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#### **Key Points:**

- Compensation day (cDOY) is the day of year when net C losses during winter are compensated by net C uptake in spring
- cDOY largely explains annual net ecosystem productivity NEP<sub>c</sub> of forests when the site has a distinct winter respiratory loss period
- cDOY and its explanatory power depends on the integration method for annual NEP<sub>c</sub> and on the forest type

Supporting Information:

Supporting Information S1

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# Winter respiratory C losses provide explanatory power for net ecosystem productivity

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Abstract Accurate predictions of net ecosystem productivity (NEP<sub>c</sub>) of forest ecosystems are essential for climate change decisions and requirements in the context of national forest growth and greenhouse gas inventories. However, drivers and underlying mechanisms determining NEP<sub>c</sub> (e.g., climate and nutrients) are not entirely understood yet, particularly when considering the influence of past periods. Here we explored the explanatory power of the compensation day (cDOY)—defined as the day of year when winter net carbon losses are compensated by spring assimilation—for NEP<sub>c</sub> in 26 forests in Europe, North America, and Australia, using different NEP<sub>c</sub> integration methods. We found cDOY to be a particularly powerful predictor for NEP<sub>c</sub> of temperate evergreen needleleaf forests ( $R^2 = 0.58$ ) and deciduous broadleaf forests ( $R^2 = 0.68$ ). In general, the latest cDOY correlated with the lowest NEP<sub>c</sub>. The explanatory power of cDOY depended on the integration method for NEP<sub>c</sub>, forest type, and whether the site had a distinct winter net respiratory carbon loss or not. The integration methods starting in autumn led to better predictions of NEP<sub>c</sub> from cDOY then the classical calendar method starting 1 January. Limited explanatory power of cDOY for NEP<sub>c</sub> was found for warmer sites with no distinct winter respiratory loss period. Our findings highlight the importance of the influence of winter processes and the delayed responses of previous seasons' climatic conditions on current year's NEP<sub>c</sub>. Such carry-over effects may contain information from climatic conditions, carbon storage levels, and hydraulic traits of several years back in time.

#### 1. Introduction

Accurate predictions of carbon dioxide ( $CO_2$ ) exchange by forest ecosystems are essential for understanding, e.g., the role of the forest mitigation in the context of the National Determined Contribution under the Paris Agreement, as well as for the required estimates of annual carbon (C) budgets to be provided at national or

©2016. American Geophysical Union. All Rights Reserved. international level. Research in the past decades focused on improving these predictions on both annual and longer (decadal) timescales, e.g., in relation to extreme events [e.g., Baldocchi and Wilson, 2001; Ciais et al., 2005; Richardson et al., 2009; Rodrigues et al., 2011; Wolf et al., 2013; Wu et al., 2013] and in relation to the length of the growing seasons or the number of carbon uptake days [e.g., Churkina et al., 2005]. Our study builds on the current understanding that some critical periods within the current or the past year (e.g., winter frost and spring drought) may explain the interannual variability of C uptake of forests better than average conditions over the current year only [Le Maire et al., 2010]. The effects of climatic conditions from previous seasonal periods on current year's annual net ecosystem productivity (NEP<sub>c</sub>) are called carry-over effects and were quantified, e.g., by Shao et al. [2016], Thomas et al. [2009], and Zielis et al. [2014]. Such carry-over effects support the influence of specific periods in the past on current year's NEP<sub>c</sub>, and their influence have been demonstrated a long time ago by tree ring analyses, e.g., for Danish forests [Holmsgaard, 1955]. Here we use positive NEP defined as net C uptake, while negative NEP is a net C release to the atmosphere (NEE is defined with the opposite sign in Aubinet et al. [2012]). Further, NEP<sub>c</sub> is defined as the cumulative sum of NEP fluxes throughout the annual cycle—not necessarily a calendar year—yielding net C flux between the atmosphere and the forest. To account for the temporal integration of the average NEP<sub>c</sub> over an annual cycle, values are expressed in  $gCm^{-2}yr^{-1}$ , whereas half-hourly NEP measurements are given in  $gCm^{-2}s^{-1}$  (as calculated from  $\mu$ mol m<sup>-2</sup> s<sup>-1</sup>).

#### 1.1. The Concept of cDOY

Following the concept of previous year's weather conditions influencing current year NEP<sub>c</sub>, we explored the information content of cDOY, defined as the day of year when the net carbon losses accumulated during the wintertime are compensated by net assimilation in spring. The timing of cDOY is assumed to change with climatic conditions of previous periods (of unknown length) and may have a direct impact on the current year NEP<sub>c</sub> [*Zielis et al.*, 2014]. Similar approaches were described in literature, e.g., the "zero-crossing time," wherein net ecosystem exchange is used to define the time when the forest ecosystem turns from a C source in winter into a C sink in spring [*Gonsamo et al.*, 2012a; *Gonsamo et al.*, 2012b]. Another approach quantifies the so-called "start of the carbon uptake period" which is determined by a sharp increase in gross primary production (GPP) [*Delpierre et al.*, 2009]. However, these approaches rely on instant net ecosystem exchange rates only and do not accumulate carbon loss over an entire wintertime, as it is the case of in the cDOY approach.

#### **1.2.** Integration Methods for NEP<sub>c</sub>

Traditionally, NEP<sub>c</sub> is integrated annually over a time period of the Gregorian calendar year (classical integration as shown in Figure 1). This is more a practical choice, but it neither reflects any particular connection to underlying carbon cycle processes nor does it take into account potential carry-over effects on NEP<sub>c</sub>. As an example, trees prepare their buds in autumn and thus the predisposition for growth (and thus NEP<sub>c</sub>) during the following season is determined in autumn already. Thus, it is important to consider the start and end of the accumulation period of NEP<sub>c</sub>. In line with these thoughts, *Urbanski et al.* [2007] introduced a method integrating NEP<sub>c</sub> at Harvard Forest from 28 October to 27 October of the following year (Urbanski integration in Figure 1), trying to come closer to a more reasonable biological time reference of the annual NEP cycle. This integration period is similar to the hydrological year as starting on 1 November in the Northern Hemisphere. *Thomas et al.* [2009] found that interannual and seasonal variations in carbon and water processes were best explained when seasonality was defined functionally within hydrological years.

More recently, a dynamic integration approach was introduced by *Zweifel et al.* [2010] in order to relate continuous stem diameter fluctuations to NEP<sub>c</sub>. The frequently occurring stem shrinkages induced by winter frost [*Zweifel and Hasler*, 2000] made the classical integration approach from 1 January to 31 December inapplicable for an unbiased analysis of annual stem growth increments (bark and wood) in relation to NEP<sub>c</sub>. An integration over a variable period was therefore proposed (Figure 1), starting with the day when NEP<sub>c</sub> of the previous calendar year reached its maximum and ending with the day in the current year when maximum NEP<sub>c</sub> was achieved (dynamic integration in Figure 1). Thus, the dynamic year corresponds more closely to the actual biological cycle, which does not exactly count 365.25 days per year. This dynamic integration method is appropriate to time series of stem increments and NEP<sub>c</sub> data from eddy covariance flux measurements. It was concluded that the application of this approach reduces distortion effects on annual sums, due to apparent interannual variations in carbon losses and stem shrinkages during wintertime [*Zweifel et al.*,



**Figure 1.** Three different methods of integrating net ecosystem productivity (NEP<sub>c</sub>) over time (real data shown: Hyytiälä, years 2010 to 2012): Classical integration runs from 1 January to 31 December of each calendar year, "current" is year 2; Urbanski integration from 28 October of the previous year to the end of 27 October of the current year; Dynamic integration runs dynamically for every site and for every year from the day of the previous year's cumulated NEP<sub>c</sub> maximum (MAX<sub>NEP<sub>c</sub></sub>) to the current year's cumulated NEP<sub>c</sub> maximum (MAX<sub>NEP<sub>c</sub></sub>). The day of compensation (cDOY) is defined as the day of the year when MAX<sub>NEP<sub>c</sub></sub> (of the previous year) is crossed by NEP<sub>c</sub> in the current year. Accordingly, cDOY depends on the integration method. For the Southern Hemisphere, i.e., for the Australian site AU-Tum, the same cuts were made, one half year later. The corresponding year started on 1 July and ended on 30 June.

