosystem biogeochemical carbon process. Quantification of GPP and NEE

and their responses to environmental changes would improve our understanding of the ecosystem carbon cycling and its feedbacks to the global climate system. At present, GPP and NEE at regional scales are often quantified using atmosphere inverse models (e.g. Prince and Goward, 1995), process-based biogeochemical models (e.g. White et al., 2000), and satellite remote sensing approaches (e.g. Running et al., 2000). The estimation of NEE based on the inverse models is limited by the sparseness of the carbon dioxide (CO₂) observation network (Tans et al., 1990); and this approach cannot differentiate the carbon source/sink contributions of each

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ecosystem (Janssens et al., 2003). Process-based models generally first derive the GPP and ecosystem respiration, which are modeled as a function of physical and biological variables and thus can be applied at regional and global scales. However, large uncertainty still exists in current quantification due to complex model structure, uncertain parameters, and model input (Chen et al., 2011). Satellite remote sensing approaches have advantages in GPP quantification that is based on a near real-time dataset, instead of broadly parameterized ecosystem models. However, they are highly dependent on the accuracy of the model algorithms and the quality of satellite imageries (Baldocchi et al., 2001).

The eddy covariance technique provides direct measurement of net carbon and water fluxes between vegetation and the atmosphere (Baldocchi et al., 1988; Foken and Wichura, 1996; Aubinet et al., 1999). At present, over 500 flux tower sites have been operated to measure the exchanges of carbon fluxes continuously over a broad range of climate and biome types (FLUXNET <http://daac.ornl.gov/FLUXNET/fluxnet.shtml>). These towers also provide calibrated, validated NEE data and the derived GPP product. To date, numerous studies have been conducted using those flux data to explore the GPP and NEE temporal or spatial variation and their controlling factors in terrestrial ecosystems (Valentini et al., 2000; Law et al., 2002; Baldocchi, 2008; Hirata et al., 2008; Kato and Tang, 2008; Lund et al., 2010; Bracho et al., 2012). Eddy flux data have also been widely used for model calibration (Baldocchi, 1997; Hanan et al., 2002; Reichstein et al., 2002; Hanson et al., 2004) and to upscale from stands to regional levels. The upscaling exercises generally employ the machine learning algorithm that uses meteorological data, vegetation properties, and remote sensing products, which have been conducted for Asia (Zhu et al., 2014; Fu et al., 2014), Europe (Jung et al., 2008; Vetter et al., 2008), the US (Yang et al., 2007; Xiao et al., 2008; Xiao et al., 2010) and even at global scales (Beer et al., 2010; Yuan et al., 2010).

However, few studies have incorporated the atmospheric CO₂ concentrations into regional GPP and NEE extrapolations when using eddy flux tower data. Increasing CO₂, which is mainly due to the burning of fossil fuels and, to a much lesser extent, land-cover change (Keeling et al., 1995; Hartmann et al., 2013), will have substantial direct and indirect effects on the carbon budget (Canadell et al., 2007). Numerous studies have been conducted to understand how plants and ecosystems will respond to elevated CO₂ (Ainsworth and Long, 2005). The large-scale free-air CO₂ enrichment experiment (FACE) showed that increasing CO₂ would reduce the stomatal conductance (Curtis and Wang, 1998; Medlyn et al., 2001; Ainsworth and Rogers, 2007) over a short time period, resulting in decreased transpiration, enhanced water-use efficiency (Conley et al., 2001; Wall et al., 2001) and increased soil moisture (Bunce, 2004). In addition, stimulated photosynthesis (e.g. Li et al., 2014) would increase the above and below-ground biomass production (Piao et al., 2006; Los, 2013; Wan et al., 2007; Deng et al., 2010) and hence accelerate the CO₂ loss from heterotrophic respiration (Luo et al., 1996). Generally, long-lived plant biomass (trees) is more responsive to increasing CO₂ than herbaceous species (Ainsworth and Long, 2005). However, increased plant production with high carbon (C) to nitrogen (N) ratio under elevated CO₂ results in low quality litter input and slows the soil N mineralization, thus triggering the negative feedback to plant biomass over time (Oren et al., 2001; Gill et al., 2002; Luo et al., 2004) if there is no extra nitrogen input. This conceptual framework about progressive N limitation is more obvious in long-lived plants (trees) (Luo et al., 2004) and has been supported by few experiments (Oren et al., 2001; Norby et al., 2010). In addition, the climate drivers and their interaction with soil resources (Reich et al., 2014) would also constrain the ecosystem response

to CO₂ fertilization. For example, increasing N mineralization due to warming temperature would further promote plant growth (Peltola et al., 2002; Dijkstra et al., 2010) whereas the drought stress may diminish this positive effect under elevated CO₂ condition (Dermody et al., 2007; Larsen et al., 2011). Moreover, altered water, energy balance (Drake et al., 1997; Keenan et al., 2013), and vegetation physiology due to increasing CO₂ would in turn affect the carbon cycling. Therefore, incorporating atmospheric CO₂ concentrations into the regional GPP and NEE estimation should be a research priority. In addition, previous upscaling flux studies that exclude the atmospheric CO₂ concentration assume that CO₂ variations were spatially and temporally uniform in a region. Although atmospheric CO₂ is generally well-mixed globally since it is chemically inert (Eby et al., 2009), it actually exhibits a large seasonal and spatial variability at the regional scale (Miles et al., 2012). The seasonal and spatial characteristics have previously been reported at site and regional levels (Davis et al., 2003; Haszpra et al., 2008; Miles et al., 2012). For example, the summer measurement of the atmospheric boundary layer CO₂ mole fraction from a nine-tower regional network deployed during the North American Carbon Program's Mid-Continent Intensive (MCI) during 2007 to 2009 shows that the seasonal CO₂ amplitude is five times larger than the tropospheric background and the spatial gradient across the region is four times the inter-hemispheric gradient (Miles et al., 2012).

In this study, we used an artificial neural network (ANN) approach to upscale the AmeriFlux tower derived GPP and NEE to the conterminous US from 2001 to 2006 by considering the atmospheric CO₂ concentrations as an independent factor. We developed two sets of ANN models linking GPP and NEE and remote sensing variables for each vegetation type: the first considered the atmospheric CO₂ concentration effects (CO₂ incorporated simulation, S0) and the other did not (non-CO₂ incorporated simulation, S1). After the training and validation procedure, we then used the two models to extrapolate GPP and NEE to the conterminous US. We hypothesized that: (1) both models would capture the GPP and NEE variation at the calibration stage; (2) the two simulations would exhibit spatiotemporal differences which associate with climate drivers; and (3) ecosystems with high productivity (e.g., forest, cropland) would show greater differences between the two simulations.

2. Method

2.1. Overview

From the AmeriFlux network, we selected the key ecosystem types in the conterminous US including: evergreen forest, deciduous forest, grassland, mixed forest, savannas, shrubland, and cropland. Day land surface temperature (D.LST) and night land surface temperature (N.LST), leaf area index (LAI), fraction of absorbed photosynthetically active radiation (fPAR), normalized difference vegetation index (NDVI), enhanced vegetation index (EVI), CO₂ measurement and derived GPP and NEE from the eddy flux towers were organized into an 8-day time step. To explore the atmospheric CO₂ concentrations' effect on the GPP and NEE, we constructed two sets of ANN models for each vegetation type: one incorporated CO₂(S0) in addition to D.LST, N.LST, LAI, fPAR, NDVI and EVI; and the second did not incorporate with CO₂ (S1) with the same explanatory variables in S0. After the training and validation of the two sets of ANN models, the regional estimation of GPP and NEE was driven by the explanatory variables at a spatial resolution of 0.05° × 0.05° on an 8-day time step. The difference between S0 and S1 was then analyzed, considering the controls of environmental variables.

