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Terrestrial biosphere models need better representation of vegetation phenology: results from the North American Carbon Program Site Synthesis

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Terrestrial biosphere models need better representation of vegetation phenology: results from the North American Carbon Program Site Synthesis

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Abstract

Phenology, by controlling the seasonal activity of vegetation on the land surface, plays a fundamental role in regulating photosynthesis and other ecosystem processes, as well as competitive interactions and feedbacks to the climate system. We conducted an analysis to evaluate the representation of phenology, and the associated seasonality of ecosystem-scale CO₂ exchange, in 14 models participating in the North American Carbon Program Site Synthesis. Model predictions were evaluated using long-term measurements (emphasizing the period 2000–2006) from 10 forested sites within the AmeriFlux and Fluxnet-Canada networks. In deciduous forests, almost all models consistently predicted that the growing season started earlier, and ended later, than was actually observed; biases of 2 weeks or more were

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typical. For these sites, most models were also unable to explain more than a small fraction of the observed interannual variability in phenological transition dates. Finally, for deciduous forests, misrepresentation of the seasonal cycle resulted in over-prediction of gross ecosystem photosynthesis by $+160 \pm 145$ g C m⁻² yr⁻¹ during the spring transition period and $+75 \pm 130$ g C m⁻² yr⁻¹ during the autumn transition period (13% and 8% annual productivity, respectively) compensating for the tendency of most models to under-predict the magnitude of peak summertime photosynthetic rates. Models did a better job of predicting the seasonality of CO₂ exchange for evergreen forests. These results highlight the need for improved understanding of the environmental controls on vegetation phenology and incorporation of this knowledge into better phenological models. Existing models are unlikely to predict future responses of phenology to climate change accurately and therefore will misrepresent the seasonality and interannual variability of key biosphere–atmosphere feedbacks and interactions in coupled global climate models.

Keywords: autumn senescence, carbon cycle, land surface model (LSM), leaf area index (LAI), model error, North American Carbon Program (NACP), phenology, seasonal dynamics, spring onset

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Introduction

Phenological transitions drive the seasonal progression of vegetation through stages of dormancy, active growth, and senescence. Although phenology has traditionally been concerned with physical changes in structure (e.g., leaf development and abscission), the inherent seasonality of mass and energy exchange between terrestrial ecosystems and the atmosphere can, more generally, be viewed as phenological in nature (Gu et al., 2003). In deciduous forests, the relationships between the phenology of canopy structure and function are obvious. In evergreen forests, physiological changes within existing foliage (and not the production of new foliage) regulate the annual rhythms of photosynthesis and transpiration (e.g., Monson et al., 2005; Richardson et al., 2009b). In both forest types, phenological switches, rather than fast responses to highfrequency variation in environmental drivers, are controlling the seasonal patterns.

Phenology is thus a key regulator of ecosystem processes and biosphere feedbacks to the climate system (Peñuelas et al., 2009). Phenology influences both spatial and temporal (at seasonal-to-interannual time scales) variability in ecosystem productivity (Baldocchi et al., 2001; Churkina et al., 2005; Richardson et al., 2009a, 2010; Dragoni et al., 2011), and it is of fundamental importance for ecosystem carbon cycling, terrestrial carbon sequestration, and mitigation of anthropogenic CO₂ emissions. Furthermore, phenology affects the following: hydrology (Hogg et al., 2000), as leaf-out is accompanied by an increase in evapotranspiration and reduced throughfall; nutrient cycling processes (Cooke & Weih, 2005), as senescence results in fresh litter inputs to the soil; and atmospheric and climate system feedbacks (Schwartz, 1992), as the amount and condition of foliage present affects albedo, surface energy balance, and surface roughness (Moore *et al.,* 1996; Sakai *et al.,* 1997; Peñuelas *et al.,* 2009).

It is, therefore, essential that terrestrial biosphere models simulating the temporal dynamics of biological processes on the land surface have an accurate representation of phenology. This is true whether the model is simple or complex (in terms of the number of biogeochemical processes it features, and the degree to which processes are coupled or interact with each other) and whether the model is being run for a single site or the entire globe. Indeed, Levis & Bonan (2004) highlight the importance of accurate prognostic modeling of phenology, and the associated seasonal patterns of canopy leaf area index (LAI), for climate model runs that couple a land surface scheme to an atmospheric general circulation model.

A number of previous studies have evaluated the phenology submodels included in state-of-the-art land surface schemes and ecosystem models and concluded that these routines tend to be overly simplistic and result in biased predictions (Kucharik et al., 2006; Ryu et al., 2008). Randerson et al. (2009) included phenological metrics as part of a systematic framework, the Carbon-LAnd Model intercomparison Project (C-LAMP), to assess the biogeochemical component of coupled climate-carbon models. They concluded that model bias toward under-predicting temperate and boreal forest uptake of CO₂ could be attributed to a 1–3 month delay in predicting the timing of maximum LAI in these ecosystems, compared to estimates derived from MODIS data. Randerson et al. (2009) also noted that the two models they evaluated tended to predict a longer growing season than was actually observed in temperate ecosystems, with photosynthetic uptake occurring too early in the spring and too late in the autumn, compared with ground observations. Errors in LAI would likely propagate to errors in partitioning the available energy to latent and sensible heat fluxes, and errors in

the timing of photosynthetic uptake would also affect the seasonality of modeled atmospheric CO_2 concentrations, emphasizing the importance of accurate representation of phenologically mediated processes.

In this study, we describe an analysis of the representation of phenology, in terms of the seasonality of LAI, gross ecosystem photosynthesis (GEP), and net ecosystem exchange (NEE), in 14 terrestrial biosphere models that contributed model runs to the North American Carbon Program (NACP) Site Synthesis. (We do not intend for this study to be considered a comprehensive analysis of all aspects of model performance; complementary NACP efforts include work by Schwalm et al. (2010) and Dietze et al. (2011), and work in preparation by K. Schaefer et al., T. Keenan et al., P. Stoy et al., and B. Raczka et al..) The five deciduous broadleaf forest (DBF) and five evergreen needleleaf forest (ENF) sites selected for the analysis are all members of either the AmeriFlux or Fluxnet-Canada networks. Our analysis draws on the continuous eddy covariance measurements of forest-atmosphere CO₂ fluxes that have been made at each site for the last decade or more. At the deciduous forest sites, the flux measurements are complemented by above- and below-canopy measurements of photosynthetically active radiation, with which the seasonal trajectory of LAI can be estimated (e.g., Turner et al., 2003).

The objectives of our analysis are as follows: (1) to assess the accuracy with which spring and autumn phenological transitions are predicted by different models; (2) to evaluate how these patterns vary between deciduous and evergreen forest types; and (3) to quantify how much of the total bias in modeling annual GEP can be attributed to errors in modeling the spring and autumn phenological transitions.

Data and method

Field measurements

The present analysis uses field measurements and model runs contributed to the NACP Site Synthesis project (http://nacp. ornl.gov/mast-dc/int_synthesis.shtml). We restrict our analysis to temperate and boreal deciduous broadleaf and evergreen needleleaf sites (five DBF and five ENF sites), with summer active/winter dormant seasonality, selected from the NACP 'Priority 1' list (Table 1).

Eddy covariance measurements of net ecosystem exchange of CO₂ (NEE; µmol CO₂ m⁻² s⁻¹) supplied by site investigators were gap filled and partitioned to GEP and ecosystem respiration according to Barr *et al.* (2004). The partitioning algorithm was compared with a variety of other approaches by Desai *et al.* (2008). Most partitioning methods were found to yield similar seasonal cycles, and estimates of annual GEP that were within 10% of each other. This gives us confidence **Table 1** Locations of deciduous broadleaf (DBF) and evergreen needleleaf (ENF) sites used in this analysis. Mean $(\pm 1 \text{ SD})$ spring and autumn transition dates (day of year), as extracted from measured data, are given for leaf area index (LAI) (20% seasonal amplitude of LAI; only reported for deciduous sites) and gross ecosystem photosynthesis (GEP)

