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# Recommended Citation

Watts, J. D., Kimball, J. S., Parmentier, F. J. W., Sachs, T., Rinne, J., Zona, D., Oechel, W., Tagesson, T., Jackowicz-Korczyński, M., and Aurela, M.: A satellite data driven biophysical modeling approach for estimating northern peatland and tundra CO2 and CH4 fluxes, Biogeosciences, 11, 1961-1980, doi:10.5194/bg-11-1961-2014, 2014

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Biogeosciences, 11, 1961-1980, 2014 [www.biogeosciences.net/ll/1961/2014/](http://www.biogeosciences.net/ll/1961/2014/)  doi:10.5194/bg-ll-1961-2014 © Author(s) 2014. CC Attribution 3.0 License.





# **A satellite data driven biophysical modeling approach for estimating northern peatland and tundra CO<sub>2</sub> and CH<sub>4</sub> fluxes**

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Received: 5 September 2013 - Published in Biogeosciences Discuss.: 25 October 2013 Revised: 25 Febraary 2014 - Accepted: 26 Febmary 2014 - Pnblished: 9 April 2014

**Abstract.** The northern terrestrial net ecosystem carbon balance (NECB) is contingent on inputs from vegetation gross primary prodnctivity (GPP) to offset the ecosystem respiration ( $R_{\text{eco}}$ ) of carbon dioxide (CO<sub>2</sub>) and methane (CH<sub>4</sub>) emissions, but an effective framework to monitor the regional Arctic NECB is lacking. We modified a terrestrial carbon flux (TCF) model developed for satellite remote sensing applications to evaluate wetland  $CO<sub>2</sub>$  and  $CH<sub>4</sub>$  fluxes over pan-Arctic eddy covariance (EC) flux tower sites. The TCF model estimates GPP,  $CO<sub>2</sub>$  and CH<sub>4</sub> emissions using in situ or remote sensing and reanalysis-based climate data as inputs. The TCF model simulations using in situ data explained  $> 70\%$  of the  $r^2$  variability in the 8 day cumulative EC measured fluxes. Model simulations using coarser satellite (MODIS) and reanalysis (MERRA) records acconnted for approximately 69 % and 75 % of the respective  $r^2$  variability in the tower CO<sub>2</sub> and CH<sub>4</sub> records, with corresponding RMSE uncertainties of  $\leq 1.3$  g C m<sup>-2</sup> d<sup>-1</sup> (CO<sub>2</sub>) and  $18.2 \text{ mg C m}^{-2} \text{ d}^{-1}$  (CH<sub>4</sub>). Although the estimated annual CH<sub>4</sub> emissions were small (<  $18 \text{ g C m}^{-2} \text{ yr}^{-1}$ ) relative to  $R_{\text{eco}}$  (> 180 g C m<sup>-2</sup> yr<sup>-1</sup>), they reduced the across-site NECB by 23 % and contributed to a global warming potential of approximately  $165 \pm 128$  g CO<sub>2</sub>eq m<sup>-2</sup> yr<sup>-1</sup> when considered over a 100 year time span. This model evalnation indicates a strong potential for nsing the TCF model approach to document landscape-scale variability in  $CO<sub>2</sub>$  and  $CH<sub>4</sub>$  fluxes, and to estimate the NECB for northern peatland and tundra ecosystems.

# **1 Introduction**

Northern peatland and tundra ecosystems are important components of the terrestrial carbon cycle and store over half of the global soil orgaruc carbon reservoir in seasonally frozen and permafrost soils (Hngelins et al., 2013). However, these systems are becoming increasingly vulnerable to carbon losses as  $CO<sub>2</sub>$  and  $CH<sub>4</sub>$  emissions, resulting from climate warming and changes in the terrestrial water balance (Kane et al., 2012; Kim et al., 2012) that can increase soil carbon decomposition. Recent net  $CO<sub>2</sub>$  exchange in northern tundra and peatland ecosystems varies from a sink of  $291 \text{ Tg} \text{Cyr} ^{-1}$  to a source of  $80 \text{ Tg} \text{Cyr} ^{-1}$ , when considering the substantial uncertainty in regional estimates using scaled flux observations, atmospheric inversions, and ecosystem process models (McGuire et al., 2012). The magnitude of a carbon sink largely depends on the balance between carbon nptake by vegetation prodnctivity and losses from soil mineralization and respiration processes. High-latitude warming can increase ecosystem carbon uptake by reducing cold-temperature constraints on plant carbon assimilation and growth (Hudson et al., 2011; Elmendorf et al., 2012). Soil warming also accelerates carbon losses due to the exponential effects of temperature on soil respiration, whereas wet and inundated conditions shift microbial activity towards anaerobic consumption pathways that are relatively slow but can result in substantial CH4 production (Moosavi and Crill, 1997; Merbold et al., 2009). Regional wetting across the Arctic (Watts et al., 2012; Zhang et al., 2012a) may increase  $CH<sub>4</sub>$  emissions, which have a radiative warming potential at least 25 times more potent than  $CO<sub>2</sub>$  per unit mass over a 100 year time horizon (Boucher et al., 2009). The northem latitudes already contain over 50 % of global wetlands and recent increases in atmospheric CH4 concentrations have been attributed to heightened gas emissions in these areas during periods of warming (Dlugokencky et al., 2009; Dolman et al., 2010). Northern peatland and tundra ( $> 50^{\circ}$  N) reportedly contribute between 8 and 79 Tg C in CH<sub>4</sub> emissions each year, but these fluxes have been difficult to constrain due to uncertainty in the parameterization of biogeochemical models, the regional characterization of wetland extent and water table depth, and a scarcity of ecosystem-scale  $CH<sub>4</sub>$  emission observations (Petrescu et al., 2010; Riley et al., 2011; Spahni et al., 2011; McGuire et al., 2012; Meng et al., 2012).