2010], i.e., shifts in uptake and loss periods that arbitrarily affect the sums calculated over a fixed calendar period. We use the terms "year" and "annual" in combination with all three integration methods, for the sake of readability, being aware that the terms usually are implicitly used for periods of Gregorian calendar years from 1 January to 31 December.

The way of splitting time series into annual integration periods also changes the potential contribution of winter carbon losses for the total annual C uptake and the cDOY timing in the following year (Figure 1). Indeed, the classical integration period splits the net carbon loss of a winter period in two parts, assigning them to two different NEP<sub>c</sub> years, while the dynamic (and Urbanski) integration method assigns net carbon loss for all the winter period entirely to the NEP<sub>c</sub> of the biological year that will last until the onset of the next winter period. Accordingly, cDOY changes with the respective integration method (Figure 1) and might have a different explanatory power for NEP<sub>c</sub>.

In this study, we used in total 26 eddy covariance forest sites with 25 sites throughout Europe and North America (Figure 2), and additionally one site from Australia, thus covering a wide range of climatic conditions (Table 1) to investigate the meaning of cDOY for NEP<sub>c</sub> and its underlying drivers. We used the cDOY timing as the key measure associated with the net carbon loss period and related it to climatic conditions and NEP<sub>c</sub>. We addressed the following specific objectives: (1) application of three different NEP<sub>c</sub> integration methods (classical, Urbanski, and dynamic) in order to calculate and compare the respective cDOYs, (2) identification of climatic and biological drivers for cDOY across sites and across different years, (3) evaluation of different cDOY as a predictor for its associated NEP<sub>c</sub>, and (4) the weight of winter net respiratory losses on current year's NEP<sub>c</sub>.

#### 2. Materials and Methods

#### 2.1. Study Sites

The study is based on carbon dioxide  $(CO_2)$  flux data from 347 site years from 26 eddy covariance (EC) forest sites (managed forest not affected by major disturbances like fire or wind throw) within Europe, North America, and Australia (Table 1 and Figure 2). The selected sites fulfilled the following criteria: (1) at least



**Figure 2.** Spatial distribution of 25 sites across North America and Europe and one Australian site (not shown). Site abbreviations are listed in Table 1. ENF = evergreen needleleaf forests, DBF = deciduous broadleaf forests, MF = mixed forests, and EBF = evergreen broadleaf forests.

4 years of continuous EC data, (2) availability of Level 4 (L4) data quality according to the European Fluxes Database [*European Fluxes Database Cluster*, 2014] or available from the FluxNet2015 data set (http://fluxnet.fluxdata.org/data/fluxnet2015-dataset/), and (3) available meteorological and forest characteristics data. The forest vegetation at the sites was classified as deciduous broadleaf forests (DBF, n = 7), mixed forest (MF, n = 3), evergreen needleleaf forests (ENF, n = 13), and evergreen broadleaved forest (EBF, n = 3).

#### 2.2. CO<sub>2</sub> Flux Measurements

Half-hourly or hourly CO<sub>2</sub> flux data (net ecosystem exchange rates summed up to net ecosystem productivity, NEP), derived from both open- and closed-path gas analyzers, were downloaded from the FluxNet2015 data set (http://fluxnet.fluxdata.org/data/fluxnet2015-dataset/) or in L4 quality from the European Fluxes Database (http://www.europe-fluxdata.eu/home). These data were already filtered and gap filled (Table 1). For three sites (CH-DAV, CH-LAE, and PL-TUC, all open-path gas analyzers) our own site-specific processing was conducted: data were filtered for unfavorable atmospheric conditions such as snow, heavy rain, and/or dust which increased window dirtiness of the infrared gas analyzer > 70%. For these three individual sites, the threshold for insufficient nocturnal turbulent mixing of the atmosphere (determined via the friction velocity  $u_*$  for mechanical turbulence) was determined with the online EC gap-filling and flux partitioning tool (Markus Reichstein and Olaf Menzer, http://www.bgc-jena.mpg.de/~MDlwork/eddyproc/) [*Reichstein et al.*, 2005] and was found to be 0.2 m s<sup>-1</sup>.

#### 2.3. CO<sub>2</sub> Flux Integration

Annual NEP<sub>c</sub> was integrated with three different methods (Figure 1): classical method: NEP<sub>c</sub> is integrated from January 1 to December 31; Urbanski method: NEP<sub>c</sub> is integrated from 28 October to 27 October 1 year later [*Urbanski et al.*, 2007]; and dynamic method: integration of NEP<sub>c</sub> from the DOY with the maximum seasonal peak of the previous year (typically in fall;  $MAX_{NEP_c}$ ) to the DOY with  $MAX_{NEP_c}$  of the current year (Figure 1). The dynamic integration method led to "annual cycles" ranging from 7 to 16 months depending on year and site; the overall average was 364 days (supporting information Figure S1). For the Southern Hemisphere site AU-Tum, the year has been shifted half a year forward, i.e., the classical year started with 1 July and the dynamic "integration period" started with the maximum peak ( $MAX_{NEP_c}$ ) before 1 July. The Urbanski integration method was not applied for this site.

#### 2.4. Statistical Analyses

Statistical analyses were performed using the statistical software R, version 3.3.1 [*R Development Core Team*, 2013]. All multiple regression models were based on linear relationships. Adjusted  $R^2$  (adj $R^2$ ) was used for the quantification of goodness of fit. Analyzed potential drivers for cDOY are listed in Table 1. Their respective impacts on cDOY were analyzed with multiple regression models based on the inclusion of explaining