| $(20\% \text{ GEP}_{m})$ | _{lax}). Further details are provided in t _i | text | | | | | | | | | |
|--------------------------|--|---------|---------|----------|-----------|------|-------------|-------------|--------------|-------------|----------------------------|
| | | | | | | | 20% LAI | | 20% GEP | | |
| Site | Name | Country | Lat. °N | Long. ∘W | Years | IGBP | Spring | Autumn | Spring | Autumn | Reference |
| Ca-Oas | SSA Old Aspen | Canada | 53.63 | 106.20 | 1997–2006 | DBF | 142 ± 9 | 279 ± 9 | 140 ± 8 | 266 ± 3 | Barr et al. (2007) |
| US-Ha1 | Harvard Forest (EMS Tower) | USA | 42.54 | 72.17 | 1992–2006 | DBF | 137 ± 6 | 308 ± 2 | 131 ± 6 | 287 ± 3 | Urbanski et al. (2007) |
| US-MMS | Morgan Monroe State Forest | NSA | 39.32 | 86.41 | 1999–2005 | DBF | 116 ± 3 | 304 ± 5 | 113 ± 2 | 290 ± 3 | Schmid et al. (2000) |
| US-UMB | Univ. of Mich. Biological Station | NSA | 45.56 | 84.71 | 1999–2006 | DBF | 142 ± 5 | 294 ± 8 | 135 ± 9 | 283 ± 5 | Gough <i>et al.</i> (2008) |
| US-WCr | Willow Creek | NSA | 45.81 | 90.08 | 2000–2005 | DBF | 138 ± 7 | 283 ± 5 | 138 ± 7 | 270 ± 7 | Cook et al. (2004) |
| Ca-Obs | SSA Old Black Spruce | Canada | 53.99 | 105.12 | 2000–2006 | ENF | | | 110 ± 11 | 288 ± 3 | Krishnan et al. (2008) |
| Ca-Ojp | SSA Old Jack Pine | Canada | 53.92 | 104.69 | 2000–2006 | ENF | | | 112 ± 10 | 289 ± 3 | Kljun et al. (2006) |
| Ca-Qfo | Chibougamau (Mature Forest) | Canada | 49.69 | 74.34 | 2004–2006 | ENF | | | 119 ± 12 | 297 ± 4 | Bergeron et al. (2007) |
| US-Ho1 | Howland Forest (Main Tower) | USA | 45.20 | 68.74 | 1996–2004 | ENF | | | 98 ± 7 | 308 ± 10 | Hollinger et al. (2004) |
| US-NR1 | Niwot Ridge | USA | 40.03 | 105.55 | 1999–2007 | ENF | | | 120 ± 8 | 298 ± 6 | Monson et al. (2002) |

in the partitioning at daily-to-annual time scales. Furthermore, we argue that the narrow moving window used by the Barr *et al.* algorithm makes it well suited to the seasonality analyses we conduct. Filtering of nocturnal NEE measurements during periods of inadequate turbulence was conducted on a site-by-site basis using a u^* change-point detection algorithm, described and evaluated by Barr *et al.* (2009; A. Barr, D.Y. Hollinger, A.D. Richardson, unpublished results).

For deciduous forests (the necessary data were generally not available for evergreen conifer sites), the seasonal trajectory of canopy leaf area index (LAI, m² m⁻²) was estimated as follows. First, we calculated the gap fraction, P, as $P = Q_t/Q_{ot}$ where Q_0 is incident solar photosynthetic photon flux density (PPFD) measured above the canopy and Q_t is the PPFD measured below the canopy. We used measurements of P when the solar zenith angle was closest to 57° (one sample in the morning and one sample in the afternoon) and then calculated LAI for each sample as $LAI = -\log(P)/K$ where K = G(57)/Kcos(57). We restricted our analyses to zenith angles nearest 57 degrees because at this point, all leaf inclination distribution functions (G) converge to 0.5. Then, we obtained daily LAI by averaging the two LAI values to consider foliar clumping effects (Rvu et al., 2010). Noise in the resulting time series, which we attribute mostly to cloud effects (variability in direct beam and diffuse PPFD), was smoothed with a spline function. We then re-scaled the seasonal trajectory of LAI so that the seasonal peak LAI derived in this manner matched with measured LAI as reported on the AmeriFlux web page (http://ameriflux.ornl.gov/) or in published manuscripts for each site (see references in Table 1), and the seasonal minimum LAI was zero.

With this approach, we obtained essentially continuous estimates of changes in LAI over time for each of the deciduous forest sites. However, the heterogeneous nature of the belowcanopy light environment raises questions about the degree to which these estimates may be representative of leaf area dynamics across the larger tower footprint. At four of the five deciduous sites, only a single below-canopy quantum sensor was used to measure Qt. At US-MMS, there were four belowcanopy sensors, which no doubt provided better sampling of spatial variability in Qt. At none of the sites were field campaigns to measure LAI (e.g., plant canopy analyzer or hemispherical photography) conducted at a sufficiently high temporal resolution (e.g., weekly) to permit accurate estimation of phenological transition dates. However, where such data are available at a lower temporal frequency (e.g., monthly), they provide a context for evaluating our LAI estimates. As shown in Fig. 1, the mean seasonal course (over multiple years) of LAI estimated from $P = Q_t/Q_0$ (solid black lines) is in good agreement with that obtained by the LAI-2000 (Li-Cor Biosciences, Lincoln, NE, USA) plant canopy analyzer instrument (open circles) across a network of plots at each site (no LAI-2000 data for US-WCr). For US-MMS and US-UMB, the timing of spring and autumn transitions was consistent between the two methods. For US-Ha1, there is a clear divergence in early autumn, with the LAI-2000 data indicating an earlier decline in leaf area. However, this could simply be an artifact of sampling in different years: although Qt data were available for 9 years and LAI-2000 for 4 years, there was only 1 year of overlap between the two data sets at US-Ha1. For Ca-Oas, LAI-2000 measurements were made in years with substantial variability in the timing of canopy development, and this variability (as shown in Fig. 1) masks the otherwise good concordance between LAI measured with the LAI-2000 and estimated from $P = Q_t/Q_o$. Both methods were in agreement, for example, on the exceptionally early springs in 1998 and 2001 and the late onset of leaf development in 1997 and 2004. The above patterns, and the overall strong correlation between LAI from the two methods at each site (r > 0.95 for Ca-Oas, US-MMS, and US-UMB; r = 0.85 for US-Ha1), give us confidence in our retrievals.

We conclude by noting that although satellite data (e.g., vegetation indices as well as more targeted products related to LAI and phenology) offer the promise of global coverage, they suffer from tradeoffs between spatial and temporal resolution and have their own substantial uncertainties (e.g., Zhang *et al.*, 2006; Garrigues *et al.*, 2008; White *et al.*, 2009), which in the context of the present analysis make them less suitable benchmarks for model evaluation.

Model runs

Participation in the NACP site synthesis was on a volunteer basis, and an open invitation was sent to the terrestrial biosphere modeling community. Modeling teams were free to choose as simple or as complex a model as they desired. Models were run on a site-by-site basis, using measured environmental drivers (gap filled as necessary using a standardized method, D. Ricciuto *et al.* in preparation), and site-specific initial conditions (as judged necessary), following a standard protocol (http://nacp.ornl.gov/mast-dc/docs/Site_Synthesis_Protocol_v7.pdf). The protocol specified spin-up of carbon pools to steady state, but the way in which this was implemented varied among modeling teams.

Of the variety of models for which output was submitted to the NACP database, we included only those that appeared to at least superficially capture the seasonal trajectory of ecosystem activity. For example, models that did not predict winter dormancy (i.e., if they instead predicted significant wintertime photosynthetic uptake) were not included in this analysis. The 14 models that were included in our analysis are listed in Table 2. Note that the LoTEC model was run in a data assimilation mode, and model parameters were optimized, on a siteby-site basis, conditional on the flux measurements. This no doubt contributes to the better performance of LoTEC compared with some of the other models in this analysis.

Model approaches to phenological variation

Critical phenological events influencing carbon uptake in deciduous forests relate to the timing of leaf appearance, leaf expansion (increase of LAI), and leaf loss. For evergreen forests, a similar classification of foliage activity may be made from winter dormant to fully active and then dormant again. The models analyzed here use a variety of methods to deter-



Fig. 1 The mean seasonal trajectory of observed and simulated leaf area index (LAI), for five deciduous broadleaf sites. The continuous observed LAI is derived from gap fraction estimates based on above- and below-canopy measurements of photosynthetically active radiation. Open circles show periodic LAI measurements made using an LAI-2000 plant canopy analyzer.

mine the amount of foliage present and active in a canopy (Table 2). The simplest approach is to prescribe a fixed seasonal course of LAI. This approach encompasses the onset and development of foliage and also the dynamics of leaf loss. The original version of SiB (Sellers *et al.*, 1986) used monthly LAI values that were specific for each plant functional type. A slightly more complicated approach to phenology is to prescribe the presence and amount of foliage based on remote sensing data. SiB2 (Sellers *et al.*, 1996) used AVHRR data to determine seasonal NDVI and then f_{PAR} and LAI. In the present analysis, BEPS relies on a global LAI dataset (Deng *et al.*, 2006) derived from SPOT4 VEGETATION images and

corrected for clumping via multi-angle POLDER observations (Chen *et al.*, 2005). ED2 is designed to operate with the MODIS LAI product or other phenological drivers (Medvigy *et al.*, 2009). Because satellite data are sometimes not available (e.g., for prognostic runs), models that use remotely sensed phenological observations may use multi-year average LAI and often maintain the flexibility of using other sources. The results presented here for the SiB class of models (as well as ISAM), for example, use a single average seasonal course of LAI determined for each site.