Ecosystem studies using chamber and tower eddy covariance (EC) methods continue to provide direct measurements of  $CO<sub>2</sub>$  and  $CH<sub>4</sub>$  fluxes and add valuable insight into the enviromnental constraints on these processes. However, extrapolating localized carbon fluxes to regional scales has proven difficult and is severely constrained by the limited number of in situ observations and the large spatial extent and heterogeneity of peatland and tundra ecosystems. Recent approaches have used satellite-based land cover classifications, photosynthetic leaf area maps, or wetness indices to "upscale"  $CO<sub>2</sub>$  (Forbrich et al., 2011; Marushchak et al., 2013) andCH4 (Tagessonetal., 2013; Sturtevant and Oechel, 2013) flux measurements. Remote sensing inputs have also been used in conjunction with biophysical process modeling to estimate landscape-level changes in plant carbon assimilation and soil CO<sub>2</sub> emissions (Yuan et al., 2011; Tagesson et al., 2012a; Yi et al., 2013). Previous analyses of regional CH<sup>4</sup> contributions have ranged from the relatively simple modification of CH4 emission rate estimates for wetland fractions according to temperature and carbon substrate constraints (Potter et al., 2006; Clark et al., 2011) to the use of more complex multi-layer wetland CH4 models with integrated hydrological components (McGuire et al., 2012; Wania et al., 2013). Yet, most investigations have not examined the potential for simultaneously assessing  $CO<sub>2</sub>$  and  $CH<sub>4</sub>$  fluxes, and the corresponding net ecosystem carbon balance (Sitch et al., 2007; McGuire et al., 2012; Olefeldt et al., 2012) for peatland and tundra using a satellite remote sensing-based model approach.

It is well recognized that sub-surface conditions influence the land-atmosphere exchange of  $CO<sub>2</sub>$  and  $CH<sub>4</sub>$  production. However, near-surface soil temperature, moisture and carbon substrate availability play a cracial role in regulating ecosystem carbon emissions. Strong associations between surface soil temperature ( $\leq 10$  cm depth) and CO<sub>2</sub> respiration have been observed in Arctic peatland and tundra permafrost systems (Kutzbach et al., 2007). Significant relationships between CH4 emissions and temperature have also been reported (Hargreaves et al., 2001; Zona et al., 2009; Sachs et al., 2010). Although warming generally increases the decomposition of organic carbon, the magnitude of  $CO<sub>2</sub>$  production is constrained by wet soil conditions (Olivas et al., 2010) that instead favor  $CH_4$  emissions and decrease methantrophy in soil and litter layers (Turetsky et al., 2008; Olefeldt et al., 2012). Oxidation by methanotrophic communities in surface soils can reduce  $CH_4$  emissions by over 90 % when gas transport occurs through diffusion (Preuss et al., 2013), but this constraint is often minimized when pore water content rises above 55-65 % (von Fischer and Hedin, 2007; Sjogersten and Wookey, 2009). Despite increases in the availability of organic carbon and accelerated  $CO<sub>2</sub>$  release due to soil warming and thickening of the active layer in permafrost soils (Dorrepaal et al., 2009), anaerobic communities have shown a preference for light-carbon fractions (e.g., amines, carbonic acids) that are more abundant in the upper soil horizons (Wagner et al., 2009). Similarly, labile carbon substrates from recent photosynthates and root exudates have been observed to increase CH<sub>4</sub> production relative to heavier organic carbon fractions (Ström et al., 2003; Dijkstra et al., 2012; Olefeldt et al., 2013) that require longer decomposition pathways to break down complex molecules into the simple compounds (i.e., acetate,  $H_2 + CO_2$ ) used in methanogenesis (Le Mer and Roger, 2001).

The objective of this study was to evaluate the feasibility of using a satellite remote sensing data driven modeling approach to assess the daily and seasonal variability in  $CO<sub>2</sub>$ and CH4 fluxes from northem peatland and tundra ecosystems, according to near-surface enviromnental controls including soil temperature, moisture and available soil organic carbon. In this paper we incorporate a newly developed  $CH<sub>4</sub>$ emissions algorithm within an existing terrestrial carbon flux (TCF) CO2 model framework (Kimball et al., 2012; Yi et al.,  $2013$ ). The CH<sub>4</sub> emissions algorithm simulates gas production using near-surface temperature, anaerobic soil fractions and labile organic carbon as inputs. Plant  $CH<sub>4</sub>$  transport is determined by vegetation growth characteristics derived from gross primary production (GPP), plant functional traits and canopy/surface turbulence. Methane diffusion is determined based on temperature and moisture constraints to gas movement through the soil column, and oxidation potential. Ebullition of CH4 is assessed using a simple gradient method (van Huissteden et al., 2006).

The integrated TCF model allows for satellite remote sensing information to be used as primary inputs, requires minimal parameterization relative to more complex ecosystem process models, and provides a framework to monitor the terrestrial net ecosystem carbon balance (NECB). Although the NECB also encompasses other mechanisms of carbon transport, including dissolved and volatile organic carbon emissions and fire-based particulates, the NECB is limited in this study to  $CO<sub>2</sub>$  and  $CH<sub>4</sub>$  fluxes, which often are primary contributors in high-latitude tundra and peatland ecosystems (McGuire et al., 2010).

To evaluate the combined  $CO<sub>2</sub>$  and  $CH<sub>4</sub>$  algorithm approach, we compared TCF model simulations to tower EC records from six northem peatland and tundra sites within North America and Eurasia. For this study, baseline simulations driven with tower EC-based GPP and in situ meteorology data were first used to assess the capability of the TCF model approach to quantify temporal changes in landscapescale carbon ( $CH_4$  and  $CO_2$ ) fluxes. Secondly,  $CO_2$  and  $CH_4$ simulations using internal TCF model GPP estimates (Yi et al., 2013) and inputs from satellite and global model reanalysis records were used to evaluate the relative uncertainty introduced when using coarser scale information in place of in situ data. These satellite and reanalysis driven simulations were then used to determine the annual  $CO<sub>2</sub>$  and  $CH<sub>4</sub>$  fluxes at the six tower sites, and the relative impact of  $CH<sub>4</sub>$  emissions on the NECB.