Table 1.	Selected Characteristics of	f the Study Site	es (See Also	o Figure 2) <sup>~</sup>										
Code	Name	Forest Type	Lat.	Long.	Year Range	MAP	MAT	Altitude	Age	Height	LAI	Z	Data Access	Data in This Table From
AU-TUM	Tumbarumba	EBF	-35.66	148.15	2002-2013	1924	9.6	1249	100	40	2.5	I	FluxNet2015	Beringer et al. [2016]; EluvMat2015 mata*: pr*
BE-VIE	Vielsalm	MF	50.31	9	1997–2014	1062	7.8	493	95	35	4.6	10.2	FluxNet2015	Flechard et al. [2011]; Elicitation et al.
CA-GRO	Ontario, Groundhog River, Boreal Mixed	MF	48.22	-82.16	2004-2013	831	1.3	340	84	32	I	ł	FluxNet2015	McCaughey et al. [2006], Gökkaya et al. [2014], and Gökkaya et al. [2014], and
CA-QFO	Quebec, Eastern Boreal, Mature	ENF	49.69	-74.34	2004–2010	962	-0.4	382	105	13.8	3.7	I	FluxNet2015	FluxNet2015 meta; pc Coursolle et al. [2012]; FluxNet2015 meta; pc
CH-DAV CH-LAE	black spruce Davos Lägeren	ENF MF	46.81 47.48	9.86 8.37	1997–2011 2004–2011	1046 1211	3.5 8.7	1662 682	240 140	25 31	3.9 3.6	1.5 14.3	own site own site	own site; FluxNet2015 meta own site; <i>Flechard et al</i> .
CZ-BK1 DE-HAI	Bily Kriz Hainich	ENF DBF	49.5 51.08	18.54 10.45	2001–2010 2000–2012	1316 720	6.7 8.3	875 430	30 125	10 33	7.5 6	10.5 12.6	download, L4 FluxNet2015	[2011]; FluxNet2015 meta Flechard et al. [2011]; pc Knohl et al. [2003], Mund et al. [2010]
DE-THA	Tharandt	ENF	50.96	13.57	1997–2014	820	7.7	385	125	26.5	7.6	12.5	FluxNet2015	Flechard et al. [2010], and Herbst et al. [2011], and FluxNet2015 meta; pc Grünwald and Bernhofer [2007]
DK-SOR	Soroe	DBF	55.49	11.65	1997–2014	660	8.2	40	100	35	4.8	14.6	FluxNet2015	and <i>Flechard et al.</i> [2011]; FluxNet2015 meta; pc <i>Pilegaard et al.</i> [2011];
ES-ES1 FI-HYY	El Saler Hyytiälä	ENF ENF	39.35 61.85	-0.32 24.3	2000–2007 1997–2014	551 709	17.9 3.8	1 181	100 40	10 14	3.5 2.5	16.9 3.5	download, L4 download, L4	FluxNet2015 meta; pc FluxNet2015 meta; pc Flechard et al. [2011];
FR-PUE	Puechabon	EBF	43.74	3.6	2000–2014	883	13.5	270	73	5.5	2.9	ł	FluxNet2015	FluxNet2015 meta; pc Flechard et al. [2011];
IT-COL	Collelongo	DBF	41.85	13.59	1997–2011	1180	6.3	1645	105	25	2	5.3	download, L4	FluxNet2015 meta; pc Scartazza et al. [2016]
IT-LAV IT-REN	Lavarone Renon	ENF	45.96 46.59	11.28 11.43	2003–2014 1999–2013	1291 809	7.8 4.7	1353 1730	100 200	28 31	8.1 5.1	15.4 4.8	FluxNet2015 FluxNet2015	FluxNet2015 meta; pc FluxNet2015 meta; pc Flechard et al. [2011];
NL-LOO	Loobos	ENF	52.17	5.74	1997–2013	996	10	25	06	18	1.9	32.4	FluxNet2015	FluxNet2015 meta; pc Flechard et al. [2011];
PL-TUC PT-ESP	Tuczno Espirra	ENF EBF	53.21 38.64	16.1 —8.6	2008–2011 2002–2008	625 665	7.8 15.4	105 85	54 12	20 20	3.1	8.5 7.2	STSM, direct download, L4	FluxNet2015 meta; pc FluxNet2015 meta; pc Pita et al. [2013],
RU-FYO	Fyodorovskoye	ENF	56.46	32.92	1998–2014	711	3.9	265	196	25.7	3.5	8.3	FluxNet2015	Rodrigues et al. [2011], and Flechard et al. [2011]; pc Flechard et al. [2011]; Flechard et al. [2011];
SE-NOR	Norunda	ENF	60.09	16.22	1996–2011	527	5.5	45	100	25	S	3.4	STSM, direct	Lagergren et al. [2008] and
US-HA1	Harvard Forest EMS Tower (HFR1)	DBF	42.54	-72.15	1992–2012	1071	6.6	340	80	18	3.7	6.4	FluxNet2015	Munger et al. [1996] and Munger et al. [1996] and Eli-MAPOTE model of
US-ME2	Metolius-intermediate aged ponderosa pine	ENF	44.45	-121.55	2002-2014	523	6.3	1253	06	14	3.2	-	FluxNet2015	ruxiverzoro mera, pc

Table 1. (c	ontinued)													
Code	Name	Forest Type	Lat.	Long.	Year Range	MAP	MAT	Altitude	Age	Height	LAI	N	Data Access	Data in This Table From
SMM-SU	Morgan Monroe State Forest	DBF	39.92	-86.41	1999–2014	1032	10.9	275	06	27	4.7	9	FluxNet2015	Schwarz et al. [2004] and Thomas et al. [2009]; FluxNet2015 meta; pc Schmid et al. [2000], Dragoni et al. [2011], Brzostek et al. [2014], and
US-NR1	Niwot Ridge Forest	ENF	40.03	-105.55	1999–2014	800	1.5	3050	115	13	4	Q	FluxNet2015	Roman et al. [2015]; FluxNet2015 meta; pc Sievering et al. [2001];
US-UMB	(LTER NWT1) Univ. of Mich. Biological Station	DBF	45.56		2000-2014	803	5.38	234	93	22	3.5	7.5	FluxNet2015	FluxNet2015 meta; pc <i>Gough et al.</i> [2008], <i>Nave et al</i> [2009], and
US-WCR	Willow Creek	DBF	45.8		1999–2014	787	4.02	520	70	24.2	5.36	I	FluxNet2015	<i>Gough et al.</i> [2013]; FluxNet2015 meta; pc <i>Cook et al.</i> [2004]; FluxNet2015 meta; pc
<sup>a</sup> Site nar EBF = everg of the indicc Age = avera stated from section 2). S *personal cc <i>et al.</i> [2011], Lagergren <i>et</i> [2015], Scart	mes are abbreviated w reen broadleaf forests. I ted years used in this sti ge age of the mature tre <i>Flechard et al.</i> [2011]; De tite characteristics were mmunication (pc). Not. <i>Flechard et al.</i> [2011], Gö al. [2008], McCaughey e azza et al. [2016], Schmi	ith the FLUXNE Further terms in udy. MAP = meai es in the stand (y ita access: NEP <sub>C</sub> , obtained from h available data ar <i>kkaya et al.</i> [2014, <i>t al.</i> [2006], <i>Mun.</i>	T codes. I the heade n annual sı rears); Heig data were i tittps://fluxr e indicatec I, Gökkaya d et al. [201	Forest type is have the f and of precip (ht = maxim obtained fr obtained fr et al. [2015] 0], Munger il. [2004], Sie	s are abbrevi ollowing meal oitation (mm). I um tree heigh: om the *Flu eferences in Ta <i>Gough et al.</i> [298], <i>M</i> t <i>et al.</i> [1998], <i>M</i> t	ated as ning: Lat MAT = m MAT = m t(m); LAI Database Database xNet201 ible 1 are 008], Goi 001], and	DBF = d t., Latituc nean ann = leaf ar = leaf ar : Cluster : Cluster 5 Metad ugh et al. 1996 3 Thoma	eciduous   de (degree: ual air tem) ea index (n in Level 4 q fata excel s ws: Beringe vws: Beringe J, Nave et a. S et al. [200	s, WGS8 s, WGS8 perature t;( <sup>2</sup> ח ח luality, fi luality, fi heet, fro <i>i net al.</i> [2 <i>i nwald o</i> <i>i</i> . [2009].	af forest: 4) and Lo e (°C) at th N = meau rom http:/ 016], Brz nnd Bernh , Pilegaau	s, ENF = bng. = Lc ne top of nannual FluxNvt2 //www.b <i>instek et c</i> <i>instek et d</i> <i>inster</i> [200	evergre nigitude nitroge 015 dat gc-jena 11. [2014 2011], <i>Pirt</i>	ten needleleaf fo (degrees, WG588 hy towers; Altitude n deposition (kg h. a set or from the re mpg.de/public/cz 1, Cook et al. [2013], Roi ta et al. [2013], Roi ta et al. [2013], Roi	rests, MF = mixed forests, and t) of the site. Year Range = data = meter above sea level (m asl); a <sup>-1</sup> yr <sup>-1</sup> ), most data and where espective PIs incl. own sites (see irboeur/sites/SITE.html, or from J, Coursolle et al. [2012], Dragoni h et al. [2012], Knohl et al. [2003], trigues et al. [2011], Roman et al.

variables in a stepwise way. The so-called standardized regression coefficients ( $\beta$  coefficients) were used to determine the relative importance of variables (var) within the models, ranging between -1 as the highest negative and +1 as the highest positive correlative importance [*Quinn and Keough*, 2002]. A  $\beta$  coefficient close to zero indicates that the variable does not add to the quality of the model.