Foliage onset and development in plants have long been related to temperature thresholds and cumulative heat sums

Table 2 Summary of models used in this analysis and their representation of phenology and seasonality of leaf area index (LAI). For models with 'prognostic' phenology, the seasonality of LAI is predicted based on climatic drivers; for those with 'prescribed' phenology, an average seasonal LAI cycle, as derived on a site-by-site basis from satellite (AVHRR) data, was used. Models with semi-prescribed and semi-prognostic phenology represent a hybrid of these approaches. GDD is growing degree days; T is temperature; C is carbon; PFT is plant functional type

| Model name | Resolution | Leaf onset | Control on LAI | Leaf loss | Source |
|------------|-------------|----------------------------|-------------------------|---------------------------------|--------------------------------|
| BEPS | Daily | Satellite | Satellite | Satellite | Ju et al. (2006) |
| Biome-BGC | Daily | GDD and radiation sum | Dynamic C allocation | Daylength and low temperature | Thornton <i>et al.</i> (2002) |
| Can-IBIS | Half-hourly | T threshold | GDD and dynamic C | Prescribed | El Maayar <i>et al.</i> (2002) |
| CN-CLASS | Half-hourly | C balance | C balance | Daylength and low temperature | Arain <i>et al.</i> (2006) |
| DLEM | Daily | $T_{7-day} > threshold$ | GDD to PFT limit | Daylength and low temperature | Tian <i>et al.</i> (2010) |
| Ecosys | Hourly | Hours above T threshold | Dynamic C allocation | Hours below T threshold | Grant <i>et al.</i> (2009) |
| ED2 | Half-hourly | Semi-prescribed | Dynamic C allocation | GDD and leaf turnover | Medvigy et al. (2009) |
| ISAM | Half-hourly | Prescribed | Prescribed | Prescribed | Jain & Yang (2005) |
| LoTEC | Half-hourly | GDD | GDD | T-dependent turnover | Hanson <i>et al.</i> (2004) |
| LPJ_wsl | Daily | GDD | GDD | Leaf longevity (prescribed) | Sitch <i>et al.</i> (2003) |
| ORCHIDEE | Half-hourly | GDD and chilling | Dynamic C allocation | Decreasing T and T threshold | Krinner <i>et al.</i> (2005) |
| SiB3 | Half-hourly | Prescribed | Prescribed | Prescribed | Baker et al. (2008) |
| SiBCASA | 10 min | Prescribed | Prescribed | Prescribed | Schaefer et al. (2008) |
| SSiB2 | Half-hourly | Prescribed | Prescribed | Prescribed | Zhan <i>et al.</i> (2003) |

(see, e.g., Shelford, 1930). This kind of approach is used by a number of the models (Table 2). The Ecosys model requires soil surfaces temperatures to exceed specified thresholds depending upon latitude for a certain number of hours. Can-IBIS operates in a similar fashion based on air temperatures while for deciduous sites DLEM uses a 7-day moving average that must exceed a threshold (for evergreen sites, DLEM uses prescribed phenology). The heat sum or growing degree day (GDD) approach is used by several models (including Biome-BGC, ED2, LoTEC, LPJ wsl, and ORCHIDEE) for deciduous broadleaf trees but details of the implementations vary widely. Biome-BGC combines GDDs with a radiation sum (White et al., 1997). ORCHIDEE requires a chilling sum (Botta et al., 2000) prior to leaf initiation; other models implicitly account for chilling by summing degree days after a particular date such as January 1. LoTEC is distinct from the other models in that it optimizes parameter values to best fit the observed data. Included in the LoTEC optimizations are a leaf initiation GDD threshold and a full canopy GDD sum. These values are determined individually for each site and across the data record (the phenological parameters do not vary by year). Some of the models (Biome-BGC, ORCHIDEE) use a GDD approach for deciduous trees only; there is no explicit phenology (removal of dormancy) for evergreens.

A final approach, exhibited by CN-CLASS, is to prognostically calculate foliage carbon balance. In this method, leaf onset starts when daily photosynthesis of virtual leaves exceeds daily respiration for seven consecutive days (Arora & Boer, 2005). Virtual photosynthesis and respiration are both functions of temperature, and photosynthesis is also a function of soil moisture. Once leaf onset occurs, a number of models (Table 2) calculate LAI based on carbon allocation principles. These include seasonal allocation rules for different tissues and net foliage C gain.

Several different schemes are used for determining foliage inactivation or shedding. These include prescribed LAI (the SiB class models, BEPS, Can-IBIS, and ISAM), prognostic leaf longevity that varies according to GDD (LPJ_wsl), and various low temperature thresholds. Several of the models (Biome-BGC, CN-CLASS, DLEM) combine daylength and temperature thresholds based on the results of White *et al.* (1997). Ecosys requires a set number of hours below plant functional type-specific thresholds and shortening photoperiods to initiate litterfall in deciduous species and foliage inactivation in evergreens. In some models (e.g., ED2, ORCHIDEE), in addition to changes in LAI through the season, there are decreases in photosynthetic capacity driven by leaf aging. Leaf loss in LOTEC is determined by a low temperature parameter that is optimized for each site.

Data processing and extraction of phenological transition dates

For the deciduous sites, we extracted phenological transition dates from measured and modeled LAI trajectories. In addition, for both deciduous and evergreen sites, we included a series of diagnostic phenological metrics extracted from the measured and modeled time series of NEE and GEP. Transition dates and thresholds were estimated from smoothing splines fit to measured and modeled data at the daily time step, as illustrated in Richardson *et al.* (2010). The phenological transition dates we estimated from the data were as follows:

- 1 The first spring and last autumn dates at which measured and modeled LAI = 20%, 50%, and 80% of the seasonal LAI amplitude (deciduous sites only);
- 2 The first spring and last autumn dates at which estimated and modeled daily GEP = 20%, 50%, and 80% of the seasonal maximum GEP; and
- 3 The first spring and last autumn dates of NEE source/sink transition (restricted to source/sink transitions that began or concluded periods of 14 continuous days of net CO₂ uptake).

The relative thresholds (20%, 50%, and 80%) were selected to correspond to a range of developmental stages, so as to evaluate the ability of models to predict not only 'start' and 'end' of the growing season but also the overall seasonal pattern. We used relative rather than absolute (e.g., LAI = $1.0 \text{ m}^2 \text{ m}^{-2}$ or GEP = 2 g C m⁻² day⁻¹) measures to account for differences in magnitude of both leaf area and CO₂ fluxes across sites (see also Richardson *et al.*, 2010 for a similar approach; alternative methods have been proposed elsewhere, e.g., Gu *et al.*, 2003).

We calculated transition date anomalies on a site-by-site basis, i.e., as interannual departures from the site mean.

For transition dates, model bias was calculated as the difference, in days, between the modeled and the observed transition date. Thus, a negative bias indicates that the model predicted the transition date too early in the year, whereas a positive bias indicates the model predicted the transition date too late in the year.