#### **2 Methods**

#### **2.1 TCF model deseription**

The combined TCF model  $CO<sub>2</sub>$  and  $CH<sub>4</sub>$  framework regulates carbon gas exchange using soil surface temperature, moisture and soil orgaruc carbon availability as inputs, and has the flexibility to run simulations at local and regional scales. TCF model estimates of ecosystem respiration *(Reco)* and net ecosystem  $CO<sub>2</sub>$  exchange (NEE) have been evaluated against tower EC data sets from boreal and timdra systems using GPP, surface  $(< 10 \text{ cm}$  depth) soil temperature  $(T_s)$  and volumetric moisture content  $(\theta)$  inputs available from global model reanalysis and satellite remote sensing records (Kimball et al., 2009; McGuire et al., 2012). A recent adjustment to the TCF model (Kimball et al., 2012; Yi et al., 2013) incorporates a light-use efficiency (LUE) algorithm that provides internally derived GPP calculations to determine *Reco* and NEE fluxes at a daily time step. The adjusted  $TCF CO<sub>2</sub>$ model also allows for better user control over parameter settings and surface meteorological inputs (Kimball et al., 2012). The  $CO<sub>2</sub>$  and newly added CH<sub>4</sub> flux model components are described in the following sections. A summary of the TCF model inputs, parameters, and the associated parameter values used in this study are provided in the Supplement (tables SI and S2; Fig. SI).

# **<sup>2</sup> .1 .1 CO2 flux component**

The internal TCF model GPP algorithm estimates daily fluxes according to a biome-dependent vegetation maximum LUE coefficient ( $\varepsilon_{\text{max}}$ ; mg CMJ<sup>-1</sup>) that represents the optimal conversion of absorbed solar energy and CO<sub>2</sub> to plant organic carbon through photosynthesis (Kimball et al., 2012). To account for daily minimum air temperature  $(T_{min})$  and atmospheric vapor pressure deficit (VPD) constraints on photosynthesis (Running et al., 2004),  $\varepsilon_{\text{max}}$  is reduced ( $\varepsilon$ ) using dimensionless linear rate scalars ranging from 0 (total inhibition) to 1 (no inhibition) that are described elsewhere (i.e., Kimball et al., 2012; Yi et al., 2013). In this study we also account for the sensitivity of shallow rooted vegetation and bryophytes, which lack vascular tissues for water transport, to changes in surface volumetric soil water (Wu et al., 2013), where  $\theta_{\min}$  and  $\theta_{\max}$  are the specified minimum and maximum parameter values:

$$
\varepsilon = \varepsilon_{\text{max}} \times f(\text{VPD}) \times f(T_{\text{min}}) \times f(\theta)
$$
  
where  $f(\theta) = (\theta - \theta_{\text{min}})/(\theta_{\text{max}} - \theta_{\text{min}}).$  (1)

Simulated GPP (g C m<sup> $-2$ </sup> d<sup> $-1$ </sup>) is obtained as

$$
GPP = \varepsilon \times 0.45 \text{SW}_{\text{rad}} \times \text{FPAR},\tag{2}
$$

where  $SW_{rad}$  (W m<sup>-2</sup>) is incoming shortwave radiation and FPAR is the fraction of daily photosynthetically active solar radiation (PAR; MJ  $m^{-2}$ ) absorbed by plants during photosynthesis. For this approach, PAR is assumed to be 45 % of SWrad (Zhao et al., 2005). Remotely sensed normalized difference vegetation index (NDVl) records have been used to estimate vegetation productivity (Schubert et al., 2010a; Parmentier et al., 2013) and changes in growing season length (Beck and Goetz, 2011) across northern peatland and tundra environments. Daily FPAR is derived using the approach of Badawy et al. (2013) to mitigate potential biases in low biomass landscapes (Peng et al., 2012):

$$
FPAR = \frac{0.94(\text{Index} - \text{Index}_{\text{min}})}{\text{Index}_{\text{range}}}.
$$
 (3)

This approach uses NDVl or simple ratio (SR; i.e.,  $(1 + NDVI)/(1 - NDVI)$  indices as input index values. The results are then averaged to obtain FPAR. Index<sub>range</sub> corresponds to the difference between the 2nd and 98th percentiles in the NDVl and SR distributions (Badawy et al., 2013).

Biome-specific autotrophic respiration  $(R_a)$  is estimated using a carbon use efficiency (CUE) approach that considers the ratio of net primary production (NPP) to GPP (Choudhury, 2000). Carbon loss from heterotrophic respiration  $(R<sub>h</sub>)$ is determined using a three-pool soil litter decomposition scheme consisting of metabolic ( $C_{\text{met}}$ ), structural ( $C_{\text{str}}$ ) and recalcitrant (Creo) organic carbon pools with variable decomposition rates. The  $C<sub>met</sub>$  pool represents easily decomposable plant residue and root exudates including amino acids, sugars and simple polysaccharides, whereas the  $C_{str}$  pool consists of litter residues such as hemi-cellnlose and lignin (Ise et al., 2008; Porter et al., 2010). The *Crec* pool includes physically and chemically stabilized carbon derived from the  $C_{\text{met}}$  and  $C_{\text{str}}$  pools and also corresponds to humified peat. A fraction of daily NPP  $(F_{\text{met}})$  is first allocated as readily decomposable litterfall to  $C_{\text{met}}$  and the remaining portion  $(1 - F_{\text{met}})$  is transferred to  $C_{str}$  (Ise and Moorcroft, 2006; Kimball et al., 2009). To account for reduced mineralization in tundra and peatland environments, approximately 70% of  $C_{str}$  ( $F_{str}$ ) is reallocated to  $C_{\text{rec}}$  (Ise and Moorcroft, 2006; Ise et al., 2008):

$$
dC_{\text{met}}/dt = \text{NPP} \times F_{\text{met}} - R_{\text{h,met}} \tag{4}
$$

$$
dC_{str}/dt = NPP (1 - F_{met}) - (F_{str} \times C_{str}) - R_{h,str}
$$
 (5)

$$
dC_{\text{rec}}/dt = (F_{\text{str}} \times C_{\text{str}}) - R_{\text{h,rec}}.
$$
 (6)