#### 3. Results

#### 3.1. Compensation of Net Carbon Loss After Wintertime

The day of year when respiratory carbon losses from the previous winter were compensated (cDOY) differed strongly across sites (Table 2 and Figure 3). cDOY varied from 3 January (AU-TUM/3 July) to 25 July (CA-QFO), with a mean of 3 May (obtained by averaging all three integration methods, Figure 3). Some sites showed no or irregular cDOY timings, meaning that they observed no distinct respiratory carbon loss period every year (Table 2). Evergreen forests ( $3 \times \text{EBF}$ ,  $13 \times \text{ENF}$ ) in general had an earlier cDOY (18 April) than deciduous forests ( $7 \times \text{DBF}$ , June 28). Only nine out of the 26 sites compensated on average their net carbon losses in the climatologically defined spring calendar months (Mar–May) (Table 2). Six sites compensated before spring, while eleven compensated after May. The yearly standard deviation of cDOY for individual sites ranged from 6 days (DE-HAI, AU-TUM) to more than 50 days (PT-ESP) (Table 2).

Further, cDOY strongly depended on the integration method. In general, the classical integration method led to a cDOY almost 3 weeks earlier than those obtained with the dynamic method (classical: 16 April; dynamic: 10 May). The average cDOY obtained from the Urbanski integration method (5 May) was almost the same as that from the dynamic method (data now shown). Much less affected were the mean differences (Urbanski versus classical:  $-54 \text{ g Cm}^{-2} \text{ yr}^{-1}$  and dynamic versus classical:  $-91 \text{ g Cm}^{-2} \text{ yr}^{-1}$ ) and the standard deviations (Urbanski versus classical:  $-7 \text{ g Cm}^{-2} \text{ yr}^{-1}$  and dynamic versus classical:  $-10 \text{ g Cm}^{-2} \text{ yr}^{-1}$ ) of NEP<sub>c</sub> between the different integration methods (see also supporting information figures for each site).

#### 3.2. Drivers of cDOY

Average cDOY was substantially correlated with mean annual air temperature ( $R^2$  between 0.4 and 0.45). The relationship was largely independent of the integration method used (Table 3), and the later cDOYs corresponded to the cooler sites (Figure 3a). Other site characteristics considered (latitude, longitude, altitude, tree age, nitrogen deposition, tree height, and mean annual precipitation) showed weak (or no) linear relationship to cDOY and did not improve the stepwise multiple linear regression models to explain cDOY (Table 3).

The meaning of mean annual site temperature (MAT) for cDOY was markedly increased when the pooled data over all sites were grouped into four forest types (Figure 3a): evergreen needleleaf forest (ENF, all included), evergreen broadleaf forest (EBF), and mixed forest (MF) showed  $R^2$  between 0.64 and 0.99. No significant correlation was found between MAT and cDOY for the deciduous broadleaf forests (DBF;  $R^2 = 0.07$ , p > 0.05).

In Figure 3b, those sites without a distinct winter respiratory loss period, and thus with no consistent cDOY timing (Table 2) were removed (all EBF and more than 50% of the ENF sites). All of these sites are evergreen, with a majority having MAT over 8–10°C; hence, in winter, these sites likely photosynthesize. The remaining six ENF sites (CA-QFO, CH-DAV, FI-HYY, RU-FYO, SE-NOR, and US-NR1), with a distinct winter respiratory loss and a latter cDOY, increase to an  $adjR^2$  of 0.90 for the linear relationship between MAT and cDOY (Figure 3b).

#### 3.3. Relationship Between cDOY and NEP<sub>c</sub>

The 26 sites analyzed in this study included C sink and C source sites (Table 2). The largest net annual respiratory loss was at RU-FYO with a consistent average C output of  $137 \text{ g Cm}^{-2} \text{ yr}^{-1}$ . The largest net C uptake was at AU-Tum with 1007 g C m<sup>-2</sup> yr<sup>-1</sup>.

Stepwise multivariate analysis showed that cDOY, among the site characteristic variables available, explained most of NEP<sub>c</sub> for all integration methods (Table 4). Sites with distinct winter respiratory loss, explained significantly more of NEP<sub>c</sub> than all other sites. cDOY obtained from the two integration approaches that initiated the NEP<sub>c</sub> year in autumn (Urbanski and dynamic) explained NEP<sub>c</sub> significantly better (adj $R^2$  = 0.35 and 0.47) than cDOY from the classical integration approach (adj $R^2$  = 0.23). When the ENF (Table 4d) and DBF (Table 4e) sites were analyzed separately (using the dynamic integration), the  $R^2$  of the linear regressions was further improved ( $R^2$  of 0.58 and 0.68, respectively).

JU,	Journal of G	eop	hy	Sİ	Ca		R	es	e	ar	'C	h:	B	ic	g	e	09	SC	ie	n	ce	S		
	Net Ecosystem	Classification	without	with	with	with	with	with	without	with	without	with	without	with	without	with	without	without	without	without	with	with	with	and the second
	nd Integrated	% Winter Resp. Loss	5.4%	-19.8%	-143.6%	-420.3%	-80.8%	-16.1%	-6.6%	-51.4%	-7.5%	-88.6%	-3.6%	-41.6%	-8.6%	-28.2%	-7.4%	-9.7%	0.1%	-6.5%	104.7%	-52.0%	-401.0%	0000
	npensated) a	Average SD NEP <sub>c</sub>	159.9	173.3	52.8	14.7	74.0	122.2	128.9	68.0	81.7	151.7	164.9	53.3	88.1	179.8	85.9	160.7	82.9	363.0	145.4	37.1	102.5	1110
	Were Com	p Dyn.	0.51	0.00	0.00	0.93	0.00	0.01	0.95	0.42	0.07	0.00	0.06	0.00	0.41	0.00	0.61	0.00	0.16	0.11	ł	0.01	0.00	000
	ory Losses	R <sup>2</sup> Dyn	0.05	0.74	0.70	0.01	0.87	0.68	0.00	0.06	0.20	0.82	0.47	0.59	0.06	0.53	0.03	0.70	0.70	0.51	1.00	0.51	0.78	50
	er Respirat	p Urb.	I	0.00	0.02	0.26	0.00	0.02	0.94	0.25	0.08	0.00	0.85	0.00	0.16	0.00	0.52	0.00	I	I	0.29	0.00	0.00	0.05
	/hen Winte	R <sup>2</sup> Urb.	1	0.73	0.52	0.24	0.88	0.72	0.00	0.12	0.18	0.84	0.01	0.53	0.19	0.53	0.04	09.0	0.00	1.00	0.19	0.86	0.75	
	/ of Year W	p Cla.	0.48	0.00	0.05	0.54	0.00	0.00	0.45	0.25	0.10	0.00	0.57	0.10	0.72	0.01	0.29	0.00	0.28	0.13	0.07	0.00	0.00	0000
	DOY = Day	R <sup>2</sup> Cla.	0.05	0.62	0.40	0.08	0.74	0.76	0.07	0.12	0.16	0.66	0.06	0.19	0.01	0.45	0.11	0.52	0.53	0.48	0.33	0.67	0.54	
	nsation Days (cl and Dynamic) <sup>a</sup>	сроу	Jul 03 ± 6	Apr 29 $\pm$ 28	Jun 27 $\pm$ 15	Jul 23 $\pm$ 23	Mai 28 $\pm$ 34	Mai 10 $\pm$ 14	Apr 04 $\pm$ 8	Jun 21 $\pm$ 6	Apr 01 $\pm$ 18	Jun 26 $\pm$ 18	Jan 27 $\pm$ 27	Mai 26 $\pm$ 10	Feb 01 $\pm$ 22	Jun 09 $\pm$ 10	Mai 10 $\pm$ 28	$Mrz 22 \pm 41$	Jan 28 $\pm$ 30	Feb 02 $\pm$ 50	Jun 13 $\pm$ 26	Apr 15 $\pm$ 31	Jul 24 $\pm$ 22	L-L-00 - 40
	Between Compe assical, Urbanski	n cDOY Dyn.	12	18	10	ε	15	80	10	13	18	16	8	16	14	13	12	17	4	9	2	12	19	ç
	id Relationships tion Method (Cl	n cDOY Urb.	:	18	10	7	15	7	10	13	18	18	9	16	12	13	12	16	-	2	8	15	19	L
	Y Averages, an ite and Integra	n cDOY Cla.	12	18	10	7	15	8	10	13	18	18	8	16	14	13	12	17	4	9	11	15	21	ç
	lEP <sub>c</sub> Averages, cDC / (NEP <sub>c</sub> ) for Each 5	Mean NEP <sub>c</sub>	$1006.7 \pm 273.4$	$461.9 \pm 155.6$	$\textbf{120.6}\pm\textbf{47.3}$	$\textbf{2.8}\pm\textbf{14.5}$	$134.1\pm70.8$	$579.7 \pm 124.5$	$798.9 \pm 132.8$	$579.2 \pm 73.2$	$617.9 \pm 81.1$	$176\pm139.4$	$415.5 \pm 154.9$	$\textbf{244.9}\pm\textbf{50.5}$	$\textbf{230.5}\pm\textbf{89.2}$	$622 \pm 179$	$248\pm96.7$	$427.5 \pm 144.6$	$667.5 \pm 76.1$	$459.7 \pm 356.6$	$-136.7 \pm 128.2$	$\textbf{52.8} \pm \textbf{33.8}$	$211.2 \pm 188$	C 10C - 001
	<b>Table 2.</b> N Productivity	oite	AU-TUM	3E-VIE	CA-GRO	CA-QFO	CH-DAV	CH-LAE	CZ-BK1	DE-HAI	DE-THA	<b>DK-SOR</b>	ES-ES1	т	-R-PUE	T-COL	T-REN	VIL-LOO	PL-TUC	PT-ESP	RU-FYO	SE-NOR	JS-HA1	IC AAFA