To evaluate the impact of errors in modeling phenological transitions on annual ecosystem productivity estimates, we calculated several metrics, beginning with the total model bias in annual GEP:

Total model bias = (annual model GEP)
- (annual tower GEP).
$$(1)$$

Three main sources of model error (which may reflect some combination of errors in model structure and errors in model parameterization) that contribute to the total model bias in GEP are (1) errors in the overall magnitude of modeled GEP; (2) errors in the seasonality of modeled GEP; and (3) errors in the sensitivity of modeled GEP to high-frequency variability in environmental drivers (e.g., incorrect representation of GEP sensitivity to vapor pressure deficit, soil water stress, or air temperature, among other factors). Errors in GEP magnitude (1), are likely due to incorrect specification of photosynthetic parameters such as A_{max} or V_{cmax} and can occur independently of errors in (2) or (3). To correct for (1) and

emphasize instead a focus on errors in seasonality, we rescaled modeled GEP (daily values) so that the seasonal maxima (as determined by the 95th percentile value) of measured and observed GEP were the same. Thus,

model
$$\text{GEP}_{\text{re-scaled}} = \text{model GEP} \times \frac{\text{tower GEP}_{\text{max}}}{\text{model GEP}_{\text{max}}}.$$
 (2)

We then calculated the total model bias in annual GEP, corrected for differences in GEP_{max} , as follows:

Total model bias (scaled for GEP_{max}) = (annual model $GEP_{re-scaled}$) – (annual tower GEP). (3)

To quantify how much of the model bias could be attributed to errors in seasonality, we defined 'spring' as the period between the first date (for each site-year of data, for each model run) when either model GEP or tower GEP rose to 20% of $GEP_{max'}$ and the last date when either model GEP or tower GEP rose to 80% of GEPmax. 'Autumn' was similarly defined as the period between the first date when either model GEP or tower GEP dropped below 80% of $\text{GEP}_{\text{max}\prime}$ and the last date when either model GEP or tower GEP dropped below 20% of GEP_{max}. Put differently, 'spring' and 'autumn' model biases were calculated from model GEP_{re-scaled} and tower GEP, with GEP integrals calculated for the period during which either model GEP or tower GEP is in the 'increasing GEP' phase (0.2 \leq daily GEP/GEP_{max} \leq 0.8) in spring or 'decreasing GEP' phase in autumn (0.8 \geq daily GEP/ $\text{GEP}_{\text{max}} \ge 0.2$). (Thus, the first and last dates of 'spring' and 'autumn' varied from year-to-year and across sites, but in each instance, the same dates were used to define the period of integration for both model GEP and tower GEP. Note also that the periods of integration could thus vary among models.)

Results

Leaf area dynamics

The modeled seasonal trajectory of deciduous forest LAI suffered from errors in the timing of both spring increases and autumn decreases in LAI, as well as errors in the amplitude of the seasonal cycle. No single model characterized LAI dynamics well at all five sites (Fig. 1). Reasonable performance at one site did not guarantee good performance at other sites: compare, for example, CN-CLASS at Ca-Oas and US-WCr with the same model at US-MMS (autumn decrease in LAI predicted approximately 3 months early) or US-UMB (virtually no seasonality to LAI, perhaps indicating that the site may have been mistyped as evergreen, rather than deciduous, forest). For some models, LAI dynamics were poor at most or all sites (e.g., Can-IBIS, DLEM, LPJ wsl, SSiB2). For Can-IBIS, deciduous forest LAI was too high ($\geq 5 \text{ m}^2 \text{ m}^{-2}$) during the winter dormant

season and did not decline in autumn. Biome-BGC captured the beginning and end of the season but LAI increased too slowly over the season. LoTEC, which optimized leaf onset, development, and abscission at each site, reproduced leaf onset and LAI values well but was surprisingly poor at capturing autumn senescence.

Models generally initiated greening up of the canopy too early in spring (Fig. 2): across all years, sites, and models, the mean (±1 SD) bias in the date at which model LAI reached 20% of maximum LAI was -10 ± 12 days, with a range from -53 to +28 days. There was, however, considerable variation among models in the magnitude of this bias. For example, across all sites, bias in the date at which the 20% LAI threshold was reached were largest for LPJ wsl and BEPS $(-19 \pm 6 \text{ days})$ and smallest for Ecosys $(-5 \pm 6 \text{ days})$ and CN-CLASS (+3 ± 13 days). Can-IBIS was the only model commonly biased toward late predictions ($+7 \pm 10$ days) of the onset of spring increases in LAI (recall however that this model retained substantial leaf area through the dormant season).

By comparison, across all years, sites, and models, the mean bias in the autumn date at which model LAI dropped to 20% of maximum LAI was $+1 \pm 32$ days, with a range from -98 to +84 days. For this indicator, model predictions were generally biased early for ED2 (-5 ± 10 days), LoTEC (-13 ± 9 days), and CN-CLASS (-30 ± 32 days), but late for all the other models (particularly LPJ_wsl, $+38 \pm 28$ days). Predictions were consistently biased late for US-WCr, US-UMB, and Ca-Oas. At the other two deciduous sites, model performance was mixed (Fig. 2).

In spite of these errors in modeling the overall seasonal trajectory of LAI, most models were able to predict some of the interannual variability in phenology (that is, year-to-year phenological anomalies) correctly in spring, but not in autumn (Table 3). Thus, although model predictions of spring onset dates were biased overall, the models did correctly represent a significant fraction of the interannual variability in canopy development associated with 'early' vs. 'late' spring onset. However, even in the best cases, with correlation coefficients typically in the range of $r \approx 0.5$ -0.8, between 30% and 75% of the observed interannual variation remained unexplained by the models. For eight models, the correlation between anomalies in observed and modeled dates at which LAI reached 20% of the seasonal maximum was highly significant in spring (all P < 0.001; Table 3). However, for only one model, Ecosys (r = 0.42, P < 0.05), was there a significant correlation between the observed and modeled anomalies in the autumn date at which this same threshold was reached. Furthermore, although models could predict some of the variation in the spring dates at which LAI reached 20% and 50% of the seasonal maximum, they were much less successful at predicting spring dates at which LAI reached 80% of the seasonal maximum. This indicates deficiencies in model representation of rates of leaf growth and the sensitivity of leaf growth to interannual climate variability. Finally, the amount of interannual variability predicted by the models was highly variable; CN-CLASS and LoTEC generally predicted too much variability in both spring and autumn developmental threshold dates, whereas in the models with prescribed phenology, there was of course no interannual variability in the dates when different thresholds were reached.

Overall, then, models were generally inadequate in their representation of the timing, and interannual variability in the timing, of both spring green-up and autumn senescence of deciduous forest sites. Better input data could rectify this problem for the subset of models using prescribed LAI (Table 2), but in those with prognostic LAI routines, either model structure or model parameters need to be improved.

Start and end of photosynthetic activity

For deciduous sites, virtually every one of the 14 models included in this analysis predicted an earlier onset of photosynthetic activity (defined as the first date at which daily GEP = 20% of maximum daily GEP) than was indicated by the eddy covariance measurements (Fig. 2). Across all models, sites, and years, the mean bias in photosynthetic onset date was -28 ± 21 days, with a range from -108 to +19 days. Relatively small biases were observed for some models (Ecosys, -3 ± 6 days; LoTEC, -4 ± 9 days), but large biases, of more than 6 weeks, were typical for other models (Can-IBIS, -52 ± 17 days; CN-CLASS, -60 ± 22 days). For evergreen sites, the same pattern was apparent – with three minor exceptions (BEPS, $+7 \pm 8$ days, ISAM, $+4 \pm 7$ days, and LPJ_wsl, $+1 \pm 14$ days), predicted dates of the onset of photosynthesis were earlier than the observed dates - but the biases tended to be somewhat smaller than for deciduous forests (mean bias, all models, sites, and years, -11 ± 15 days). Larger errors were observed for ORCHIDEE (-18 ± 12 days), Biome-BGC (-25 ± 10 days), and ED2 (-29 ± 20 days).