Daily  $CO_2$  loss from the  $C_{\text{met}}$  pool (i.e.,  $R_{h, \text{met}}$ ) is determined as the product of  $C_{\text{met}}$  and an optimal decomposition rate parameter  $(K_p)$ . The realized decomposition rate  $(K_{\text{met}})$ results from the attenuation of  $K_p$  by dimensionless  $T_s$  and  $\theta$ multipliers ( $T_{\text{mult}}$  and  $W_{\text{mult}}$ , respectively), that vary between <sup>0</sup> (fully constrained) and 1 (no constraint):

$$
K_{\text{met}} = K_{\text{p}} \times T_{\text{mult}} \times W_{\text{mult}} \tag{7}
$$

$$
T_{\text{mult}} = \exp\left[308.56\left(66.02^{-1} - (T_s + T_{\text{ref}} - 66.17)^{-1}\right)\right]
$$
 (8)

$$
W_{\text{mult}} = 1 - 2.2(\theta - \theta_{\text{opt}})^2
$$
\n(9)

The temperature constraints are imposed using an Arrheniustype function (Lloyd and Taylor, 1994; Kimball et al., 2009) where decomposition is no longer limited when average daily  $T_s$  exceeds a user-specified reference temperature ( $T_{ref}$ ; in K) that can vary with carbon substrate complexity, physical protection, oxygen availability and water stress (Davidson and Janssens, 2006). The *Wmuit* modifier accounts for the inhibitory effect of dry and near-saturated soil moisture conditions on heterotrophic decomposition (Oberbaner et al., 1996). For this study,  $\theta_{\text{opt}}$  is set to 80% of pore saturation to account for ecosystem adaptations to wet soil conditions (Ise et al., 2008; Zona et al., 2012) and near-surface oxygen availability provided by plant root transport (Elberling et al., 2011). Decomposition rates for  $C_{str}$  and  $C_{rec}$  ( $K_{str}$ ,  $K_{rec}$ ) are determined as 40% and 1% of  $K_{\text{met}}$ , respectively (Kimball et al., 2009), and  $R<sub>h</sub>$  is the total  $CO<sub>2</sub>$  loss from the three soil organic carbon pools:

$$
R_{\rm h} = K_{\rm met} \times C_{\rm met} + K_{\rm str} \times C_{\rm str} + K_{\rm rec} \times C_{\rm rec}.
$$
 (10)

Finally, the TCF model estimates NEE (g C m<sup> $-2$ </sup> d<sup>-1</sup>) as the residual difference between  $R_{\text{eco}}$ , which includes  $R_{\text{a}}$  and  $R_{\text{h}}$ respiration components, and GPP. Negative  $(-)$  and positive (+) NEE fluxes denote respective terrestrial  $CO<sub>2</sub>$  sink and source activity:

$$
NEE = (R_a + R_h) - GPP.
$$
 (11)

### **2.1.2 CH4 flux component**

A CH4 emissions algorithm was incorporated within the TCF model to estimate  $CH<sub>4</sub>$  fluxes for peatland and tundra landscapes. The model estimates  $CH<sub>4</sub>$  production according to  $T_s$ ,  $\theta$ , and labile carbon availability. Plant CH<sub>4</sub> transport is modified by vegetation growth and production, plant functional traits, and canopy aerodynamic conductance that takes into account the influence of wind turbulence on moisture/gas flux between vegetation and the atmosphere. The CH4 module is similar to other process models (e.g., Walter and Heimann, 2000; van Huissteden et al., 2006), but reduces to a one-dimensional near-snrface soil profile following Tian et al.  $(2010)$  to simplify model parameterization amenable to remote sensing applications. For the purposes of this study, the soil profile is defined for surface  $(< 10 \text{ cm}$  depth) soil layers as most temperature and moisture retrievals from satellite remote sensing do not characterize deeper soil conditions. Althongh this approach may not account for variability in carbon fluxes associated with deeper soil constraints, field studies from high-latitnde ecosystems have reported strong associations between CH<sub>4</sub> emissions and near-surface conditions including  $T_s$  and soil moisture (Hargreaves et al., 2001; Sachs et al., 2010; von Fischer et al., 2010; Sturtevant et al., 2012; Tagesson etal., 2012b).

#### **CH4 production**

Soil moisture in the upper rhizosphere is a fundamental control on CH4 production and emissions into the atmosphere. Methanogenesis ( $R_{\text{CH}_4}$ ) within the saturated soil pore volume  $(\varphi_s, m^{-3})$ ; the aerated pore volume is denoted as  $\varphi_a$ ) is determined according to an optimal CH4 production rate *(Ro ',*  $\mu$ M CH<sub>4</sub> d<sup>-1</sup>) and labile photosynthates:

$$
R_{\rm CH_4} = (R_0 \times \phi_s) \times C_{\rm met} \times Q_{10p}^{(T_s - T_p)/10}.
$$
 (12)

For this study,  $CH_4$  production was driven using the soil *Cmet* pool to reflect contributions by lower weight carbon substrates (Reiche et al., 2010; Corbett et al., 2013) in labile organic carbon-rich environments. Carbon from the  $C_{str}$ pathway may also be allocated for  $CH_4$  production in ecosystems with lower labile organic carbon inpnts and higher contributions by hydrogenotrophic methanogenesis (Alstad and Whiticar, 2011). The  $Q_{10p}$  temperature modifier is used as an approximation to the Arrhenius equation and describes the temperatme dependence of biological processes (Gedney and Cox, 2003; van Huissteden et al., 2006). The reference temperature  $(T_p)$  typically reflects mean annual or non-frozen season climatology. Both  $Q_{10p}$  and  $T_p$  can be adjusted, in addition to  $R<sub>o</sub>$ , to accommodate varying temperature sensitivities in response to ecosystem differences in substrate quality and other environmental conditions (van Hulzen et al., 1999; Inglett et al., 2012). Methane additions from  $R_{\text{CH}_4}$  are first allocated to a temporary soil storage pool  $(C<sub>CH4</sub>)$  prior to determining the CH<sub>4</sub> emissions for each 24 h time step;  $C_{\text{met}}$ 

is also updated to account for carbon losses due to  $CH<sub>4</sub>$  production.