PL-TUC **RU-FYO** 

FI-HYY IT-COL IT-REN

Site

**BE-VIE** 

with with with EDOY = show Average SD ification into	-48.3% -80.5% -80.7% (d dynamic); <i>n</i> - ard deviations, age NEP <sub>c</sub> ; class	31.8 31.4 164.6 Jrbanski, an Iding stand s from aver	0.19 0.02 0.00 (classical, L gated inclu biratory los:	0.12 0.34 0.67 methods rrs investig winter resp	0.25 0.05 0.01 ntegration of the yea	0.09 0.27 0.55 all three ii all three ii age perce	0.71 0.10 0.04 0.04 iged over compensa loss = aver	0.01 0.19 0.35 0.35 an NEP <sub>c</sub> ( spiratory l	Jun 02 $\pm$ 9 Jul 04 $\pm$ 10 Jun 29 $\pm$ 41 eviation (g m <sup>-2</sup> yr Mean cDOY = me thods; % winter re	16 15 11 nd standard d ion methods; ion methods; inatory loss (th	16 15 12 m productivity ai of the integrati over all three in tinct winter resp	16 15 12 net ecosyster ated for each EP <sub>c</sub> averaged or without dis	170.7 $\pm$ 30.1 268.9 $\pm$ 65.8 271.9 $\pm$ 153.6 IEP <sub>c</sub> = mean annual Ys could be calcula dard deviation of N arte is a site with	-NR1 -UMB -WCR -WCR -Mean N any cDO -ether the
with	-89.7%	164.6	0.00	0.67	0.01	0.55	0.04	0.35	Jun 29 $\pm$ 41	11	12	12	271.9±153.6	S-WCR
with	48.3% 80.5%	31.8 31.4	0.02	0.12 0.34	cz.u 0.05	0.27 0.27	0.10	0.19	Jul 04 $\pm$ 10	15	15	15	$1/0.7 \pm 30.1$ 268.9 $\pm$ 65.8	5-UMB
	04.6.00-				10.0	00		<u>-</u>		2 ;	2 ;	2 ;		
without	-0.9%	316.6	0.00	0.67	0.05	0.76	0.00	0.59	Feb 08 $\pm$ 40	13	5	13	$588 \pm 291.2$	S-ME2
with	-401.0%	102.5	0.00	0.78	0.00	0.75	0.00	0.54	Jul 24 $\pm$ 22	19	19	21	$211.2 \pm 188$	S-HA1
with	-52.0%	37.1	0.01	0.51	0.00	0.86	0.00	0.67	Apr 15 $\pm$ 31	12	15	15	$\textbf{52.8} \pm \textbf{33.8}$	-NOR
with	104.7%	145.4	ł	1.00	0.29	0.19	0.07	0.33	Jun 13 $\pm$ 26	2	8	11	$-136.7 \pm 128.2$	J-FYO
	-6.5%	363.0	0.11	0.51	I	1.00	0.13	0.48	Feb 02 $\pm$ 50	D	2	9	$459.7 \pm 356.6$	r-esp
without										ų				

CH-LAE



**Figure 3.** Relationship between cDOY (integrated dynamically) and mean annual temperature (MAT) grouped for the four forest types. (a) All sites included (n = 26) and (b) sites with distinct winter respiratory losses only, i.e., where winter net respiration loss accounts for more than 10% of the annual net ecosystem productivity and n was > 2 for all integration methods.

Drivers of cDOY	<i>R</i> <sup>∠</sup> Alone	1 Var	2 Vars	3 Vars	4 Vars	5 Vars	6 Vars	7 Vars
		(	Classical Integ	ration Metho	d			
MAT	0.4***	-0.63	-0.58	-0.57	-0.58	-0.53	-0.55	-0.48
LAI	0.11		0.25	0.25	0.23	0.13	0.14	0.03
Age	0.06			0.03	-0.02	-0.06	-0.03	0.06
Height	0.05				0.1	0.29	0.27	0.23
Ν	0.03					0.08	0.07	-0.05
Altitude	0.01						-0.07	-0.19
MAP	0							0.37
Total adjR <sup>2</sup>		0.4***	0.39***	0.36**	0.34**	0.17*	0.12*	0.21*
		L	Irbanski Integ	ration Metho	d			
MAT	0.45***	-0.67	-0.61	-0.58	-0.59	-0.61	-0.43	-0.5
Height	0.17*		0.29	0.29	0.31	0.34	0.35	0.27
MAP	0.1			0.21	0.21	0.21	0.28	0.33
Age	0.07				-0.06	-0.09	-0.04	0.06
LAI	0.06					-0.01	-0.03	-0.01
Ν	0.05						-0.06	-0.12
Altitude	0.01							-0.24
Total adj <i>R</i> <sup>2</sup>		0.45***	0.48***	0.51***	0.49***	0.45***	0.23*	0.21*
		Ľ	ynamic Integ	ration Metho	d			
MAT	0.4***	-0.63	-0.57	-0.57	-0.57	-0.5	-0.43	-0.52
LAI	0.1		0.23	0.23	0.22	0.1	0.01	0.04
Age	0.06			0.03	0	-0.06	-0.03	0.1
Height	0.03				0.06	0.36	0.36	0.24
Ν	0.03					0.04	-0.02	-0.09
MAP	0.01						0.23	0.3
Altitude	0							-0.33
Total adjR <sup>2</sup>		0.4***	0.37**	0.34**	0.31**	0.2*	0.2*	0.22*

**Table 3.** Stepwise Multiple Linear Regression Models to Determine the Drivers of the Day of Compensation cDOY for the Classical, Urbanski, and Dynamic Integration Method<sup>a</sup>

<sup>a</sup>The variables were included one by one in the models (MAT = mean annual air temperature at the top of the eddy towers; LAI = leaf area index; Age = average age of the mature trees in the stand; Height = maximum tree height; N = mean annual nitrogen deposition; MAP = mean annual sum of precipitation). The  $\beta$  coefficients (var) indicate the relative importance of the variable, ranging from -1 (highest importance, negative correlation) to +1 (highest importance, positive correlation). The first column gives the  $R^2$  for individual site characteristics (see Table 1).