Table 1
AmeriFlux Sites used in ANN models.

Site name	Site ID	Vegetation type	Data range
ARM SGP Control	US-Arc	Grasslands	2005–2006
ARM SGP Main	US-ARM	Cropland	2003–2006
Bartlett Experimental Forest	US-Bar	Deciduous broadleaf forest	2004–2005
Blodgett Forest	US-Blo	Evergreen needleleaf forest	2000–2006
Duke Forest Hardwoods	US-Dk2	Mixed Forest	2001–2005
Duke Forest Loblolly Pine	US-Dk3	Evergreen needleleaf forest	2001–2005
Fermi Agricultural	US-IB1	Cropland	2005–2006
Fermi Prairie	US-IB2	Grasslands	2004–2006
Fort Dix	US-Dix	Mixed Forest	2005–2006
Freeman Ranch Mesquite Juniper	US-FR2	Woody savannas	2004–2006
Harvard Forest	US-Ha1	Deciduous broadleaf forest	2000–2006
Harvard Forest Hemlock	US-Ha2	Evergreen needleleaf forest	2004
Howland Forest Main	US-Ho1	Evergreen needleleaf forest	2000–2004
Kendall Grassland	US-Wkg	Grasslands	2004–2006
Kennedy Space Center Scrub Oak	US-KS2	Shrubland	2000–2006
Mead Rainfed	US-Ne3	Cropland	2001–2005
Mead Irrigated	US-Ne1	Cropland	2001–2005
Mead Irrigated Rotation	US-Ne2	Cropland	2001–2005
Metolius Intermediate Pine	US-Me2	Evergreen needleleaf forest	2003,2005
Metolius Second Young Pine	US-Me3	Evergreen needleleaf forest	2004–2005
Morgan Monroe State Forest	US-MMS	Deciduous broadleaf forest	2000–2005
Lost Creek	US-Los	Shrubland	2001–2005
Missouri Ozark	US-MOz	Deciduous broadleaf forest	2004–2006
North Carolina Loblolly Pine	US-NC2	Evergreen needleleaf forest	2005–2006
Ohio Oak Openings	US-Oho	Deciduous broadleaf forest	2004–2005
Santa Rita Mesquite Savanna	US-SRM	Woody savannas	2004–2006
Sky Oak Old	US-SO2	Shrubland	2005–2006
Sky Oak Young	US-SO3	Shrubland	2005–2006
Sylvania Wilderness	US-Syv	Mixed Forest	2002–2006
Tonzi Ranch	US-Ton	Woody savannas	2001–2006
Univ of Mich Biological Station	USMS	Mixed Forest	2000–2003
Vaira Ranch	US-Var	Grasslands	2001–2006
Willow Creek	US-Wcr	Deciduous broadleaf forest	2000–2006
Wind River Field Station	US-Wrc	Evergreen needleleaf forest	2000–2004
Wisconsin Intermediate Hardwoods	US-Wi1	Deciduous broadleaf forest	2003
Wisconsin Mature Red Pine	US-Wi4	Evergreen needleleaf forest	2002–2005

2.2. Explanatory variables

NEE is the difference between photosynthesis and ecosystem respiration, which is influenced by a variety of physical, physiological, hydrological, and atmospheric variables. As for the plant photosynthesis, current ecosystem models have incorporated various mechanisms related to these factors to simulate GPP, such as the bio-chemical model (e.g. Farquhar et al., 1980), light use efficiency equation (e.g. Running et al., 2000), and the semi-empirical relationship (e.g. Raich et al., 1991). Ecosystem respiration (R_e) includes autotrophic (R_a) and heterotrophic respiration (R_h). R_a is generally modeled as a function of temperature and plant biomass, while R_h is empirically modeled as a function of soil temperature, moisture, and soil organic carbon stocks. Many of these factors influencing GPP and NEE can be assessed by satellite remote sensing products. NDVI is closely correlated to the photosynthetic activity, vegetation biomass, and fractional vegetation cover, which have been widely used in production efficiency models. We also incorporated the EVI to improve the ANNs performance due to the several limitations of normalized difference vegetation index (NDVI), including the saturation in a multilayer closed canopy and sensitivity to atmosphere aerosols and soil background. The LST derived from the Moderate Resolution Imaging Spectroradiometer (MODIS) corresponds the surface soil temperature, which is known to strongly control the R_e as both R_a and R_h . Previous studies demonstrated that satellite-derived LST was strongly correlated with R_e , especially in dense-vegetated ecosystems (Rahman et al., 2005). The estimation of LAI and fPAR, characterizing the vegetation canopy function and energy absorption capacity, are key parameters in most biogeochemical models due to the high correlation with GPP. Taken together, we selected LST, NDVI, EVI, fPAR,

and LAI as explanatory variables to account for the GPP and NEE variation.

2.3. Data organization

We organized AmeriFlux site-level data of carbon flux and meteorology, MODIS land-cover, spatially and seasonally explicit CO_2 , and the remote sensing products derived from MODIS. The AmeriFlux network coordinates regional analyses of observations from eddy covariance flux towers across the US. The Level-3 and Level-4 data for 36 AmeriFlux sites over the period 2000–2006 that contained CO_2 measurements were ultimately organized (Table 1 & Fig. 1). In the Level-3 product, the half-hourly CO_2 was aggregated into 8-day time step values. The Level-4 product consists of two types of GPP and NEE data: standardized (GPP_{st} and NEE_{st}) and original (GPP_{or} and NEE_{or}). Here, we used the same method in Xiao et al. (2010) to select the type of GPP and NEE product. Explanatory variables, GPP and NEE values derived from the selected sites were organized to develop the ANN models for each ecosystem type.

We used the following four MODIS data products (Collection 4), including daytime and nighttime LST (MOD11A2; Wan et al., 2002), EVI (MOD13A1; Huete et al., 2002), and LAI/fPAR (MOD15A2; Myneni et al., 2002). EVI is at a spatial resolution of 500 m × 500 m, while LAI, fPAR, and LAI are at spatial resolutions of 1 km × 1 km. These MODIS data were resampled to the same resolution of 0.05° × 0.05° (longitude × latitude). The explicitly spatial and seasonal CO_2 data was obtained from the National Oceanic and Atmospheric Administration (NOAA) gridded CO_2 product (<ftp://afftp.cmdl.noaa.gov/products/carbontracker/co2/>). The product is based on global CO_2 observation network data, employing

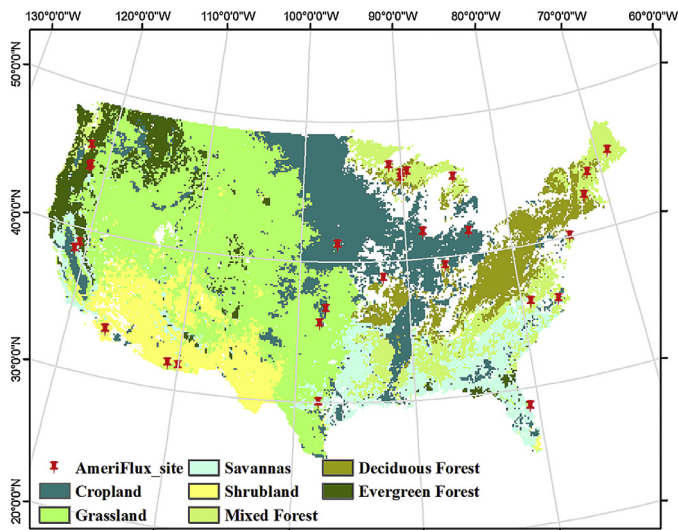


Fig. 1. Selected AmeriFlux tower sites in this study.