In autumn, most models predicted that the photosynthetic activity of deciduous sites (here judged as the last date at which daily GEP = 20% of maximum daily GEP) persisted later than was actually observed (mean error, all models, sites, and years, $\pm 15 \pm 17$ days) (Fig. 2). Biases were again largest for Can-IBIS ($\pm 45 \pm 16$ days) and CN-CLASS ($\pm 32 \pm 23$ days); the



Fig. 2 Difference in number of days (*y* axis) between observed and modeled start (left panels) and end (right panels) of photosynthetically active period for five deciduous broadleaf and five evergreen needleleaf sites. The start and end of photosynthetic uptake are defined as the first and last dates at which daily gross ecosystem photosynthesis (GEP) = 20% of maximum daily GEP. For each of 14 models (*x* axis), bars indicate the mean bias, with error bars indicating the standard deviation across multiple years. Circles indicate bias in the spring and autumn dates at which 20% of seasonal amplitude of leaf area index (LAI) was reached (deciduous sites only). Negative values indicate that the modeled transition occurred *prior* to the observed transition.

smallest errors were observed for LoTEC ($+2 \pm 5$ days) and ED2 (-2 ± 13 days). By comparison, for evergreen sites there was not a strong bias one way or the other (mean bias, all models, sites, and years, $+3 \pm 14$ days) with respect to predicting the end of photosynthetic uptake. On average, some models were too early, and some models were too late, but for a given model and a

given site, even the sign of the bias could vary from year-to-year. However, ORCHIDEE (+18 \pm 12 days) and SSiB2 (+18 \pm 14 days) were notable exceptions to this general pattern, because both models predicted that photosynthetic uptake would continue, on average, for two and a half weeks longer than was actually observed at the evergreen sites.

| | | 1, | | | | |
|-----------|---------------|--------------|--------------|-------------|--------------|------------|
| | Spring LAI th | resholds | | Autumn LAI | thresholds | |
| Model | 20% | 50% | 80% | 80% | 50% | 20% |
| BEPS | 0.18 | 0.48 | 0.12 | -0.25 | -0.20 | 0.01 |
| Biome-BGC | 0.64^{***} | 0.64^{***} | 0.24 | 0.23 | 0.02 | -0.29 |
| Can-IBIS | 0.57^{***} | 0.46^{*} | -0.19 | | | |
| CN-CLASS | 0.69^{***} | 0.61^{***} | 0.40^{*} | -0.10 | -0.14 | 0.05 |
| DLEM | -0.41^{*} | -0.03 | -0.06 | -0.39^{*} | -0.11 | 0.02 |
| Ecosys | 0.71^{***} | 0.71^{***} | 0.66*** | 0.36 | 0.47^{**} | 0.42^{*} |
| ED2 | 0.86^{***} | 0.75^{***} | 0.41^{*} | 0.48^{**} | 0.59^{***} | 0.26 |
| LoTEC | 0.82*** | 0.92*** | 0.77^{***} | 0.02 | 0.09 | -0.06 |
| LPJ_wsl | 0.64^{***} | 0.79^{***} | 0.59^{***} | 0.19 | 0.00 | 0.20 |
| ORCHIDEE | 0.62*** | 0.58*** | 0.37* | 0.00 | -0.02 | -0.12 |

Table 3 Correlation coefficient between observations and model predictions of leaf area index (LAI) transition date anomalies (i.e., years with 'earlier' vs. 'later' spring). Anomalies were calculated on a site-by-site basis, across all years of data for each site (deciduous sites only). LAI transition dates were estimated based on dates at which specific relative thresholds of seasonal development. i.e., 20%, 50%, 80% of seasonal LAI amplitude, were reached

Note: SiB3, SiBCASA, SSiB2, and ISAM were excluded from this analysis because prescribed phenology did not vary from year-toyear. No autumn correlations are reported for Can-IBIS because predicted LAI did not decrease in autumn.

Asterisks denote statistical significance:

***P* < 0.01;

***P < 0.001.

For deciduous sites, only about half of the models could predict interannual variation in the timing of the spring onset of photosynthesis with any degree of success, and none of the models consistently captured observed variation in the end of photosynthetic uptake. Six models showed statistically significant correlation with observed anomalies in spring dates at which GEP = 20% of maximum GEP (Table 4, top). However, even for the two best models (ED2, r = 0.66; LoTEC, r = 0.67), these correlation coefficients, although statistically significant, indicate that models capture no more than 50% of the observed interannual variation in the onset of photosynthesis. For the 50% and 80% spring thresholds of maximum GEP, the number of correlations significant at P < 0.001 was even lower (three and two models, respectively). For evergreen sites, models (with the exception of ED2) were generally more successful at predicting interannual variation in the timing of the spring onset of photosynthesis (Table 4, bottom). Once more, however, the 80% threshold of maximum GEP was predicted less well than the 20% threshold. At the end of the growing season, most models, whether simulating deciduous or evergreen sites, were unable to explain more than a very small proportion of the interannual variation in the timing of autumn declines in GEP (Table 4).

For deciduous sites, errors in modeling the seasonal dynamics of GEP could largely be attributed to errors in modeling seasonal dynamics of LAI. For both spring (most models) and autumn (some models), biases in

modeled dates at which 20% thresholds of LAI and GEP were reached were strongly correlated (for each model, across all sites, and years) with each other (Fig. 2; Table 5). Thus, errors in modeling the beginning of canopy development, and the end of canopy senescence, typically translated directly to corresponding errors in modeling the timing of seasonal dynamics of GEP. Errors in modeling the dates at which 80% thresholds of LAI and GEP were reached were less strongly correlated, probably reflecting a decoupling between photosynthesis and leaf area in models once the canopy is more than half-full. There were, however, some obvious exceptions to these patterns (Fig. 2). For example, for Can-IBIS, the deciduous sites were modeled with a large LAI in winter, and no autumn decline in LAI. CN-CLASS retained varying amounts of leaf area through winter, and for some sites, GEP could increase in spring before any new foliage was formed. For LPJ_wsl, leaf area was retained in autumn much longer than photosynthesis was sustained, resulting in large biases for LAI, but not GEP, threshold dates.

Source/sink transition dates

As was the case with GEP, virtually every model predicted an earlier spring source/sink transition than was actually observed for the deciduous sites (Fig. 3; mean across all models, sites, and years, -32 ± 36 days). Biases of more than 6 weeks were typical for some models (Can-IBIS, -54 ± 42 days; SSiB2, -71 ± 27 days;

^{*}P < 0.05;

CN-CLASS, -90 ± 18 days), but biases were negligible for other models (e.g., LoTEC, -4 ± 7 days and Ecosys, -5 ± 9 days). Among deciduous sites, the largest biases (30 days or more) in modeling the spring source/sink transition data were observed for Ca-Oas, US-UMB, and US-Ha1.

Model bias in predicting spring source/sink transition dates was smaller for evergreen sites (mean across all models, sites, and years, -8 ± 33 days). Several models (ED2, -36 ± 28 days; SSiB2, -71 ± 25 days) predicted the transition more than a month early. However, other models (most notably Can-IBIS, $+17 \pm 32$)

were generally late in predicting the spring source/sink transition date for evergreen sites.

Across all models, sites, and years, autumn sink/ source transition dates were modeled somewhat better than spring source/sink transition dates for deciduous (mean bias, -1 ± 37 days) sites, but considerably worse for evergreen (+42 ± 58 days) sites. However, within each forest type, large variation was observed. For example, for deciduous sites, some models predicted excessively early autumn source/sink transition dates (SSiB2, -68 ± 35 days) while others were late (Can-IBIS, +45 ± 24 days). For evergreen sites, mean model

Table 4 Correlation between observed and modeled gross ecosystem photosynthesis (GEP) transition date anomalies (e.g., years with 'earlier' vs. 'later' transition). Transition dates were determined as the first (spring) and last (autumn) day at which daily GEP = 20%, 50%, or 80% of maximum daily GEP. Anomalies were calculated on a site-by-site basis, across all years of data for each site