\*p < 0.05, \*\*p < 0.01,

\*\*\*\**p* < 0.001.

Table 4. Stepwise Multiple Linear Regression Models to Determine the Drivers of Net Ecosystem Productivity NEPc for the Classical (All Sites), Urbanski (All Sites), and Dynamic Integration Method (All Sites)<sup>a</sup>

Drivers of $NEP_c$	R <sup>2</sup> Alone	1 Var	2 Vars	3 Vars	4 Vars	5 Vars	6 Vars	7 Vars	8 Vars
			Classical	Integration Met	hod (All Sites) <sup>b</sup>				
cDOY	0.23*	-0.48	-0.48	-0.39	-0.44	-0.38	-0.36	-0.59	-0.6
MAP	0.22*		0.47	0.47	0.41	0.33	0.26	0.45	0.38
MAT	0.14			0.14	0.14	0.27	0.25	0.14	0.1
Height	0.05				0.21	0.24	0.36	0.39	0.33
Altitude	0.04					0.24	0.38	0.38	0.35
Age	0.04						-0.31	-0.39	-0.38
N	0.02							0.04	0.07
LAI	0.02								0.21
Total adi <sup>2</sup>		0.23*	0.45**	0.47**	0.51**	0.55**	0.61**	0.54	0.57
, , , , , , , , , , , , , , , , , , ,			Urbanks	i Integration Met	hod (All Sites) <sup>c</sup>				
cDOY	0.35**	-0.59	-0.7	-0.74	-0.67	-0.64	-0.83	-0.85	-0.85
MAP	0.23*		0.38	0.38	0.33	0.31	0.34	0.42	0.37
MAT	0.15			-0.06	0.06	0.06	0	0.01	-0.01
Altitude	0.05				0.18	0.24	0.34	0.31	0.29
Age	0.04					-0.14	-0.33	-0.33	-0.33
Height	0.03						0.45	0.38	0.34
N	0.02							-0.03	-0.01
LAI	0								0.14
Total adjR <sup>2</sup>		0.35**	0.47**	0.48**	0.5**	0.51**	0.65***	0.69**	0.71**
			Dynamic	Integration Met	hod (All Sites) <sup>d</sup>				
cDOY	0.47***	-0.68	0.39	-0.66	-0.6	-0.58	-0.6	-0.84	-0.86
MAP	0.22*		-0.64	0.39	0.35	0.34	0.23	0.45	0.39
MAT	0.14			-0.04	0.06	0.05	0.06	0	-0.04
Altitude	0.06				0.15	0.2	0.29	0.27	0.24
Age	0.04					-0.13	-0.25	-0.31	-0.3
Height	0.02						0.29	0.38	0.33
N	0.02							0.01	0.03
LAI	0.01								0.18
Total adj $R^2$		0.47***	0.62**	0.62***	0.63***	0.65***	0.7***	0.72**	0.74**
			Dynamic	Integration Meth	od (ENF Only) <sup>e</sup>				
cDOY	0.58**	-0.76	-0.4	-1.03	-1.08	-1.16	-1.09	-1.04	-0.71
MAT	0.17		-1.07	-0.4	-0.42	-0.38	-0.36	-0.22	0.43
Age	0.13			-0.09	-0.11	-0.06	-0.04	-0.2	-0.72
LAI	0.06				0.35	0.2	0.23	0.15	-0.04
MAP	0.04					0.36	0.36	0.46	0.56
Ν	0.03						0	-0.1	-0.25
Height	0.01							0.22	0.71
Altitude	0.01								0.53
Total adjR <sup>2</sup>		0.58**	0.65	0.66**	0.78**	0.87***	0.87**	0.88*	0.94*
			Dynamic	Integration Meth	od (DBF Only)'				
cDOY	0.68*	-0.83	0.37	-0.44	-0.4	-0.37			
Altitude	0.53		-0.61	0.43	0.44	0.16			
Age	0.33			0.23	0.22	0.29			
LAI	0.27				0.07	0.24			
MAP	0.17					0.38			
Ν	0.11								
MAT	0.03								
Height	0.02								
Total adjR <sup>2</sup>		0.68*	0.78*	0.81*	0.81	0.86			

<sup>a</sup>The variables were included one by one in the models (cDOY = compensation day; MAT = mean annual air temperature at the top of the eddy towers; LAI = leaf area index; Age = average age of the mature trees in the stand; Height = maximum tree height; N = mean annual nitrogen deposition; MAP = mean annual sum of precipitation). The  $\beta$  coefficients (var) indicate the relative importance of the variable, ranging from -1 (highest importance, negative correlation) to +1 (highest importance, positive correlation). The first column gives the  $R^2$  for individual site characteristics (see Table 1).

<sup>c</sup>Urbanski (all sites). <sup>d</sup>Dynamic (all sites).

<sup>e</sup>The analysis for the dynamic integration for evergreen needleleaf forests (ENF) only.

<sup>f</sup>The analysis for the dynamic integration for deciduous broadleaf forests (DBF) only. Other forest types had too low replications (*n* = 3) for a separate analysis. \*p < 0.05. \*\*p < 0.01.

\*\*\*\**p* < 0.001.



**Figure 4.** Linear regressions between compensation days (cDOY) and annual sums of net ecosystem productivity (NEP<sub>c</sub>) (Table 1) for each site and integration method: (a) classical, (b) Urbanski, and (c) dynamic. Solid regression lines are shown for  $R^2 \ge 0.4$ , broken lines for the rest. The frequency columns at the bottom of each panel indicate the number of site years occurring at a specific cDOY (color coded for the four forest types).

Mean annual temperature (MAT) was the secondary determinant variable of NEP<sub>c</sub> in stepwise multiple linear regression models (Table 4). The ranking of site factors, with minor contributions, such as leaf area index (LAI), mean annual precipitation (MAP), and stand age followed next; however, the ranking depended on the integration method. An exception was the DBF sites (Table 4e): MAT had no explanatory weight for NEP<sub>c</sub> at these sites, in line with the finding that cDOY of these forests was not determined by MAT (Figure 3).

When analyzing individual sites instead of pooled data, the site-specific relationships between cDOY and  $NEP_c$  showed a high variability and ranged from not existing to excellent (annual resolution, Table 1 and Figures 4 and 5). There appeared clear clusters of points (in the scatterplot of cDOY versus  $NEP_c$ , according



**Figure 5.** Differences between modeled annual net ecosystem productivity (NEP<sub>c</sub>), as a linear function of cDOY and measured NEP<sub>c</sub> for each site in a leave-one-yearout cross evaluation for each integration method: (a) Classical, (b) Urbanski, and (c) dynamic. The results are grouped for sites with and without distinct winter net respiratory losses. The Urbanski method was left out for AU-Tum, there, the "year" begins on 1 July and goes to 30 June one year later.

to the forest types in Figure 4): evergreen forests (ENF and EBF) had the lowest cDOY with the highest NEP<sub>c</sub>. Deciduous broadleaf forest (DBF) had the highest cDOYs with on average lower NEP<sub>c</sub>. Mixed forest (MF) had average cDOYs with relatively high NEP<sub>c</sub>.