a novel ensemble assimilation method to accurately model atmospheric CO₂ mole fractions (Peters et al., 2007). Land cover was obtained from the MODIS product Land Cover Types Yearly L3 Global 0.05 Deg CMG (MOD12C1) (Year 2005) from the NASA Goddard Space Flight Center website (<http://modis-land.gsfc.nasa.gov>). The classification of the International Geosphere and Biosphere (IGBP) land-cover classification system was used to reclassify the land-cover map of the conterminous United States into seven major vegetation types in the ANN model. To be more specific, the evergreen needleleaf forest and evergreen broadleaf forest were combined into evergreen forest (same for deciduous forest); and closed shrubland and open shrubland were merged into shrubland. Therefore, the model calibration and simulation were based on the seven vegetation types: evergreen forest, deciduous forest, grassland, mixed forest, savannas, shrubland, and cropland.

2.4. Neural network development

We used a General Regression Neural Network (GRNN) algorithm (Specht, 1991) to represent the input-output relationship between the independent variables and dependent variables. The GRNN scheme can model the nonlinear system without specifying the algebraic relationship or internal mechanism. With the advantage of a fast-learning speed and good convergence with a large number of training datasets, GRNN can be a useful tool to estimate GPP and NEE based on the explanatory environmental variables. The training data set, including input and output values of measurements, is fed into the multilayer neural network. The network is trained to obtain a set of optimized interconnected network weights, which are used to produce the most probable value for the outputs. More details about the GRNN algorithm and optimization procedure are provided in the supplementary information (GRNN algorithm and Fig. S1). To develop the GRNN model for each ecosystem type, we first randomly divided each dataset into two sets: a training set (75% of whole data for each vegetation type) used to construct the GRNN model and a testing set (25% of whole data for each vegetation type) used to evaluate the performance of constructed GRNN. MATLAB codes were used for developing the model (The MathWorks Inc., Natick, MA, USA).

The uncertainty of GRNN model prediction is mainly from three sources: input variables, ANN model structure, and model parameters. Here we focused on the uncertainties associated with the model structure because we do not have sufficient knowledge about the uncertainty of input variables. We used the “delete-one”

method (Zhuang et al., 2012) through developing a number of alternative ANN models to quantify the uncertainty of GPP and NEE due to uncertain model structures. To be more specific, we randomly selected 75% of the observed data for each ecosystem type to develop a new ANN model. The new model was then scaled up to obtain the new GPP and NEE estimation at the regional scale. The process was repeated for 100 times in S0 and S1 for each ecosystem type, respectively. The averaged GPP and NEE of all estimations were used to analyze the differences among simulations. The 95% confidence intervals were considered to be the range of model uncertainty to define the upper and lower bounds of the GPP and NEE estimation.

3. Results

3.1. Annual GPP and NEE and spatial difference

The simulated GPP, NEE with the two sets of ANN models were both close to the observed GPP and NEE for each ecosystem type (Fig. S2). We applied the two sets of models to calculate the annual averaged GPP and NEE of terrestrial ecosystems for the conterminous US at an 8-day time step from 2001 to 2006 (Fig. 2a and c). Both models showed a high spatial variability with a clear gradient from west to east. The Gulf Coast and parts of the Southeastern US were the most productive regions, with GPP greater than 2000 g C m⁻² yr⁻¹. Most parts of the Southeast and coastal Pacific Northwest also possessed high GPP (1250–1500 g C m⁻² yr⁻¹). The Northeast had intermediate GPP (500–1000 g C m⁻² yr⁻¹), where was mainly dominated by deciduous forest. The Midwest, Southwest and Rocky Mountain areas generally had relatively low GPP (<500 g C m⁻² yr⁻¹). As for spatial patterns of NEE, the greatest net carbon uptake took place in the Gulf Coast and Mideast, with the annual averaged value of -800 g C m⁻² yr⁻¹. By contrast, the Rocky Mountain area had the highest contribution to carbon release across the conterminous US. The uncertainty analysis based on the 100 ANN models (Fig. S3) demonstrated that our ANNs models predicted GPP and NEE with a small uncertainty across the whole ecosystem type. Overall, both simulations indicated that the East and Pacific Northwest showed high ecosystem productivity, acting as a carbon sink, while the Rocky Mountain region had the lowest GPP value and acted as a carbon source to the atmosphere.

The two simulations however, diverged in regions including the Pacific Northwest coast, Gulf coast and most parts of Southeast dominated by evergreen forests, savannas and mixed forest (Fig. 2b and d). S0 exhibited lower GPP (-200 to -100 g C m⁻² yr⁻¹) and more positive NEE (100–250 g C m⁻² yr⁻¹) value in most Southeastern US than that of S1 (Fig. 2b). In contrast, west North Central region had higher GPP (100–200 g C m⁻² yr⁻¹) and more negative NEE (-150 to -50 g C m⁻² yr⁻¹) in S0. From the vegetation perspective, the relative difference of GPP and NEE in the mixed forest and cropland showed a spatial variability. Specifically, the lower estimation of GPP in mixed forests and cropland was mainly found in the Southeast and east of South Central, while a higher GPP was observed in the North Central in S0. These patterns were similar to the NEE estimation (Fig. 2d).

3.2. Seasonal GPP and NEE difference

Our 8-day GPP and NEE estimation based on the two ANN models both captured the seasonal flux variability in the conterminous US (Fig. 3 and Fig. S4). Take the GPP seasonal pattern, for example; in spring (March to May), the Gulf Coast and the Pacific Northwest region began to assimilate carbon at 100–200 g C m⁻² yr⁻¹, followed by the Southeast and the South Central. In contrast, most of the Northeast and Rocky Mountain area possessed a low GPP due to the relatively cold weather or late leaf-out. Plants reached