#### **CH4 emission**

The magnitude of daily CH<sub>4</sub> emissions ( $F_{\text{CH}_4}$ ) from the soil profile is determined through plant transport  $(F_{\text{plant}})$ , soil diffusion ( $F_{\text{diff}}$ ) and ebullition ( $F_{\text{ebull}}$ ) pathways:

$$
F_{\rm CH_4} = F_{\rm plant} + F_{\rm diff} + F_{\rm cbull}.
$$
\n(13)

Vegetation plays an important role in terrestrial  $CH<sub>4</sub>$  emissions by allowing for gas transport through the plant stracture, avoiding slower diffusion through the soil column and often reducing the degree of CH<sub>4</sub> oxidation (Joabsson et al., 1999). Daily  $F_{\text{plant}}$  is determined using a rate constant  $(C_p)$ modified by vegetation growth and production  $(f_{\text{grow}})$ , and an aerodynamic term ( $\lambda$ ) and a rate scalar ( $P_{trans}$ ) that account for differences in  $CH<sub>4</sub>$  transport ability according to plant functional type:

$$
F_{\text{plant}} = (C_{\text{CH}_4} \times C_p \times f_{\text{grow}} \times \lambda \times P_{\text{trans}})(1 - P_{\text{ox}}). \tag{14}
$$

A fraction of  $F_{\text{plant}}$  is oxidized ( $P_{\text{ox}}$ ) prior to reaching the atmosphere and can be modified according to plant functional characteristics (Frenzel and Rudolph, 1998; Strom et at., 2005; Kip et al., 2010). Plant transport is further reduced under frozen surface conditions to account for pathway obstraction by ice and snow or bending of the plant stem following senescence (Hargreaves et al., 2001; Sun et al., 2012). The magnitude of  $f_{\text{grow}}$  is determined as the ratio of daily GPP to its annual maximum and is used to account for seasonal differences in root and above-ground biomass (Chanton, 2005).

Aerodynamic conductance  $(g_a)$  represents the influence of near-surface turbulence on energy/moisture fluxes between vegetation and the atmosphere (Roberts, 2000; Yan et al., 2012) and gas transport within the plant body (Grosse et al., 1996; Sachs et al., 2008; Wegner et al., 2010; Sturtevant et al.,  $2012$ :

$$
g_{\rm a} = \frac{k^2 \mu_m}{\ln[(z_m - d)/z_{\rm om}] \ln[(z_m - d)/z_{\rm ov}]}.
$$
 (15)

Values for  $z_m$  and  $d$  are the respective anemometer and zero plane displacement heights (m);  $z_{\text{om}}$  and  $z_{\text{ov}}$  are the corresponding roughness lengths (m) for momentum, heat and vapor transfer. The von Karman constant  $(k; 0.40)$  is a dimensionless constant in the logarithmic wind velocity profile (Högström, 1988),  $\mu_m$  is average daily wind velocity (m  $s^{-1}$ ), *d* is calculated as 2/3 of the vegetation canopy height,  $z<sub>om</sub>$  is roughly 1/8 of canopy height (Yang and Friedl, 2002), and  $z_{\text{ov}}$  is 0.1  $z_{\text{om}}$  (Yan et al., 2012). The estimated  $g_{\text{a}}$  is then scaled between 0 and 1 to obtain  $\lambda$  using a linear function for sites with a lower observed sensitivity to surface turbulence; for environments with a higher sensitivity to surface turbulence, a quadratic approach is used when  $\mu_m$  exceeds  $4 \text{ m s}^{-1}$ :

$$
\lambda = 0.0246 + 0.5091g_{a}
$$
  

$$
\lambda = 0.0885 - (3.28g_{a}) + (44.51g_{a}^{2}), \mu_{m} > 4 \,\text{ms}^{-1}.
$$
 (16)

Although this approach focuses on the influence of wind turbnlence on plant gas transport within vegetated wetlands, it is also applicable for inundated microsites where increases in surface water mixing can stimulate CH4 degassing (Sachs et al., 2010). In addition, Eq. (15) reflects near-neutral atmospheric stability and adjustments may be necessary to accommodate unstable or stable atmospheric conditions (Ranpach, 1998).

The upward diffusion of  $CH_4$  within the soil profile is determined using a one-layer approach similar to Tian et al. (2010). The rate of CH<sub>4</sub> transport ( $D_e$ ; m<sup>-2</sup> d<sup>-1</sup>) is considered for both saturated ( $D_{\text{water}}$ ;  $1.73 \times 10^{-4}$   $\mu$ M CH<sub>4</sub> d<sup>-1</sup>) and aerated ( $D_{\text{air}}$ ; 1.73 µM CH<sub>4</sub> d<sup>-1</sup>) soil fractions:

$$
D_e = (D_{\text{water}} \times \phi_s)(D_{\text{air}} \times \phi_a). \tag{17}
$$

Potential daily transport through diffusion ( $P_{diff}$ ) is estimated as the product of  $D_e$  and the gradient between  $C_{\text{CH}_4}$  and the concentration of  $CH_4$  in the atmosphere (Air $_{\text{CH}_4}$ ). This is further modified by soil tortuosity ( $\tau$ ; 0.66), which increases exponentially for  $T_s < 274$  K to account for slower gas movement at colder temperatures and barriers to diffusion resulting from near-surface ice formation (Walter and Heimann, 2000; Zhnang et al., 2004), and pathway constraints within the saturated pore fraction  $(1 - \theta)$ :

$$
P_{\text{diff}} = \tau \times D_e (C_{\text{CH}_4} - \text{Air}_{\text{CH}_4}) (1 - \theta)
$$
  
\n
$$
T_s \ge 274, \quad \tau = 0.66
$$
  
\n
$$
T_s < 274, \quad \tau = 0.05 + 10^{-238} \times T_s^{97.2}.
$$
\n(18)