The site-specific quality of the relationships between cDOY and NEP<sub>c</sub> was largely explicable by grouping the pooled data according to sites with and without a distinct net respiratory carbon loss over wintertime (Table 2). The separation criterion for the two groups was a net respiration loss of 10% of the annual NEP<sub>c</sub> (Table 2). Sites with distinct winter respiration loss had on average a stronger correlation between cDOY and NEP<sub>c</sub> ( $R^2$  0.53 versus 0.37; dynamic method) and were on average 4°C cooler than sites with no distinct winter respiration loss (Tables 1 and 2).

#### 3.4. Quality of NEP<sub>c</sub> Predictions From cDOY

The quality of NEP<sub>c</sub> predictions from cDOY were tested by comparing measured and modeled NEP<sub>c</sub> per site and year with a leave-one-year-out cross evaluation (Figure 5). There were two very clear results: 1. The Urbanski and the dynamic integration methods led to distinctly better NEP<sub>c</sub> prediction than the classical integration method over the Gregorian/orbital calendar year. 2. The NEP<sub>c</sub> predictions from cDOY were stronger for sites with a distinct respiratory carbon loss over wintertime. Thereby, sites where the forest did not become a C source for a distinct period, and thus did not lose at least 10% of annual NEP<sub>c</sub> every year, failed to show a strong prediction of NEP<sub>c</sub> from cDOY.

#### 4. Discussion

There is increasing evidence that a considerable proportion of the interannual variability of  $NEP_c$  cannot be explained by the current year's climatic variability alone but needs considering previous periods' weather factors. Predispositions of growth by the determination of buds in autumn of the past year [Thomas et al., 2009; Zweifel et al., 2006], carry-over effects on physiology in years following climate extremes [Law et al., 2002; Thomas et al., 2009; Wu et al., 2012; Zielis et al., 2014], C-storage pools accumulated over several years [Campioli et al., 2009; Hoch et al., 2003], sapwood-related hydraulic traits [Zweifel et al., 2006], and winter chilling effects (vernalization) [Cook et al., 2012] are examples of potential causalities between conditions back in time and the current year NEP<sub>c</sub>. In order to better understand the intraannual variability of NEP<sub>c</sub> and its drivers, we introduced the day of compensation (cDOY), i.e., the day of the current year (typically in spring) when net carbon losses during wintertime are compensated by carbon assimilation in spring or early summer (Figure 1). cDOY reflects the complete winter conditions and the related accumulated CO<sub>2</sub> losses, in combination with the onset and rate of  $CO_2$  assimilation in spring (Table 2). Therefore, cDOY is not directly comparable with studies focusing on the onset of GPP or the change in NEP/NEE from a C source to a sink in spring [Delpierre et al., 2009; Gonsamo et al., 2012a; Gonsamo et al., 2012b] since these approaches do not account for the amount of accumulated respiratory C losses over wintertime. In the following we discuss the meaning of cDOY and its impact on the interpretation of NEP<sub>c</sub>.

#### 4.1. Mean Annual Site Temperature Determining cDOY

The loss of C during wintertime and the respective cDOY was found to be statistically highly independent of most of the site characteristics like mean annual precipitation, nitrogen deposition, leaf area index, age, or tree height (Table 3). Only MAT was significantly related to cDOY ( $R^2$  about 0.4, pooled data for all sites, Table 3) particularly when the sites were grouped according to their forest types ( $R^2$  up to 0.99, Figure 3, with one exception, see below).

The importance of air and soil temperatures for the recovery of trees from the inactive physiological winter dormancy back into a physiologically active status is well documented [*Baldocchi et al.*, 2005] and covers issues such as rehydration of tissues [*Koike*, 1990; *Lundmark et al.*, 1988; *Suni et al.*, 2003; *Zweifel et al.*, 2000], bud burst [*Basler and Körner*, 2014], assimilation [*Monson et al.*, 2011a], flowering [*Cook et al.*, 2012], length of the vegetation/growth period [*Aurela et al.*, 2004; *Baldocchi and Wilson*, 2001; *Churkina et al.*, 2005; *Monson et al.*, 2011a], growth [*Zweifel et al.*, 2010], and probably many more. All these processes are, finally, determining cDOY with different weights, since they are influencing quantities and timing of ecosystem respiration and assimilation, explaining the influence of MAT on cDOY well.

#### 4.1.1. One Exception: The Deciduous Broadleaf Forests

There was one exception from the generally close relationship between MAT and cDOY: MAT had no impact on cDOY for deciduous broadleaf forests (DBF, n = 7) (Figure 3), but cDOY had a high explanatory power for NEP<sub>cr</sub> particularly with the dynamic integration method (Table 4e). This finding was unchanged when considering DBF filtering for those sites with a distinct winter respiratory carbon loss of more than 10% of the annual NEP<sub>c</sub> (Figure 3b, negative sign = respiratory loss/positive NEP<sub>c</sub>). Overall, this means that cDOY is strongly forest type specific and that cDOY includes information not covered by the site characteristics investigated and thus offers a new dimension in interpreting NEP<sub>c</sub>. This seems to be particularly true for the DBF sites. The seven DBF forests included in this study (IT-COL, US-MMS, DE-HAI, US-HA1, US-WCR, DK-SOR, and US-UMB) consisted of beech (Fagus sylvatica), maple (Acer spp.), oak (Quercus spp.), ash (Fraxinus spp.), basswood (Tilia americana), and sourwood trees (Oxydendrum arboreum). We suggest two potential explanations why the cDOY of these forests does not depend on MAT. First, (i) the group of DBF sites might still be too heterogeneous in terms of their species composition to show a concise MAT-cDOY relationship. The limited number of replications (n = 7) for this group does, however, not allow for further differentiations. And second, (ii) cDOY reflects processes which are indeed independent of MAT for this forest type, e.g., due to biological predispositions of water and carbon storage which have their origin before the time period investigated [Keenan et al., 2012; Urbanski et al., 2007; Zielis et al., 2014; Zweifel et al., 2010], or due to genetic predispositions which determine the regulation of physiological activity independently of temperature [Basler and Körner, 2014], or in a way that positive and negative temperature effects level each other off. A convincing chain of arguments for the second explanation was recently brought up by Cook et al. [2012]. They showed that increasing temperatures during winter and spring induce opposite effects in certain species. Warmer winter conditions can lead to an insufficient vernalization, i.e., chilling requirements that must be met before a plant is able to respond to spring warming, which in turn leads to a delayed initiation of phenological processes in spring despite the positive effect of increased spring temperatures. Further, for beech trees Basler and Körner [2014] recently reported a codetermination of beech bud burst by the photoperiod and, therefore, a partial decoupling from temperature. Such a partial decoupling from temperature in terms of physiological processes could be, in terms of physiological processes, a species-specific explanation for a predisposition disturbing the generally valid relationship between MAT and cDOY. The effect of climate change on the relationship between cDOY and NEP might thus also depend on species-specific physiological responses and acclimation potentials. It is however difficult to understand how heterotrophic respiration ( $R_{\rm H}$ ) in the soil is triggered by the mentioned tree physiological processes. Apart from temperature, R<sub>H</sub> might be stimulated by rhizosphere processes such as root exudates and mycorrhiza, which in turn might be more closely coupled to the tree physiological status in DBF. Further field studies are needed to test this hypothesis.

#### 4.2. Timing of cDOY

In general, evergreen forests (EBF and ENF) had earlier cDOYs than the deciduous forests, and mixed forests with evergreen and deciduous species were in between (Figure 4). Photosynthesis of evergreens during winter varies with climatic region, but can be substantial. Thus, the early cDOYs of the evergreens may be explained by the ability of evergreen trees to start earlier in the season with assimilation [*Richardson et al.*, 2010] or even maintain it during mild winters [*Pallardy*, 2010]. Photosynthetic capacity can be attained after just a few days of sufficient environmental conditions [*Ottander et al.*, 1995; *Ottander and Oquist*, 1991; *Suni et al.*, 2003].