| | Spring GEP thresholds | | | Autumn GEP thresholds | | |
|-------------------|-----------------------|--------------|--------------|-----------------------|------------|-------------|
| | 20% | 50% | 80% | 80% | 50% | 20% |
| Deciduous forests | | | | | | |
| BEPS | 0.45^{**} | 0.51^{**} | 0.63*** | -0.10 | 0.19 | 0.11 |
| Biome-BGC | 0.56^{***} | 0.36^{*} | 0.43^{*} | 0.19 | 0.26 | -0.16 |
| Can-IBIS | 0.20 | 0.56*** | 0.32 | 0.18 | 0.05 | -0.28 |
| CN-CLASS | 0.08 | 0.14 | 0.38^{*} | 0.21 | -0.15 | -0.20 |
| DLEM | 0.53^{***} | 0.27 | 0.23 | 0.47^{**} | 0.37^{*} | -0.21 |
| Ecosys | 0.61*** | 0.72*** | 0.53*** | 0.31 | 0.15 | 0.45^{**} |
| ED2 | 0.66*** | 0.40^{*} | 0.10 | 0.22 | -0.02 | 0.27 |
| ISAM | 0.23 | 0.06 | 0.42^{*} | 0.13 | -0.02 | -0.23 |
| LoTEC | 0.67^{***} | 0.77^{***} | 0.50^{**} | 0.00 | -0.09 | 0.28 |
| LPJ_wsl | 0.41^{**} | 0.28 | -0.01 | -0.06 | 0.03 | 0.32^{*} |
| ORCHIDEE | 0.51^{***} | 0.30 | 0.11 | 0.45^{**} | 0.37^{*} | 0.03 |
| SiB | -0.03 | 0.16 | 0.42^{**} | 0.33^{*} | 0.27 | -0.34^{*} |
| SiBCASA | -0.07 | 0.34^{*} | 0.13 | 0.33^{*} | 0.31^{*} | -0.29 |
| SSiB2 | 0.12 | -0.18 | 0.36^{*} | 0.17 | 0.31* | -0.29 |
| Evergreen forests | | | | | | |
| BEPS | 0.71** | 0.37 | 0.55^{*} | 0.34 | 0.48 | 0.32 |
| Biome-BGC | 0.47^{**} | 0.67*** | 0.57*** | 0.32 | 0.38^{*} | 0.24 |
| Can-IBIS | 0.75^{***} | 0.63*** | 0.26 | 0.59^{***} | 0.09 | 0.42^{*} |
| CN-CLASS | 0.65^{***} | 0.71^{***} | 0.02 | 0.37^{*} | 0.29 | 0.38^{*} |
| DLEM | 0.88^{***} | 0.56*** | 0.05 | 0.18 | 0.14 | 0.38^{*} |
| Ecosys | 0.51^{**} | 0.58^{***} | 0.49^{**} | 0.12 | 0.27 | 0.41^{*} |
| ED2 | 0.21 | 0.49^{**} | 0.12 | 0.36^{*} | -0.26 | 0.18 |
| ISAM | 0.80^{***} | 0.56^{**} | -0.05 | 0.45^{*} | 0.21 | 0.27 |
| LoTEC | 0.67^{**} | 0.69** | 0.49 | -0.04 | 0.35 | 0.42 |
| LPJ_wsl | 0.64^{***} | 0.66*** | 0.43** | 0.20 | 0.01 | 0.31 |
| ORCHIDEE | 0.59^{***} | 0.46^{**} | 0.65^{***} | 0.45^{**} | 0.40^{*} | 0.18 |
| SIB | 0.57^{***} | 0.53^{***} | 0.37^{*} | 0.47^{**} | 0.30 | 0.29 |
| SiBCASA | 0.45^{**} | 0.61*** | 0.16 | 0.41^{*} | 0.32 | 0.34^{*} |
| SSiB2 | 0.65^{***} | 0.64^{***} | -0.09 | -0.06 | 0.24 | 0.37^{*} |

Asterisks denote statistical significance: *P < 0.05;

**P < 0.01;

***P < 0.001.

bias in predicting autumn sink/source transition dates was relatively small (although somewhat variable) for some sites (Ca-Qfo, $+5 \pm 23$ days) but very large for other sites (Ca-Ojp, $+80 \pm 37$ days; US-NR1, $+80 \pm 52$ days). Both ED2 ($+87 \pm 38$ days) and Can-IBIS ($+72 \pm 59$ days) models predicted autumn source/sink transition dates for evergreen sites that were more than 2 months later than actually observed.

Although many models predicted a statistically significant proportion of the interannual variation in the first date at which daily GEP = 20% of maximum daily GEP (Table 4), only a handful of models could predict the interannual variation in spring source/sink transition dates (Table 6). Of those, only Biome-BGC and LoTEC could explain at least half of the observed variability in spring source/sink transition dates for both deciduous and evergreen sites. Model skill in predicting autumn sink/source transition date anomalies was consistently poor (Table 6).

Just as errors in modeling the seasonal cycle of LAI explained much of the error in modeling the seasonal cycle of GEP, errors in modeling the seasonal cycle of GEP explained, for deciduous sites in both spring and autumn, a sizable fraction of the error in modeling the observed NEE source/sink and sink/source transitions (Table 6). Only for BEPS and LPJ_wsl were errors in predicting NEE transition dates not significantly correlated with errors in predicting GEP transition dates at deciduous sites. But, for evergreen sites, errors in modeling GEP transition dates generally did not explain the errors in modeling NEE transition dates transition dates generally did not explain the errors in modeling NEE transition transition the errors in modeling NEE transition dates generally did not explain the errors in modeling NEE transition transition transition the errors in modeling NEE transition dates generally did not explain the errors in modeling NEE transition transition the errors in modeling NEE transition transition transition the errors in modeling NEE transition transition transition transition transition the errors in modeling NEE transition transition the errors in modeling NEE transition transition transition the errors in modeling NEE transition transi

sition dates, presumably indicating that interannual variability in the seasonality of ecosystem respiration may contribute substantially to variability in NEE source/sink and sink/source transition dates in this forest type.

Errors in GEP integrals from incorrect representation of seasonality

The total bias in modeled annual GEP was +35 \pm 365 g C m^{-2} yr^{-1} for deciduous forests and +70 \pm 335 g C m⁻² yr⁻¹ for evergreen forests (mean \pm 1 SD across all sites, models, and years; for reference, mean annual GEP was 1250 ± 200 g C m⁻² yr⁻¹ in deciduous forests and 950 \pm 375 g C m⁻² yr⁻¹ in evergreen forests). By comparison, the total bias in modeled annual GEP, after correcting for model bias in GEP_{max} , was +260 \pm 250 for deciduous forests and \pm 55 \pm 130 g C m⁻² yr⁻¹ for evergreen forests. Thus, for deciduous sites, re-scaling GEP generally increased the total model bias in annual GEP, indicating that biases in GEP_{max} were effectively compensating for other model deficiencies. As an example, ED2 consistently under-estimated annual GEP in deciduous sites because model GEP_{max} was much smaller than the observed GEP_{max}. However, when modeled daily GEP was re-scaled to account for differences in GEPmax, the model typically over-estimated annual GEP because the model predicted a longer growing season than was actually observed (Figs 2 and 4).

Biases in annual simulated GEP were driven nearly equally by misrepresenting the timing of spring and fall

| | Errors in spi | ring threshold | | Errors in aut | umn thresholds | |
|-----------|---------------|----------------|-------------|---------------|----------------|--------------|
| Model | 20% | 50% | 80% | 80% | 50% | 20% |
| BEPS | 0.31 | -0.06 | 0.34 | 0.65^{*} | 0.22 | 0.82*** |
| Biome-BGC | 0.87^{***} | 0.38 | 0.23 | 0.00 | 0.24 | 0.50^{*} |
| Can-IBIS | 0.67^{***} | 0.36 | -0.05 | | | |
| CN-CLASS | 0.38^{*} | 0.55^{**} | 0.50^{**} | 0.38^{*} | 0.25 | -0.25 |
| DLEM | 0.54^{**} | 0.66**** | 0.57^{**} | 0.05 | 0.33 | 0.44^{*} |
| Ecosys | 0.84^{***} | 0.87^{***} | 0.31 | 0.23 | 0.43^{*} | 0.66*** |
| ED2 | 0.77*** | 0.50^{**} | 0.51** | 0.23 | 0.36 | -0.02 |
| ISAM | 0.27 | 0.39^{*} | -0.04 | -0.02 | -0.06 | 0.14 |
| LoTEC | 0.94*** | 0.82*** | 0.03 | 0.59^{**} | 0.00 | 0.74^{***} |
| LPJ_wsl | 0.56^{**} | 0.31 | 0.42^{*} | 0.00 | -0.08 | 0.10 |
| ORCHIDEE | 0.85^{***} | 0.55^{**} | 0.46^{**} | 0.22 | 0.25 | 0.69*** |
| SiB3 | 0.49^{**} | 0.37^{*} | 0.07 | 0.25 | 0.50^{**} | 0.50^{**} |
| SiBCASA | 0.41 | 0.18 | -0.25 | 0.08 | 0.57^{**} | 0.53^{*} |
| SSiB2 | 0.73*** | 0.35 | 0.06 | 0.26 | 0.55^{**} | 0.53** |

Table 5 Correlation between errors in modeled dates at which gross ecosystem photosynthesis (GEP) and leaf area index (LAI)thresholds (20%, 50%, 80% of seasonal maximum) were reached, across all deciduous sites

Asterisks denote statistical significance: *P < 0.05;

**P < 0.01;

***P < 0.001.

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Fig. 3 Difference in number of days (*y* axis) between observed and modeled start (left panels) and end (right panels) of carbon uptake period for five deciduous broadleaf and five evergreen needleleaf forest sites. The start and end of the carbon uptake period are defined as the first spring date and last autumn date, respectively, on which daily net ecosystem exchange of CO_2 (NEE) crossed from a source to a sink, or vice versa. For 13 of 14 models (*x* axis), bars indicate the mean bias, with error bars indicating the standard deviation across multiple years. Negative values indicate that the modeled transition occurred *prior* to the observed transition. No results shown for ISAM, which did not provide NEE output.

transitions and by biases in simulated GEP_{max} . We showed (Fig. 2) that for most deciduous sites, models generally predicted an earlier spring onset (first date at which GEP = 20% of GEP_{max}) of GEP than was actually observed, and a later autumn termination (last date at which GEP = 20% of GEP_{max}) of GEP than was actually observed. Thus, not surprisingly, models overestimated GEP during both the spring and autumn transition periods (Fig. 4). What is perhaps surprising is the magnitude of this bias; across all sites, models,

and years, the mean (\pm 1 SD) total model bias (scaled for GEP_{max}) in deciduous forests was +160 \pm 145 g C m⁻² yr⁻¹ in spring and +75 \pm 130 g C m⁻² yr⁻¹ in autumn (Fig. 4). Indeed, together (+235 \pm 230 g C m⁻² yr⁻¹) these two biases essentially offset the model error that could be attributed to differences in modeled vs. observed GEP_{max} (-225 \pm 440 g C m⁻² yr⁻¹), and accounted for virtually all of the model bias that remained after correcting for differences in GEP_{max} (+260 \pm 250 g C m⁻² yr⁻¹) for deciduous sites.