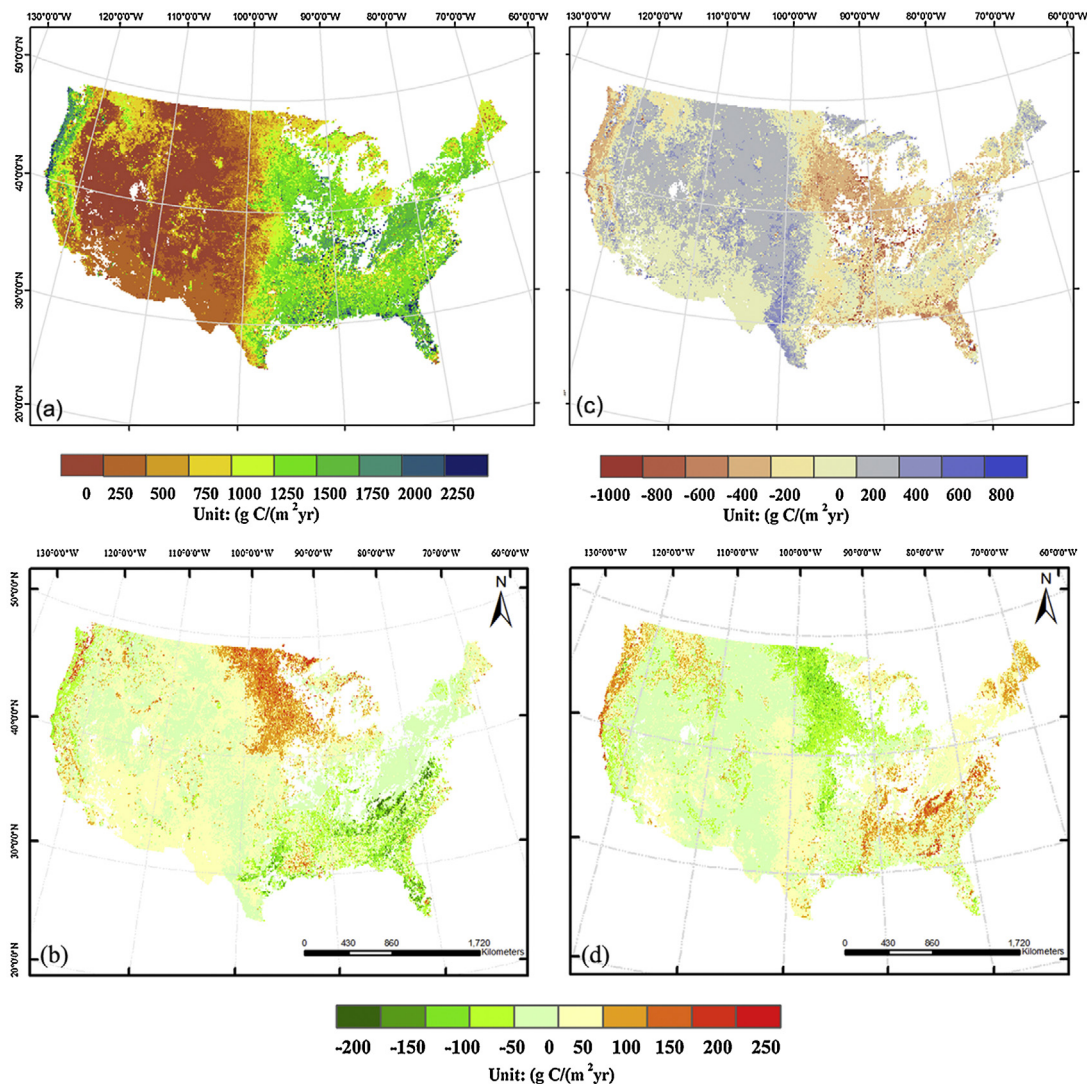


Fig. 2. Annual averaged GPP (a) and NEE (c) in S0 and their differences (GPP (b), NEE (d)) between S0 and S1 across the conterminous US scale from 2001 to 2006 (Unit: $\text{g C}/(\text{m}^2 \text{yr})$). The difference is calculated as S0 minus S1. S0 and S1 are the simulations with and without considering the atmospheric CO_2 concentration, respectively.

the greatest productivity in summer (June–August) with the highest GPP of $300\text{--}400 \text{ g C m}^{-2} \text{ yr}^{-1}$ in the Northeast regions. From fall (September–November) to winter (December–February), GPP decreased drastically to either a low level ($50\text{--}100 \text{ g C m}^{-2} \text{ yr}^{-1}$) in the Gulf Coast, Pacific Northwest and South Central or nearly zero in most parts of Rocky Mountains throughout the US.

The obvious difference between seasonal GPP and NEE generally occurred in mixed forests and croplands (Fig. 4 and Fig. S5). In the early spring (March–April), the mixed forest in the Southeast began to exhibit lower GPP and more positive NEE in S0, and this difference was much stronger from late spring (May) to early fall (September). However, for the mixed forest in North Central and the Northeast, the difference only took place in June, July, and August. And this was similar for cropland in North Central, which only showed higher GPP and more negative NEE in these three months. No obvious difference was observed between the two simulations in winter.

3.3. GPP and NEE response to CO_2 concentrations under drought condition

Annual GPP and NEE both exhibited positive and negative anomalies each year. The anomalies could have been due to

climate variability, disturbances, management practices, or model errors. The severe drought affected over 50% of the country, including the Southwest, the Great Plains, the Gulf Coast, the coastal Southeast, and particularly Texas and Oklahoma. Therefore, we evaluated the GPP and NEE response to the drought condition in 2006. Both models showed negative anomalies in most regions, like the Great Plains (Fig. 5). However, as for the Southeast and parts of North Central, the S0 exhibited the negative GPP and positive NEE anomalies when compared with S1. According to the US Drought Monitor (<http://www.drought.unl.edu>) and the PRISM database, these regions showed around 250 mm less precipitation in 2006. Without considering the CO_2 effect in upscaling GPP and NEE, the S1 failed to show the ecosystem response to severe drought condition.

4. Discussion

4.1. Comparison of S0 and S1 with other studies

Our estimation of GPP and NEE and their spatial variations agreed well with previous published results. For example, Xiao et al. (2010) estimated that the spatially averaged GPP across the conterminous US scaled at $907 \text{ g C m}^{-2} \text{ yr}^{-1}$. Our simulation showed that the Gulf Coast and parts of the Southeast were the