Forest types, excepting DBF, and cDOY are both found to be linked to MAT (Figure 3 and Table 3). Evergreen broadleaf forest (EBF), for instance, grows at relatively warm sites and do not have a consistently occurring winter respiratory carbon loss period and thus show no consistently cDOY timings in each year (Figure 3). Typical examples are the eucalypt sites in Australia and Portugal (Table 2 and supporting information Figure S AU-TUM and S PT-ESP). These sites show an almost full year growth period or at least do not turn into C sources once every year, and the cDOYs, which can hardly occur, happen thus hardly any year. At the other end of the biological scale appear the deciduous broadleaf tree sites (DBF) with the latest cDOYs (Figure 4) at the generally cooler sites (Figure 3). The existence of a cold season is the main reason for forming a deciduous canopy. Deciduous forests need more time in spring for bud burst and leaf flushing, for the development of the photosynthetic apparatus, and for the onset of photosynthetic activity [*Basler and* 

*Körner*, 2014; *Epron et al.*, 1996; *Jurik*, 1986; *Koike*, 1990; *Reich et al.*, 1991]. The evergreen needleleaf forests (ENF) have the widest temporal range for cDOY (Figure 4), again in line with the widest range of occurring MAT (Figure 3a).

#### 4.3. Strengths and Limitations of cDOY to Predict NEP<sub>c</sub>

The explanatory power of cDOY as a predictor of NEP<sub>c</sub> was strongly depending on whether the site had a net carbon respiratory loss higher than 10% of the annual NEP<sub>c</sub> or not (Figures 4 and 5). For sites with a distinct net carbon loss over wintertime (Figure 5a) the estimated annual NEP<sub>c</sub> from cDOY reached accuracies of  $\pm$ 75 g C m<sup>-2</sup> yr<sup>-1</sup> which are comparable to some of the most successful (but much more complex) NEP models [*Keenan et al.*, 2012]. For the other sites without a distinct winter respiratory loss, the standard deviation between modeled and measured NEP<sub>c</sub> was a factor 2 to 3 higher (Figure 5b), which leads to the conclusion that cDOY is of limited explanatory power in these cases. This could be explained partly by the large variation in winter photosynthesis in temperate evergreens, and by the fact that evergreen needleleaf species grow in some of the harshest conditions, such as the western U.S. where summer drought is the norm [*Law and Waring*, 2015].

Besides the importance of the winter net respiratory C loss, the forest type had a strong influence on the predictive power of cDOY on NEP<sub>c</sub>. Pooled data reached an  $R^2$  of 0.47 (dynamic integration) for the linear regression between cDOY and NEP<sub>c</sub> (Table 4), whereas the grouped data for ENF ( $R^2 = 0.58$ , dynamic integration, Table 4e) and DBF ( $R^2 = 0.68$ , dynamic integration, Table 4d) were much higher. This again indicates that the information content of cDOY, i.e., the net effect of winter and spring processes, depends on the forest type and the respective species composition (Figure 4). Both winter respiratory loss and vegetation type are related to temperature and therefore linked to each other (Figure 3). It is therefore not surprising that besides cDOY as the variable with the highest explanatory power for NEP<sub>c</sub>, mean annual temperature appeared as the second best driver in our stepwise multiple regression analyses (Table 4). The addition of other site factors, namely precipitation, age, or LAI improved the multiple regressions further. Generally, the goodness of fit between cDOY and NEP<sub>c</sub> increased with the timing of later cDOYs and with decreasing air temperatures (Table 4).

We conclude that lower mean annual temperatures lead to generally more pronounced winter net respiratory losses and it appears plausible that this is linked to later cDOYs. This is also in line with studies analyzing the onset of forests as a C sink in relation to winter and spring temperatures [*Baldocchi et al.*, 2005; *Cook et al.*, 2012; *Delpierre et al.*, 2009; *Monson et al.*, 2011b]. Or the other way around, the warmer the site the less distinct the carbon loss period may be the earlier cDOY happens and the less likely the influence of cDOY on annual carbon uptake. Furthermore, we conclude that latter cDOYs are linked to lower annual NEP<sub>c</sub>, and thus, the influence of cDOY on the annual NEP<sub>c</sub> increases with its timing.

cDOYs of deciduous broadleaf forests (DBF) showed the highest prediction quality for NEP<sub>c</sub> (Table 4e) despite the fact that the respective cDOY did not correlate with mean annual temperature (Figure 3) nor other site variables like for other forests types (Table 4d) or the pooled data (Tables 4a–4c). Sites at higher altitudes (e.g., US-NR1 and US-Me2) experience large interannual variation in the physiological active period, for example, 45 days at US-Me2 [*Thomas et al.*, 2009], and studies in the mountains of the western U.S. have shown declining snowpack for decades and its correlation with warm temperature anomalies. Further at US-NR1, longer growing seasons were correlated with low snow water equivalent and resulted in less annual net carbon uptake [*Hu et al.*, 2010]. Overall, such processes may confound an explanatory power of MAT for cDOY and NEP<sub>c</sub> in certain cases; however, we found no generally convincing explanation for the relationship between cDOY (its not found drivers) and NEP<sub>c</sub>. Even when not understanding why DBF sites appear as a special case, we conclude that cDOY timing must in general depend on variables (eventually beyond the ones we analyzed) containing information about the site and its past (climatic) history, including genetic predispositions leading to this high predictive power for NEP<sub>c</sub>.

In summary, there are many indications for winter effects on NEP<sub>c</sub> of forests and related to it on the cDOY timing. The compensation day (cDOY) is suggested to capture air temperature and intrinsic forest type-dependent differences, leading to a specific date in the first part of the calendar year, with a high explanatory power for the upcoming annual NEP<sub>c</sub> values of the entire year for forest sites under distinct respiratory net carbon losses during wintertime.

#### 4.4. Starting the NEP<sub>c</sub> Year in Autumn

Three different ways of integrating NEP over a year were applied: the static "classical" calendar-year method (1 January to 31 December), the static "Urbanski" method (28 October to 27 October), and the more processoriented "dynamic" method, defining the "biological" year as the period between two annual NEP<sub>c</sub> peaks. There appeared distinctly better fits between cDOY and NEP<sub>c</sub> for the two methods starting the NEP<sub>c</sub> year in autumn (Table 4). The classical method performed generally worse for all types of analyses (Tables 3 and 4 and Figure 5). The additional gain of predictive quality for the dynamic method over the static Urbanski method was relatively small. This means that it is important to include the complete autumn and winter period before the actual C sink period for interpreting NEP<sub>c</sub>, but doing so with a static approach captures more or less the same information as when doing so with the site- and year-specific dynamic method (which can be more labor intensive to deal with).

#### 4.5. Conclusions

The compensation day cDOY reflects processes, which take place before the net C-sink period begins in forests in spring and early summer. The fact that cDOY explains more of NEP<sub>c</sub> when starting the NEP<sub>c</sub> year in autumn shows that the (autumn-winter) period already before 1 January plays an important role for the following NEP<sub>c</sub> performance. cDOY analysis takes seasonal and interannual variations of the carbon cycle dynamics into account and is therefore suggested to take up carry-over effects of climate and carbon storage in temperate forests [*Keenan et al.*, 2012; *Urbanski et al.*, 2007; *Zielis et al.*, 2014; *Zweifel et al.*, 2010]. Such carry-over effects seem to be less important in forests with no distinct winter net respiratory loss of C (C loss less than 10% of annual NEP<sub>c</sub>). This is in line with the finding that cDOY gains explanatory power for NEP<sub>c</sub> at sites with distinct winter respiratory C losses. The fact that biological processes, occurring before the annual net assimilation period begins, are able to explain more than 50% of the annual NEP<sub>c</sub> should change our view on the drivers of NEP<sub>c</sub>. Thus, an accurate NEP<sub>c</sub> interpretation additionally needs to include the conditions that affected a forest before this period.

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