Table 6 (Left columns) Correlation between observed and modeled NEE source/sink transition date anomalies (e.g., years with 'earlier' vs. 'later' transition). Transition dates were determined as the first (spring) and last (autumn) day at which NEE source/ sink transition occurred. Anomalies were calculated on a site-by-site basis, across all years of data for each site. (Right columns) Correlation between errors in modeled dates at which daily GEP = 20% maximum GEP and NEE source/sink transition dates, in spring and autumn. NEE is net ecosystem exchange of CO₂, and GEP is gross ecosystem photosynthesis

| | NEE source/sink transition | | Errors in GEP and | NEE transitions |
|-------------------|----------------------------|------------|-------------------|-----------------|
| | Spring | Autumn | Spring | Autumn |
| Deciduous forests | | | | |
| BEPS | -0.27 | -0.06 | 0.20 | -0.16 |
| Biome-BGC | 0.70^{***} | 0.34 | 0.59** | 0.83^{***} |
| Can-IBIS | 0.11 | 0.42** | 0.46** | 0.60^{***} |
| CN-CLASS | 0.24 | -0.07 | 0.58*** | 0.24 |
| DLEM | -0.12 | 0.04 | 0.29 | 0.67^{***} |
| Ecosys | 0.53*** | 0.38^{*} | 0.79*** | 0.64^{***} |
| ED2 | 0.77*** | 0.00 | 0.76** | 0.34 |
| LoTEC | 0.71^{***} | 0.08 | 0.83*** | 0.77^{***} |
| LPJ_wsl | 0.24 | 0.06 | 0.24 | 0.28 |
| ORCHIDEE | 0.25 | 0.90*** | 0.73*** | 0.47^{**} |
| SiB3 | -0.01 | -0.19 | 0.33* | 0.47^{**} |
| SiBCASA | 0.14 | 0.19 | 0.38^{*} | 0.36^{*} |
| SSiB2 | -0.16 | -0.01 | 0.57*** | 0.46^{**} |
| Evergreen forests | | | | |
| BEPS | 0.38 | -0.07 | 0.03 | 0.31 |
| Biome-BGC | 0.79*** | 0.37 | -0.10 | -0.11 |
| Can-IBIS | -0.03 | -0.28 | 0.51** | -0.01 |
| CN-CLASS | 0.60**** | 0.16 | -0.10 | 0.08 |
| DLEM | 0.35 | 0.09 | 0.19 | -0.26 |
| Ecosys | 0.45^{*} | -0.24 | 0.95*** | 0.52^{**} |
| ED2 | 0.07 | 0.28 | 0.88*** | -0.05 |
| LoTEC | 0.76^{***} | 0.15 | 0.46 | 0.76^{***} |
| LPJ_wsl | -0.36 | 0.04 | 0.07 | -0.06 |
| ORCHIDEE | 0.85^{***} | 0.40^{*} | 0.54^{**} | -0.10 |
| SiB3 | 0.23 | 0.03 | 0.22 | 0.26 |
| SiBCASA | 0.21 | -0.16 | 0.26 | 0.25 |
| SSiB2 | -0.26 | 0.22 | 0.46 | -0.18 |

Note: ISAM was excluded from this analysis because model NEE was not provided.

Asterisks denote statistical significance:

*P < 0.05;

**P < 0.01;

***P < 0.001.

Biases in model predictions of spring and autumn GEP transition dates were generally much smaller for evergreen sites than for deciduous sites (Fig. 3). As a result, model biases in GEP during seasonal transition periods (after re-scaling to account for differences in GEP_{max}) tended to be smaller than those for deciduous sites (Fig. 4): +40 \pm 80 and -5 \pm 65 g C m⁻² yr⁻¹ in spring and autumn, respectively. And, whereas for deciduous sites, spring (≈20% of all sites, models, and years) and autumn (≈10% of all sites, models, and years) GEP errors of \geq 250 g C m⁻² yr⁻¹ were common (Fig. 4), they were rare to nonexistent ($\approx 1\%$ and 0%, respectively) for everyreen sites.

Discussion

Overview and relevance to modeling climate system feedbacks

The above analysis of predictions from 14 different terrestrial biosphere models has identified four key weaknesses in the representation of phenology and phenologically mediated processes:

- 1 Most models are biased toward predicting a growing season that is substantially longer than the observed growing season for deciduous forest sites, but biases are smaller and less consistent for evergreen forest sites.
- 2 Models are typically unable to capture more than a small fraction (albeit, in more than a few cases, a statistically significant fraction) of the observed interannual variability in phenological transition dates. This problem is more pronounced for deciduous forest sites than evergreen forest sites and more pronounced in autumn than in spring.
- 3 For deciduous sites, errors in modeling the seasonality of LAI often appear to propagate to errors in modeling the seasonality of GEP. This, in turn, leads to errors in modeling the seasonality of NEE.
- 4 Accumulated biases in GEP during spring and autumn transition periods, attributed to misrepresentation of the seasonality of GEP, are large and highly variable for deciduous sites: $+160 \pm 145$ and $+75 \pm$ 130 g C m⁻² yr⁻¹ ($\approx 13\%$ and $\approx 8\%$ of total annual GEP), respectively. These tend to offset errors associated with under-estimation of the magnitude of the seasonal peak GEP in deciduous sites. Thus, compensating errors may lead to erroneous conclusions about model performance at the annual time step.

From the perspective of global change science, the results presented here are important because (1) phenology is sensitive to climate change and variability; and (2) phenology controls many vegetation feedbacks to the climate system (Morisette et al., 2009). Analyses of diverse data sets provide compelling evidence for phenological shifts toward earlier spring onset and delayed autumn senescence over the last four decades (Peñuelas et al., 2002; Badeck et al., 2004; Schwartz et al., 2006; Parmesan, 2007; Parry et al., 2007). These patterns have largely been attributed to climate change, particularly recent warming trends. However, this analysis suggests that current models are unable to portray adequately the seasonality of either LAI or processes related to ecosystem carbon cycling under present climate scenarios. The analysis by Desai (2010) showed that accurate representation of interannual variability in phenology is important if the corresponding variability in net uptake of CO_2 is to be predicted correctly. We expect that most models (especially those in which phenology is prescribed) will not accurately predict the associated phenological responses to future climate change and variability either, which limits the usefulness of these models for prognostic studies. As will be discussed below, this is an outstanding challenge for phenological modeling in general.

In terms of the second point, our analysis was limited to the seasonality of LAI and ecosystem-atmosphere fluxes of CO₂. In all likelihood, however, these and similar models would also misrepresent other key feedbacks of terrestrial vegetation to the climate system during spring and autumn transition periods, e.g., through changing albedo, surface energy balance adjustment, and the changing partitioning of available energy to latent and sensible heat fluxes. This is of great importance because in addition to an influence on microclimate (e.g., ambient surface temperature, humidity, and radiative transfer through the canopy), phenology has effects on the planetary boundary layer, regional-to-global circulation patterns, and thus continental-scale climatic patterns (Hayden, 1998; Pielke et al., 1998; Chapin et al., 2000; Hogg et al., 2000; Fitzjarrald et al., 2001). Failure to represent phenology accurately in models that couple the land surface to the atmosphere could lead to large errors in the seasonal evolution of regional weather patterns, for example. The study by Levis & Bonan (2004) demonstrated on a regional scale that when phenology was prescribed, model runs using the Community Land Model coupled to the Community Atmosphere Model could not replicate observations that document a reduction in the rate of increase in surface air temperature that occurs coincident with spring leaf emergence and associated increases in transpiration. By comparison, when a prognostic phenology scheme was implemented, the important coupling between biological processes on the land surface and feedbacks to the atmosphere was restored, thereby improving model performance for this diagnostic. Thus, accurate model representation of phenology is critical because of the multitude of climate system feedbacks that are mediated by phenology.