A portion of diffused CH<sub>4</sub> is oxidized  $(R_{ox})$  before reaching the soil surface, nsing a Michaelis-Menten kinetics approach that is scaled by  $\phi_a$ :

$$
R_{\text{ox}} = \frac{(V_{\text{max}} \times \phi_{\text{a}}) P_{\text{diff}}}{(K_m + \phi_{\text{a}}) P_{\text{diff}}} \times Q_{10d}^{(T_s - T_d)/10},\tag{19}
$$

where  $V_{\text{max}}$  is the maximum reaction rate and  $K_m$  is the substrate concentration at 0.5  $V_{\text{max}}$  (van Huissteden et al., 2006). Oxidation during soil diffusion is modified by soil temperature  $Q_{10}$  constraints ( $Q_{10d}$ );  $T_d$  is the reference temperature and can be defined using site-specific mean annual  $T<sub>s</sub>$  (Le Mer and Roger, 2001). Total daily CH<sub>4</sub> emission ( $F_{diff}$ ) from the soil diffusion pathway is determined by substracting  $R_{\text{ox}}$ from  $P_{\text{diff}}$ .

The CH4 algorithm uses a gradient-based approach to account for slow or "steady-rate" ebullition from inundated micro-sites in the landscape (Rosenberry et al., 2006; Wania et al., 2010), whereas episodic events originating deeper within the soil require more complex modeling techniques and input data requirements (Kettridge et al, 2011) that are beyond the scope of this study. Emission contributions due to ebullition occur when  $C_{\text{CH}_4}$  exceeds a threshold value  $(v_e)$  of 500 pM (van Huissteden et al, 2006). The magnitude of gas release is determined by steady-rate bubbling  $(C_e)$  applied within the saturated soil pore space  $(\varphi_s)$ :

$$
F_{\text{ebull}} = (C_e \times \phi_s) (C_{\text{CH}_4} - v_e), C_{\text{CH}_4} > v_e.
$$
 (20)

#### **2.2 Study sites and in situ data records**

Tower EC records from six pan-Arctic peatland and tundra sites in Finland, Sweden, Russia, Greenland and Alaska were used to assess the integrated TCF model  $CO<sub>2</sub>$  and  $CH<sub>4</sub>$ simulations (Fig. 1; Table 1). The Scandinavian tower sites include Siikaneva (SK) in southern Finland and Stordalen Mire (SM) in northem Sweden near the Abisko Scientific Research Station. The Lena River (LR) delta site is located on Samoylov Island in northem Siberia and EC measurements from the Kytalyk (KY) fiux tower were collected near Chokurdakh in northeastem Siberia. The Zackenberg (ZK) fiux tower is located within the Northeast Greenland National Park, and tower data records for Alaska were obtained from a water table manipulation experiment (Zona et al., 2009; 2012; Sturtevant et al., 2012) approximately 6 km east of Barrow (BA). With the exception of Siikaneva, the EC tower footprints represent wet permafrost ecosystems with complex, heterogeneous terrain that includes moist depressions, drier, elevated hummocks and inundated microsites. Vegetation within the tower footprints (Rinne et al., 2007; Riutta et al., 2007; Sachs et al., 2008; Jackowicz-Korczyhski et al., 2010; Parmentier et al., 2011a; Zona et al., 2011; Tagesson et al., 2012b) consists of *Carex* and other sedges, dwarf shrubs (e.g., *Dryas* and *Salix),* grasses (e.g., *Arctagrostis)* and *Sphagnum* moss (with the exception of Zackenberg).

Mean daily  $T_s$  and  $\theta$  site measurements corresponding to near-surface  $(\leq 10 \text{ cm})$  soil depths were selected when possible (Table 1), to better coincide with the soil penetration depths anticipated for upcoming satellite-based microwave remote sensing missions (Kimball et al., 2012). For Siikaneva, reanalysis  $\theta$  was used in place of in situ measurements as only water table depth information was available to describe soil wetness (Rinne et al., 2007). At the Lena River site  $T_s$  and  $\theta \leq 12$  cm) observations were obtained from the nearby Samoylov meteorological station and represent tundra polygon wet center, dry rim and slope conditions (Boike et al., 2008; Sachs et al., 2008). Although *0* was also measured during the summer of 2006, the in situ records are limited to the wet polygon center location (Boike, personal communication, 2012) and were not used in this study due to the potential for overestimating saturated site conditions. For Zackenberg, site  $T_s$  measurements were obtained at a 2 cm depth (Tagesson et al., 2012a, b) within the tower footprint, while near-surface  $\theta$  (< 20 cm) and  $\geq$  5 cm  $T_s$  measurements were collected adjacent to the site (Sigsgaard et al., 2011). At Stordalen, site *0* measurements were not avail-



Fig. 1. Locations of the flux tower sites (circles) used in this study, including Barrow (BA), Kytalyk (KY), Lena River (LR), Siikaneva (SK), Stordalen Mire (SM) and Zackenberg (ZK). The Arctic Circle is indicated by the dashed line.

able at the time of this study (Jackowicz-Korczyński et al., 2010). Barrow (Zona et al., 2009; Sturtevant et al., 2012) includes southern (S), central (C) and northem (N) tower locations; in 2007 only  $CO<sub>2</sub>$  and CH<sub>4</sub> EC measurements from the northem tower were used in the analysis, due to minimal EC data availability for the other tower sites following data processing (Zona et al., 2009). Many of the Barrow  $CO<sub>2</sub>$  measurements were also rejected for the 2009 period; as a result NEE was not partitioned into  $R_{\text{eco}}$  and GPP (Sturtevant et al., 2012).