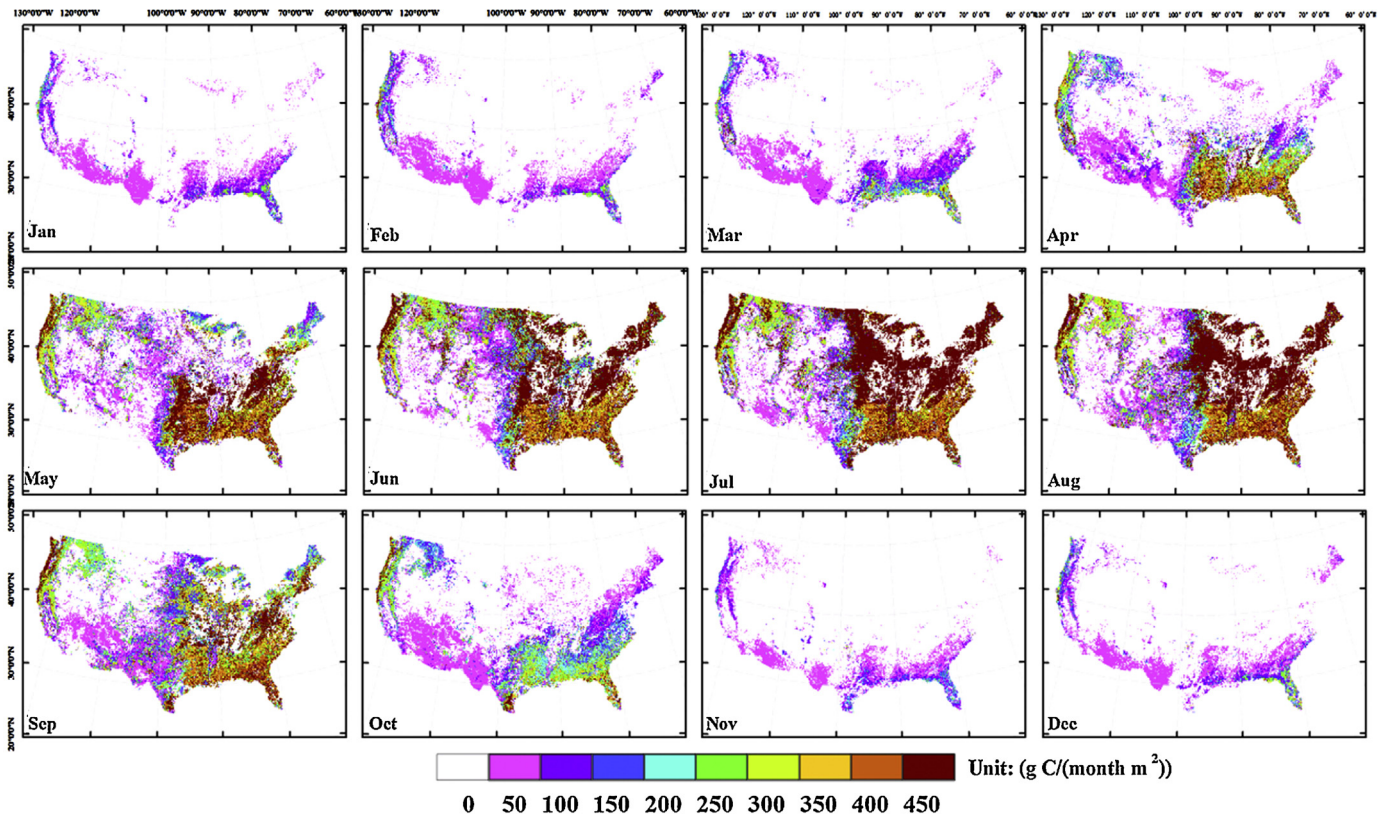


Fig. 3. Averaged monthly GPP (g C/(month m²)) for the conterminous U.S. from January through December from 2001 to 2006 in S0. S0 is the simulation with considering the atmospheric CO₂ concentration.

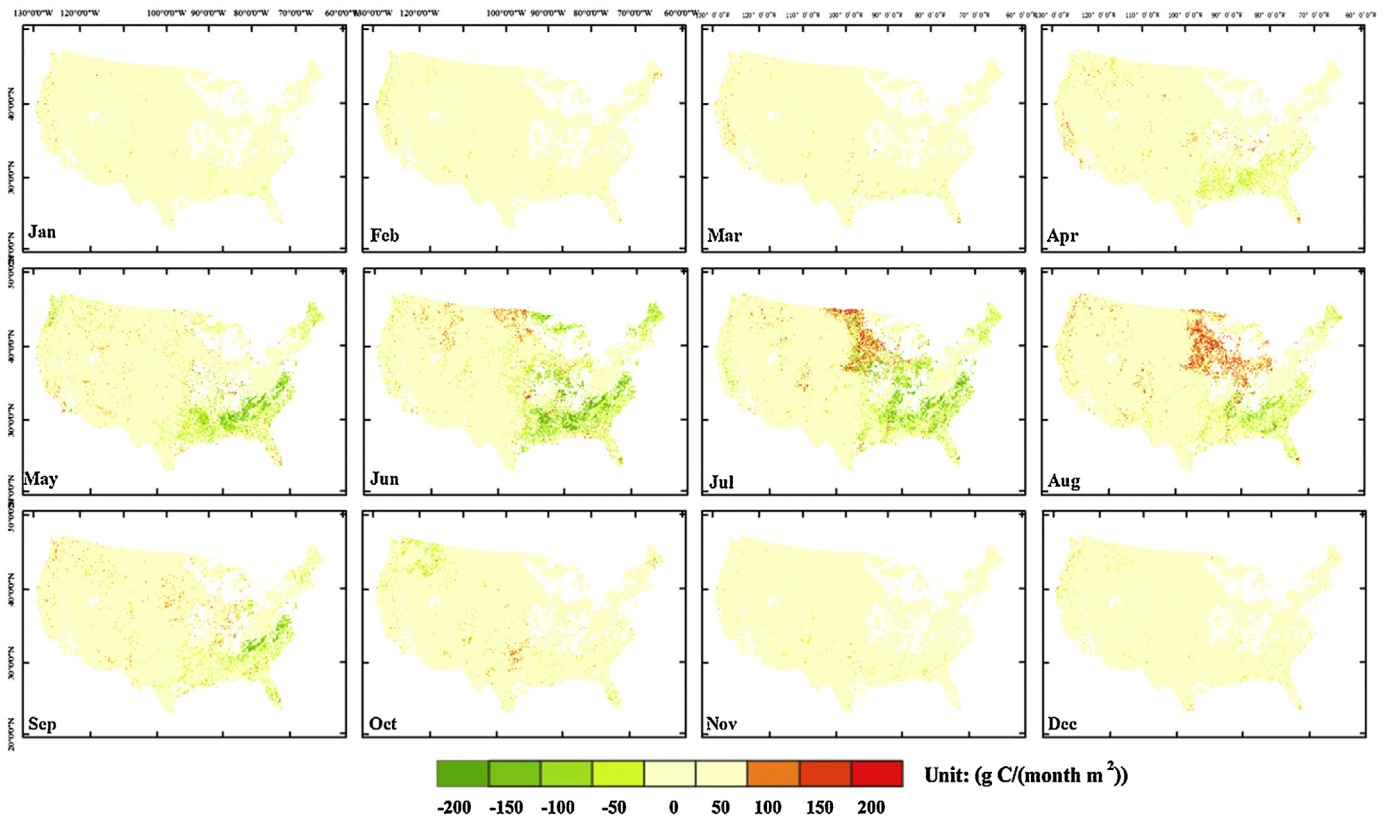


Fig. 4. Averaged monthly GPP difference (g C/(month m²)) between S0 and S1 for the conterminous U.S. from January through December from 2001 to 2006. The difference is calculated as S0 minus S1. S0 and S1 are the simulations with and without considering the atmospheric CO₂ concentration, respectively.