Improving models

What steps are needed to improve phenological submodels in terrestrial biosphere models? For deciduous sites, large biases in predicting the start and end of the growing season need to be resolved, but models also need to do a better job of reproducing the interannual variability in phenology as well. Undoubtedly, progress requires better understanding of the controls on vegetation phenology, and the phenology of ecosystem processes, in different biomes and across plant functional types. For example, although the phenology of temperate, deciduous forests is well studied, there is remarkably little agreement regarding the degree to which photoperiod, cold temperatures, and warm temperatures combine to regulate spring budburst in these ecosystems (Chuine et al., 2010; Körner & Basler, 2010). Consequently, numerous models to predict budburst



Fig. 4 Bias in modeled gross ecosystem photosynthesis (GEP) for deciduous broadleaf (top) and evergreen needleleaf (bottom) forests. Left panels show bias, by model (means and standard deviations across multiple years of data for n = 5 sites), as follows: 'Total model bias' is (annual model GEP) – (annual tower GEP); 'Total model bias (scaled for GEP_{max})' re-scales the modeled GEP to account for order-of-magnitude differences between model and tower GEP_{max} and then calculates the bias as (annual model GEP_{re-scaled}) – (annual tower GEP); 'spring' and 'autumn' model biases are calculated from model GEP_{re-scaled} and tower GEP, with sums calculated for the period during which either model GEP or tower GEP is in the 'increasing GEP' phase ($0.2 \le daily GEP/GEP_{max} \le 0.8$) in spring or 'decreasing GEP' phase in autumn ($0.8 \ge daily GEP/GEP_{max} \ge 0.2$). The right panels show the frequency distribution of these spring and autumn biases in re-scaled model GEP, across all models, sites, and years of data, for each forest type: models show a strong and consistent bias in both spring and autumn in deciduous, but not evergreen, forests. The sign convention is that positive bias means that modeled GEP > tower GEP.

have been described in the literature (e.g., Chuine, 2000; Hänninen & Kramer, 2007; Richardson & O'Keefe, 2009), but there is no consensus on which model works best across species or across geographically distinct populations of a given species.

Forecasts of budburst dates for future climate scenarios are highly uncertain because the predicted response to warming depends strongly on the underlying model structure. Failure to incorporate photoperiodic control and chilling requirements, as is the case for most of the phenology submodels in the 14 models analyzed here, will likely result in over-estimation of the response to future warming (Körner & Basler, 2010). Even ignoring the impact that such errors would have on other aspects of biogeochemical cycling and climate system feedbacks, it is obvious that such biases would only further exaggerate the patterns for LAI and GEP reported here.

In general, neither prescribed nor prognostic schemes did well in simulating site-level phenology and its impact on LAI and GEP. However, this general pattern obscures some notable differences in model structure and performance, especially as related to the representation of environmental controls on photosynthesis and why the models tend to do better at evergreen sites than deciduous sites. Importantly, the LAI at evergreen sites stays nearly constant with time whereas the deciduous sites lose (nearly) all their leaves each winter. The timing of spring uptake at evergreen sites is controlled primarily by temperature, whereas at deciduous sites spring uptake is controlled by both temperature and the production of new foliage. Correspondingly, there are a variety of prognostic schemes for each of several aspects of phenology; (1) the start of leaf onset for deciduous species and removal of dormancy for evergreens, (2) the progress to full LAI in deciduous species

and full photosynthetic capacity in evergreens, and (3) the onset and schedule of leaf loss in deciduous species and entry into dormancy for cold winter evergreens.

Models using GDD sums which explicitly or implicitly included a chilling requirement did relatively well in capturing the onset of LAI and GEP for deciduous and evergreen forests. Models that used the GDD approach for deciduous but not evergreen forests (Biome-BGC, ORCHIDEE) consequently did relatively better for the deciduous type. LoTEC demonstrated that optimization of GDD parameters to multi-year mean values improved the model's ability to capture interannual variability in spring LAI. However, optimized phenological parameters were site specific, giving this approach limited power in long-term climate change simulations.

For autumn leaf loss in deciduous species and photosynthetic deactivation in evergreens, temperature thresholds combined with a shorter photoperiod have some predictive utility although there was a range of success in the models employing this approach (Ecosys, Biome-BGC, CN-CLASS, DLEM). In general, most of the variance in site-level autumn decreases in LAI, GEP, and NEE were unaccounted for by the models. Historically, there has been much less of an emphasis on developing models for autumn phenology, but the analysis presented here illustrates the need for new efforts in this direction.

Data needs

Increasing recognition of the importance of phenology should motivate progress toward disentangling the mechanisms and environmental drivers of annual phenological cycles. As a starting point, Stöckli et al. (2008) and Randerson et al. (2009) have emphasized the importance of developing better data sets with which to test and evaluate model predictions. Comparative analyses of different phenological models have typically used data from only a single site (e.g., Richardson & O'Keefe, 2009), which hinders the development and parameterization of generalized models. Only in a very few studies (e.g., Schaber & Badeck, 2003) have attempts been made to constrain phenological models using data across a wide geographic range. Thus, results from most analyses reflect over-fitting of models to individuals from a particular population, when in reality there may be genetic variation across the native range of a species with respect to the phenological sensitivity to climatic drivers (Chuine et al., 1998). An additional drawback is that single-site, short-term observational data often do not span sufficiently wide ranges of environmental or climatic conditions to falsify model predictions and thus distinguish among competing model structures (Hänninen, 1995). Satellite data offer the promise of global coverage, but are hindered by issues of both spatial and temporal resolution. Longterm, spatially extensive ground observations (ideally characterizing the entire seasonal trajectory of canopy development and senescence) are therefore urgently needed to elucidate environmental controls on phenology and improve phenological models. Ecosystem-scale modeling provides an additional challenge, in that it requires methods for scaling up from the phenology of individual species to correctly represent the aggregate phenology of mixed-species stands.

More than a decade ago, Baldocchi et al. (1996) recognized the value of eddy covariance time series of CO₂ and H₂O exchanges for evaluating and improving model representation of seasonal vegetation dynamics - i.e., phenology, and its role in regulating ecosystem processes related to carbon and water cycling. Indeed, this has motivated the present analysis. Beyond the NACP Site Synthesis, there are opportunities for related analyses that are even broader in scope. With close to a thousand site-years of measurements, from ecosystems spanning much of the globe's climate and vegetation space, the FLUXNET 'La Thuile' database (http:// www.fluxdata.org) is a virtual goldmine for the earth system modeling community (e.g., Williams et al., 2009). Also offering promise for improving phenological models are ground observations from continentalscale monitoring networks. For example, citizen science efforts, such as the USA National Phenology Network (http://www.usanpn.org) or webcam-based efforts such as PhenoCam (http://phenocam.sr.unh.edu; see Richardson et al., 2007, 2009b), could potentially yield spatially extensive data on the phenology of key plant functional types. Phenological observations from multifactor, manipulative global change experiments (e.g., Cleland et al., 2006) would be valuable for constraining model predictions under novel climatic or environmental conditions. Combining these diverse observations within a model-data fusion framework (as described in Williams et al., 2009), and in conjunction with objective model selection criteria (as previously applied to phenology models by Richardson & O'Keefe, 2009), it should be possible to develop and parameterize new phenological models and make substantial progress toward reducing biases and uncertainties of the type that have been documented here.

Conclusion

This analysis has shown that errors in simulating phenology and the seasonality of GEP result in large biases in modeling the productivity of deciduous broadleaf forests, whereas model performance was better for evergreen forests. Improving deciduous forest phenological models, particularly the controls on the seasonality of LAI and relationships among LAI, canopy photosynthesis and environmental drivers, should therefore be seen as a priority for the terrestrial biosphere modeling community. This is a prerequisite to better forecasts of vegetation responses to climate change and variability and is also essential for reducing errors in model representation of many biosphere– atmosphere interactions and climate system feedbacks.

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