# **2.3 Remote sensing and reanalysis inputs**

Daily input meteorology was obtained from the Goddard Earth Observing System Data Assimilation Version 5 (GFOS-5) MERRA archive (Rienecker et al., 2011) with  $1/2 \times 2/3$ ° spatial resolution. The MERRA records were recently verified for terrestrial  $CO<sub>2</sub>$  applications in highlatitude systems (Yi et al., 2011, 2013; Yuan et al., 2011), and provide model enhanced  $T_s$  and surface  $\theta$  information similar to the products planned for the NASA Soil Moisture Active Passive (SMAP) mission (Kimball et al., 2012). In addition to near-surface ( $\leq 10$  cm)  $T_s$  and  $\theta$  information from the MERRA-Eand reanalysis (Reichle et al., 2011) required for the  $R_{\text{eco}}$  and CH<sub>4</sub> simulations, daily MERRA SW<sub>rad</sub>,  $T_{\text{min}}$ and VPD records were used to drive the intemal GPP calculations. The MERRA near-surface (2 m) wind parameters were also used to obtain mean daily  $\mu_m$  for the CH<sub>4</sub> simulations. The MERRA-Eand records for Greenland are spatially limited due to land cover/ice masking inherent in the reanalysis product, and MERRA  $T_s$  and  $\theta$  were not available

**Table 1.** Description of flux tower locations and site characteristics including permafrost (PF) cover and climate. The length (days) of each tower site  $CO<sub>2</sub>$  and  $CH<sub>4</sub>$  record is provided in addition to the observation year.



for the Zackenberg tower site. As a proxy,  $T_s$  was derived from reanalysis surface skin temperatures by applying a simple Crank-Nicholson heat diffusion scheme that accounts for energy attenuation with increasing soil depth (Wania et al., 2010); for  $\theta$ , records from a nearby grid cell were used to represent moisture conditions at Zackenberg.

For the daily LUE-based GPP simulations, quality screened cloud-filtered 16 day 250 m NDVI values from MODIS Terra (M0D13A1) and Aqua (MYD13Q1) data records (Solano et al., 2010) were used as model inpnts. Differences between the M0D13A1 and MYD13Q1 retrievals were minimal at the tower locations, and the combination of Terra and Aqua MODIS records rednced the retrieval gaps to approximate 8 day intervals. The NDVl retrievals correspond to the center coordinate locations for each flux tower site, and temporal linear interpolation was used to scale the 8 day NDVl records to daily inpnts. Coarser (500-1000 m resolution) NDVl records were not used in this study due to the close proximity of water bodies at the tower sites, which can substantially reduce associated FPAR retrievals. In addition, 250 m MODIS vegetation indices have been reported to better capture the overall seasonal variability in tower EC flux records (Schubert et al., 2012).

#### **2.4 TCF model parameterization**

A summary of the site-specific TCF model parameters is provided in the Supplement (Table S2). Parameter values associated with grassland biomes were selected for the LUE model VPD and  $T_{min}$  modifiers used to estimate GPP (Yi et al., 2013), as more specific values for tnndra and mossdominated wetlands were not available. Parameter values for  $\theta_{\text{max}}$  were obtained using growing-season maximum  $\theta$  measurements for each site and  $\theta_{\min}$  was set to 0.15 for scaling purposes. Model  $\varepsilon_{\text{max}}$  was specified as 0.82 mg CMJ<sup>-1</sup> for the duration of the growing season, althongh actual LUE can vary throughout the summer due to differences in vegetation growth phenology and nutrient availability (Connolly et al., 2009; King et al., 2011). The tnndra CUE ranged from  $0.45$  to  $0.55$  (Choudhury, 2000); a lower CUE value of 0.35 was used for the moss-dominated Siikaneva site due to a more moderate degree of carbon assimilation occurring in bryophytes that has been observed in other sub-Arctic communities (Street et al., 2012). For the TCF model  $F_{\text{met}}$ parameter, the percentage of NPP allocated to  $C_{\text{met}}$  varied between 70 % and 72 % for tower tnndra sites (Kimball et al., 2009) compared to 50% and 65% for Siikaneva and Stordalen, where moss cover is more abundant. The TCF model  $R_0$  parameter ranged from 4.5 and 22.4  $\mu$ MCH<sub>4</sub> d<sup>-1</sup> (Walter and Heimarm, 2000; van Huissteden et al., 2006). Values for  $Q_{10p}$  varied between 3.5 and 4 due to an enhanced microbial response to temperature variability under colder climate conditions (Gedney and Cox, 2003; Inglett et al., 2012). A  $Q_{10d}$  of 2 was assigned for CH<sub>4</sub> oxidation (Zhnang et al., 2004; van Huissteden et al., 2006). Parameter values for  $P_{trans}$ , which indicates relative plant transport ability, ranged from 7 to 9 (dimensionless); lower values were assigned to tower locations with a higher proportion of shrab and moss cover, whereas higher  $P_{trans}$  corresponds to sites where sedges are more prevalent (Ström et al., 2005; Rinne et al., 2007). For  $\lambda$ , the scaled conductance for lower site wind sensitivity was used in the  $CH<sub>4</sub>$  model simulations, except for the Lena River, which showed higher sensitivity to surface turbulence. Values for  $P_{\text{ox}}$  ranged from 0.7 in tundra to 0.8 in *Sphagnum*-dominated systems to account for higher CH<sub>4</sub> oxidation by peat mosses (Parmentier et al., 2011c). Due to a lack of detailed soil profile descriptions and heterogeneous tower footprints, soil porosity was assigned at 75 % for sites with more abundant fibrous surface layer peat (i.e., Siikaneva and Stordalen) and 70% elsewhere to reflect more humified or mixed organic and mineral surface soils (Elberling et al., 2008; Veny etal., 2011).

# **2.5 TCF model simulations**

The TCF model was first evaluated against tower EC records using simulations driven with in situ environmental data including EC-based GPP,  $T_s$ ,  $\theta$  and  $\mu_m$ . This step allowed for baseline TCF model  $R_{\text{eco}}$  and CH<sub>4</sub> flux estimates to be assessed without introducing additional uncertainties from input reanalysis meteorology and LUE model derived GPP calculations. Four additional TCF model simnlations were conducted using reanalysis  $\theta$ ,  $T_s$ ,  $\mu_m$  (in the CH<sub>4</sub> module), or intemal model GPP in place of the in sitn data. A final TCF model mn included only satellite and reanalysis-based data, and was used to establish annual GPP,  $R_{\text{eco}}$  and CH<sub>4</sub> carbon budgets for each site. Baseline carbon pools were initialized by continuously cycling ("spinning-up") the model for the tower years of record (described in Table 1) to reach a dynamic steady-state between estimated NPP and surface soil organic carbon stocks (Kimball et al., 2009). In situ data records were used during the model spin-np to establish baseline organic carbon conditions for the first five TCF model simulations, although it was often necessary to supplement these data with reanalysis information to obtain a continuons armnal time series. The final model simulation did not include in situ data in the spin-up process.