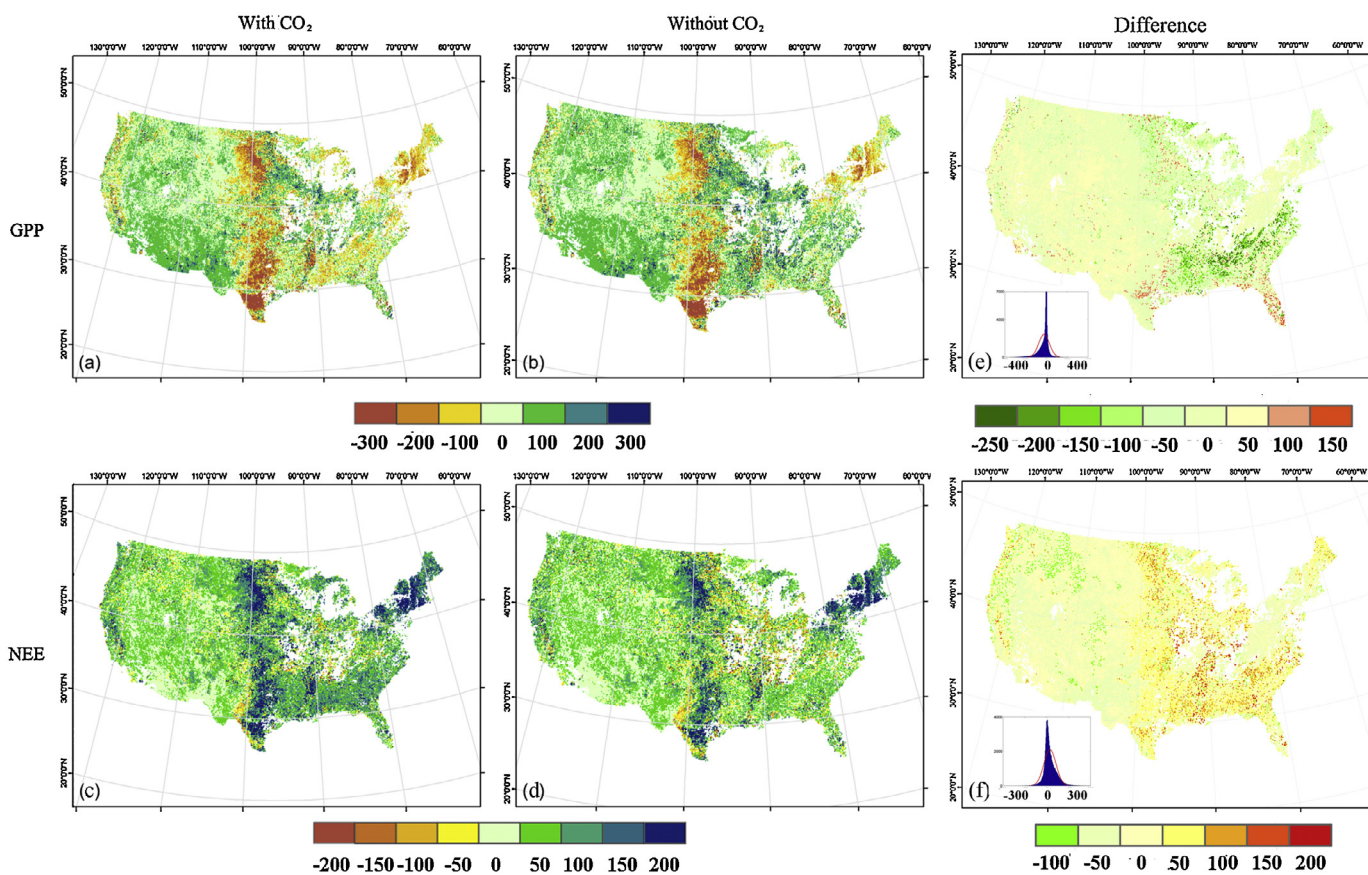


Fig. 5. Annual GPP (a and b), NEE (c and d), and anomalies in S0 and S1 for the conterminous U.S. at 2006. The anomalies of annual GPP and NEE at 2006 were relative to the 6-year average value. The anomalies difference (e and f) is calculated as S0 minus S1. S0 and S1 are the simulations with and without considering the atmospheric CO₂ concentration, respectively.

most productive regions, with GPP greater than $2000 \text{ g C m}^{-2} \text{ yr}^{-1}$, and was consistent with previous studies (Yang et al., 2007; Xiao et al., 2010; Chen et al., 2011). The total gross carbon uptake estimated in both models was 4.00 and 3.95 Pg yr^{-1} , respectively, and was lower than the results of Xiao et al. (2010) because we did not reclassify the cropland/natural vegetation mosaic in MODIS land cover product as cropland ecosystems. In addition, the estimation of GPP ($1250\text{--}1500 \text{ g C m}^{-2} \text{ yr}^{-1}$) in the Southeast was much lower than Xiao's et al. (2010) prediction ($>2250 \text{ g C m}^{-2} \text{ yr}^{-1}$), but was comparable with Chen's et al. (2011) study and the MODIS GPP product (Running et al., 2004). The spatially averaged GPP in two simulations generally agreed well with MODIS GPP (Fig. S6) ($R^2 > 0.8$). However, MODIS GPP ($<800 \text{ g C m}^{-2} \text{ yr}^{-1}$) was markedly underestimated in cropland compared with our models ($>1000 \text{ g C m}^{-2} \text{ yr}^{-1}$). This large difference may be attributable to that MODIS GPP, which was based on the light use efficiency algorithm while ANN was based on the site-level flux tower data. The key parameter, the maximum light use efficiency in the MODIS GPP algorithm, was always underestimated and thus led to lower estimation in high productivity areas (Turner et al., 2006; Xiao et al., 2010). Recently, Madani et al. (2014) used eddy flux carbon data to show that the optimum light use efficiency in cropland was twice as much as that in the MODIS GPP algorithm.

4.2. Climate controls on the CO₂ effects on GPP and NEE

Previous studies have evaluated the interactions between elevated CO₂ and other environmental variables, such as increasing temperature (Peltola et al., 2002; Tingey et al., 2003; Hovenden

et al., 2008; Dijkstra et al., 2010) and hydrological conditions (Morgan et al., 2004; Nowak et al., 2004). Increasing CO₂ with enhanced N mineralization at higher temperatures may further promote plant growth (Kirschbaum et al., 1994), whereas the positive effect may be ameliorated in a reduced precipitation environment due to the slow turnover rate of N belowground (Emmett et al., 2004; Sowerby et al., 2008). The seasonal GPP and NEE difference in our simulations also indicated there were interactions between CO₂ effect and climate drivers. Obvious GPP and NEE difference can only be observed from late spring to summer in the Northern forest ecosystem (Fig. 4 and Fig. S5), suggesting that the temperature controlling the plant phenology may play a key role in CO₂ effects on the carbon fluxes. In addition, evergreen forest in Pacific Northwest exhibited lower GPP in May and October (Fig. 4), indicating that drought or the soil water deficit (Law et al., 2000; Schwarz et al., 2004) may impact the CO₂ effects. The seasonal variation of NEE difference was slightly different from GPP (Fig. 4 and Fig. S5). NEE differences were shown earlier than GPP in spring, especially in the Northern forest. The enhanced heterotrophic respiration due to soil warming (Raich and Schlesinger, 1992; Lloyd and Taylor, 1994) may result in the asynchronicity of GPP and NEE's response to the atmospheric CO₂ effect.

We analyzed the causes for the spatial variability of GPP difference using annual averaged air temperature (T_a) and vapor pressure deficit (VPD) (Fig. 6). We obtained the T_a and VPD from North American Regional Reanalysis dataset (Mesinger et al., 2006). The atmospheric CO₂ concentration showed negative effect on the evergreen forest and the effect was more obvious under high temperature condition ($>285 \text{ K}$). Both deciduous forest and shrubland

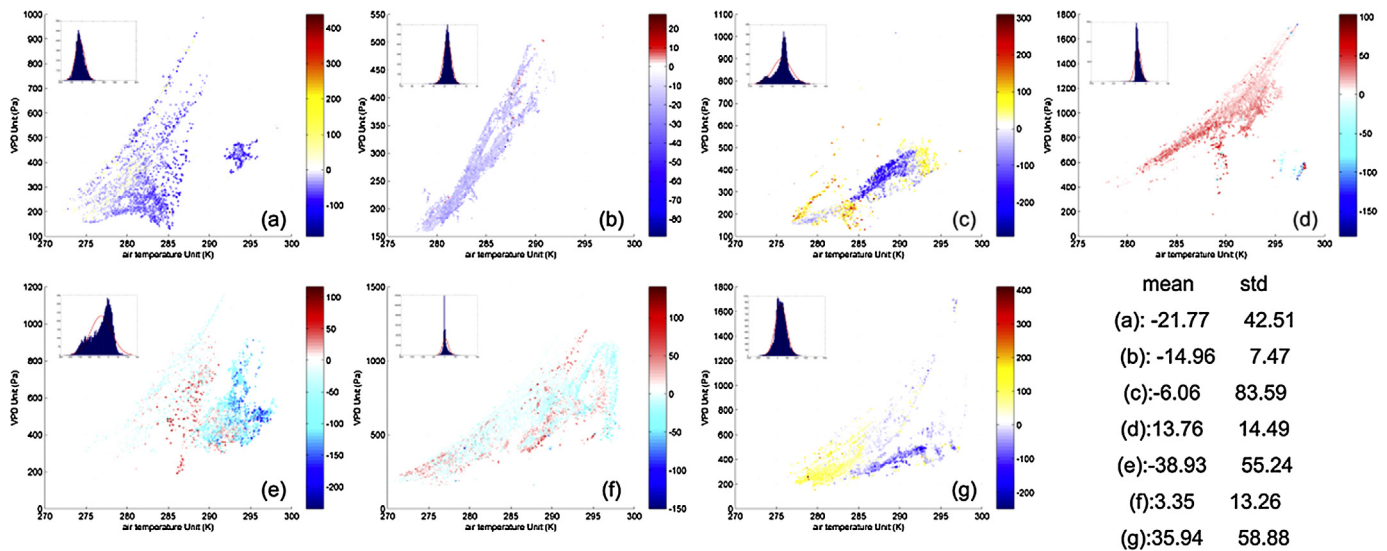


Fig. 6. Air temperature and VPD controls on the annual GPP difference (the difference is calculated as S0 minus S1) spatial variability for each vegetation type: (a) Evergreen forest; (b) Deciduous forest; (c) Mixed forest; (d) Shrubland; (e) Savannas; (f) Grassland; and (g) Cropland. The mean values and standard deviations of GPP difference in each vegetation types are presented. S0, and S1 are the simulations with and without considering the atmospheric CO₂ concentration, respectively.

did not show a obvious difference across the T_a and VPD ranges. However, for the mixed forests, the difference could be divided by the T_a value around 285 K. The GPP estimation in S0 was generally lower when $T_a > 285$ K but higher when $T_a < 285$ K, when compared with S1. This was similar for the cropland; a lower GPP estimation occurred in S0 when $T_a > 285$ K and it had a higher estimation when $T_a < 285$ K. In addition, the negative effect was much stronger in lower VPD ranges (VPD < 600 Pa), indicating that the CO₂ had stronger effect in the humid area. As for the savannas, no obvious T_a threshold was observed and the negative effect generally located in the high T_a range. CO₂ concentration showed complex effect on grassland, but overall, had a small influence on GPP estimation. Fig. 5 also gave the statistical values (mean and standard deviation) of the GPP difference between the two simulations in each ecosystem type. Although the spatially averaged GPP difference was generally small (<50 g C m⁻² yr⁻¹), the coefficient of variation (the ratio of standard variation to mean value) was great in mixed forest and evergreen forest, followed by cropland and savannas.

4.3. GPP and NEE response to atmospheric CO₂ under drought condition

Remote sensing has been widely used to monitor the spatial patterns of drought and its related changes in global ecosystems. Previous studies implemented the vegetation indices (NDVI, NDWI) from Advanced Very High Resolution Radiometer (AVHRR) (Tucker, 1979) and MODIS to detect drought conditions over the Great Plains in the US (Gu et al., 2007), East Asia (Song et al., 2004), and Amazonian forest (Anderson et al., 2010). Due to the lagged response of vegetation to the drought events, these vegetation indices were combined with land surface thermal infrared information to improve drought detection (Singh et al., 2003; Park et al., 2004; Wan et al., 2004; Son et al., 2012). In this study, the negative anomalies of GPP in the Southeast only occur in S0, although both simulations captured the vegetation response to drought events in 2006, especially in the low biomass ecosystem (grassland, shrubland), by exploring the vegetation indices and thermal information. The spatial pattern of GPP anomalies in S1 was similar with Xiao et al. (2010), which also incorporated MODIS products using the regression tree model. Park et al. (2004) demonstrated that soil water-holding capacity affected the drought detection when using

MODIS vegetation indices and thermal information. Gu et al. (2008) also pointed out that the various correlations between MODIS, NDVI, NDWI and drought events were highly dependent on the vegetation structure and soil types. Drought stress with reduced soil moisture can constrain plant nutrient uptake by reducing nutrient supply through mineralization (Emmett et al., 2004; Sowerby et al., 2008; Larsen et al., 2011), also by reducing nutrient diffusion and mass flow in soils (Lambers et al., 2008). Therefore, the induced drought condition may potentially decouple the soil microbial carbon and soil CO₂ fluxes with canopy photosynthesis (Ruehr et al., 2009), increasing the residence of recent assimilated carbonate in vegetation (Deng et al., 1990; Li and Wang, 2003) and thus affecting the whole ecosystem carbon and nitrogen cycling. Some studies also pointed out the great contribution of drought to the surface CO₂ variability (Knorr et al., 2005; Keppel-Aleks et al., 2014). The climate of the Southeast is predicted to be warmer and drier in future (Dai, 2013). Therefore, incorporating the atmospheric CO₂ concentration into an upscaling practice using the flux site data should improve the qualification of GPP and NEE and strengthen our understanding of vegetation's response to environmental condition.

4.4. Implication of the CO₂ effect on GPP and NEE

Since atmospheric CO₂ is an important factor related to plant carbon assimilation and water balance, incorporating it is expected to improve our understanding on the ecosystems' response to environmental variables. However, previous studies concerning the degree of climate controls on the spatial and temporal patterns of the eddy covariance derived GPP and NEE (e.g. Yu et al., 2013) or upscale the site-level data to regional estimation (e.g. Xiao et al., 2010) always ignored the atmospheric CO₂ concentration. As our study statistically demonstrated above, including CO₂ as an explanatory variable in upscaling site-level eddy flux data to regional GPP and NEE made an obvious seasonal and spatial difference. Taking the North Central region of US as an example, our results demonstrated that GPP was about 200 g C m⁻² yr⁻¹ higher in S0 due to the overestimation in the growing season; this implied that larger GPP and NEE differences may occur between the two simulations when atmospheric CO₂ showed a greater variability during these periods. Furthermore, excluding the atmospheric CO₂

concentration, the regional GPP and NEE variation may be overly attributed to other climate variables.

Up to now, most of the regional and global estimations of terrestrial carbon uptake come from various ecosystem biogeochemistry models, which are mathematical representations of biophysical processes (Bonan, 2008). The model constrained with eddy flux towers observation would strengthen the understanding of ecosystem mechanisms. This data-assimilation method provides an alternative way to quantify regional carbon exchanges between the terrestrial biosphere and the atmosphere at regional and global scales (Xiao et al., 2012). Generally, flux measurements are used to calibrate the parameters in ecosystem models, and then the optimized parameters are extrapolated for the estimation over a region. In some process-based ecosystem models which use CO₂ data as an input to compute GPP, the carbon fluxes are routinely calibrated and simulated based on annual CO₂ time series (or to be more specific, the background CO₂ concentration) across a region or globe (Turner et al., 2003; Chen et al., 2011). However, although CO₂ is generally well-mixed globally since it is chemically inert (Eby et al., 2009), it may exhibit a large local variability at short time scales. A recent study by Keppel-Aleks et al. (2014) also underscored the need to consider the full spatial and temporal distribution of atmospheric CO₂ to accurately separate the climate events' (e.g., warming and drought) influence on carbon cycling. Thus, GPP and NEE estimation should be improved if spatially- and temporally-varied atmospheric CO₂ data are used.

4.5. Limitation

There are a number of limitations in our ANN estimations. First, the gridded CO₂ product was developed using a global inversion model, whose accuracy is highly dependent on the prior CO₂ fluxes (Masarie et al., 2011). In addition, inaccurate weather data in complex terrain also introduces errors by using the transport model (TM5) at coarse spatial resolutions (Geels et al., 2007). The uncertain CO₂ product may bias our GPP and NEE estimations. To assess the uncertainty of the NOAA CO₂ products, we did a comparison analysis between the 8-daily averaged NOAA gridded CO₂ values and the in-situ CO₂ observation in summer season at seven sites (Fig. S7). Generally, the extracted CO₂ seasonal variations from NOAA products were comparable with the CO₂ measurement at forests (Fig. S7a–S7c) and crop sites (Fig. S7g), with the differences ranging from 5 to 15 ppmv. The site-level sensitivity analysis (Fig. S8) showed that these CO₂ input data differences would change annual GPP by 8–50 gC (2–6% relative change) using ANN models. In particular, the GPP changes in the mixed forest and croplands were almost two times larger than in other ecosystems. Despite the uncertainty of the gridded CO₂ product, we used it to gain insight into the effects of CO₂ on regional GPP and NEE prediction when upscaling site-level data. High frequency, stable CO₂ measurements (Andrews et al., 2014) and high-density observation networks (Lauvaux et al., 2012) are needed to accurately quantify the atmospheric CO₂ effects on GPP.

Second, the selected AmeriFlux sites can contribute a large uncertainty to our model development, because ANN training and validating are highly associated with the eddy flux data. The representativeness of the AmeriFlux data is crucial to scaling up site-level measurement to large regions. In our study, the criterion to select sites is based on the available CO₂ measurement from 2000 to 2006. The representativeness of the flux sites is crucial to scaling up site-level measurement and to understand the carbon dynamics across large spatial scales and quantify uncertainty. Although the global sites well represent variations in climate and ecological regions (Reichstein et al., 2014), whether the major ecoregions from North America are well represented by the AmeriFlux network has not been examined (Hargrove et al., 2003). Using a cluster analysis with

25 climatic and physiological forcing variables based on 59 AmeriFlux sites, Hargrove et al. (2003) showed that the vast interior of the US was well-represented by the AmeriFlux network, but the Pacific Northwest, Southern, and Southwestern were under represented. In addition, the uncertain eddy flux data – due to system errors, random errors, site selection criteria (Loescher et al., 2006), and the gap-filling method (Moffat et al., 2007; Desai et al., 2008) – may affect the GPP and NEE uncertainty in the model parameterization and simulation.

Third, our approach has not considered other controlling factors on the carbon fluxes. The vegetation distribution was constant without any shifts or disturbances during the study period, which might have biased estimation. Lacking the effects of land-cover changes, natural disturbances (e.g., fire, hurricane, insect infestation), economic influences (e.g., harvest) and demographic factors (Lambin et al., 2003) would also introduce uncertainties to our quantifications. Especially in the US, about half of the forest area is disturbed each decade, including the timber harvesting and grazing (Birdsey and Lewis, 2003). The forest age reflects the past disturbance history. Use of the North American forest stand age database (Pan et al., 2011) derived from forest inventory, historical fire data, and remote sensing imagery could help reduce our quantification uncertainty (Deng et al., 2013; Xiao et al., 2014). For cropland, without considering the effects of crop rotation (such as from corn to soybeans) may also introduce uncertainty. Since C4 photosynthetic capacity is different from C3 plant, explicit C3 versus C4 plant distribution data shall also help reduce the uncertainty of carbon quantification. In addition, the MODIS ecosystem distribution is not consistent with flux tower footprints (Xiao et al., 2008) and can be significantly different from the towers' vegetation type. Recent studies by Melton and Arora (2013) showed that the plant function type variability within the one-grid scale is important in simulating ecosystem responses to changing climate and atmospheric CO₂ concentration.

Fourth, using only seven plant functional types (PFTs) derived from the MODIS land use product may also introduce uncertainties to the carbon flux quantification. In the simulations, we assumed that, at each PFT level, the response of carbon fluxes to environmental condition changes is same. The usefulness and accuracy of such assumption has been questioned by field ecologists and there is a significant overlap between PFTs (Reich et al., 2007). The PFT-level parameterization could be biased because it may not properly represent the functional and structural characteristics of the species within a region. The parameter values obtained at the PFT level led to diverse significant carbon dynamic quantifications (Alton, 2011; He et al., 2013). Additionally, Reichstein et al. (2014) showed that the functional biogeographical variations of ecosystem properties were only partially explained by climate and PFTs. The large variability of biogeochemical processes within each PFT (Kattge et al., 2011) depended even more on plant traits, such as allocation and stabilization of carbon in the soil. Future studies should explicitly include the site-level information and trait database to test the relation between ecosystem functions and organismic traits to accurately quantify carbon fluxes.

Finally, the simplistic aggregation of CO₂ hour by hour into 8-day averages may bias our quantification without considering the influence of the planetary boundary layer (PBL) development. The PBL depth and turbulence intensity have a strong impact on the vertical and horizontal distribution of CO₂ in the atmosphere (Yi et al., 2001). The covariance between surface fluxes of CO₂ and the atmospheric mixing process has a strong seasonal character (Helliiker et al., 2004). Since the rectifier effect influences the horizontal and vertical distributions of CO₂ in the atmosphere, it can bias carbon flux quantification (Denning et al., 1995). We further noted that we have not explicitly differentiated CO₂ fertilization effects and its indirect effects on CO₂ uptake through affecting stomata openness

in these ANN simulations. Process-based ecosystem models shall incorporate both fertilization and indirect effects into regional GPP and NEE quantification.

5. Conclusion

When scaling site-level eddy flux data to a region, the atmospheric CO₂ concentration on GPP and NEE has rarely been incorporated. Here we made a step forward by considering this effect in quantifying regional GPP and NEE in the conterminous US. We constructed two sets of artificial neural networks incorporating remote sensing variables to upscale the AmeriFlux site-level data to the region for seven ecosystem types: one that incorporated CO₂ and a second that did not. Our results showed that both sets of models showed good performance and were able to capture the GPP and NEE variation when compared with previous published results and MODIS products. However, the GPP and NEE difference between two models exhibited a great spatial and seasonal variability, which was closely related with climate drivers (air temperature and vapor pressure deficit). In addition, the simulation without considering CO₂ effects fails to detect the ecosystem response to droughts in the southeastern US in 2006, indicating that drought-induced surface CO₂ variation could have more constraint on the regional GPP and NEE qualification. This study is among the first to explore the CO₂ influence on the regional GPP and NEE estimation in upscaling eddy flux data to regional scales. Our study suggests that the spatially and temporally varied CO₂ concentration should be factored into GPP and NEE quantification when scaling eddy flux data to a region.

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Appendix A. Supplementary data

Supplementary material related to this article can be found, in the online version, at <http://dx.doi.org/10.1016/j.agrformet.2016.01.007>.

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