The temporal agreement between the tower EC records and TCF model simulations was assessed nsing Pearson correlation coefficients  $(r; \pm$ one standard deviation) for the daily, 8 day, and total-period (EC length of record) cumulative carbon fluxes and corresponding tests of significance at a  $0.05$  probability level. The 8 day and total-period cumulative fluxes were evaluated, in addition to the daily fluxes, to account for differences between the model estimates and tower EC records stemming from temporal lags between changing environmental conditions and resulting carbon  $(CO<sub>2</sub>, CH<sub>4</sub>)$ emissions (Lnnd et al., 2010; Levy et al., 2012). The mean residual error (MRE) between the tower EC records and TCF modeled  $CO<sub>2</sub>$  and  $CH<sub>4</sub>$  fluxes was used to identify potential positive (underestimation) and negative (overestimation) biases in the simulations; root-mean-square-error (RMSE) differences were used as a measure of model estimate uncertainty in relation to the tower EC records.

# **3 Results**

# **3.1 Surface organie carbon pools**

The TCF model generated surface soil organic carbon pools represent steady-state conditions obtained through the continuous cycling of in situ or satellite and reanalysis enviromnental data for the years of record associated with each tower site (described in Table 1). Approximately 600 and 1000 years of model spin-up were required for  $C_{\text{rec}}$  to reach dynamic steady state conditions. Over 95% of the resnlting total carbon pool was allocated to  $C_{\text{rec}}$  by the TCF model, with  $2-3\%$  stored as  $C_{\text{met}}$  and the remainder partitioned to  $C_{str}$ . The estimated carbon pools from the in situ (reanalysis-based) model spin-np ranged from approximately  $3.3 \text{ kg C m}^{-2}$  (2.3 kg C m<sup>-2</sup>) for Zackenberg and Stordalen to  $1.3 \text{ kg C m}^{-2}$  (2.1 kg C m<sup>-2</sup>) for the other tower sites.

Differences in carbon stocks, resnlting from the use of satellite remote sensing and reanalysis information in the TCF model, reflect warm or cold biases in the input *Ts* records relative to the in situ data that modified the rate of  $CO<sub>2</sub>$  loss during model initialization. The larger carbon stocks at Zackenberg, compared to the other tnndra sites, resulted from higher tower EC-based GPP inputs that often exceeded  $5 \text{ g C m}^{-2} d^{-1}$  in mid-summer, and a short (< 50 day) peak growing season (Tagesson et al., 2012a) that minimized TCF modeled  $R<sub>h</sub>$  losses. Although it was necessary to use intemal LUE-based GPP calculations for Stordalen in the absence of available  $CO<sub>2</sub>$  records, the resulting  $C<sub>met</sub>$  and  $C_{\text{rec}}$  carbon stocks were similar in magnitude to surface litter measurements at this site (Olsrad and Christensen, 2011). The TCF model simulated carbon stock for Lena River was less than a  $2.9 \text{ kg C m}^{-2}$  average determined from in situ  $($   $\leq$  10 cm depth) measurements of nearby river terrace soils  $(Zubrzvcki et al., 2013)$ , but this could have resulted from site spatial heterogeneity and the use of recent climate records in the model spin-np that may not reflect past conditions.



**Fig. 2.** TCF model simulations for GPP (lines) using input remote sensing and reanalysis information as compared with flux tower EC records (circles). Site GPP records were not available for SM and BA 2009.

# **3.2 LUE-based GPP**

The GPP simulations using reanalysis and satellite-based inputs captured the overall seasonality observed in the tower records (Fig. 2; Table 2) and explained 76%  $(r^2;$  $p < 0.05$ ,  $N = 7$ ) of variability in the total EC period-ofrecord fluxes (Eig. 3). The across-site RMSE and MRE were  $1.3 \pm 0.51$  and  $-0.1 \pm 0.7$  g C m<sup>-2</sup> d<sup>-1</sup>, respectively. Although the 8 day cumulative flux correspondence between the tower EC and TCE model GPP estimates was strong  $(r^2 = 75 \pm 16 \%)$ , the model-tower agreement decreased considerably for daily GPP ( $r^2 = 57 \pm 22$  %). These differences may reflect a delayed response in vegetation productivity following changes in atmospheric and soil conditions (Lund et al., 2010), and short term fluctuations in the reanalysis  $SW_{rad}$ inputs. For Kytalyk, the large RMSE  $(2.2 \text{ g C m}^{-2} \text{ d}^{-1})$  observed for the TCE model GPP simulations resulted from warm spring air temperatures that reduced  $T_{\text{min}}$  constraints on carbon assimilation, although a similar increase in GPP did not occur in the EC-based records. This lack of response likely resulted from a shallow (< 14 cm) early season thaw depth at this site, that limited bud break activity in deeper rooted shrubs (e.g., *Betula nana* and *Salix pulchra).* To address this, an additional simulation was conducted using a temperature driven phenology model described in Parmentier et al.  $(2011a)$  to better inform the start of growing season in the TCE model. This step reduced the corresponding RMSE



Fig. 3. Correspondence between TCF model and tower EC records for cumulative  $(g \text{ C m}^{-2})$  GPP,  $R_{\text{eco}}$ , NEE, and CH<sub>4</sub> fluxes from six pan-Aretic tower locations. The TCF model simulations include those derived from in situ measurements (open circles) or MODIS remote sensing and MERRA reanalysis inputs (MDMR; in black). A 1 : 1 relationship is indicated by the dashed line. The  $r^2$  agreement is significant at a 0.05 probability level, except for MDMRbased  $R_{\text{eco}}$  and NEE ( $p = 0.16$  and 0.27), and excludes NEE fluxes for KY (circled) due to large differences in the  $CO<sub>2</sub>$  response relative to the other sites.

difference for Kytalyk by 56% (to  $1 \text{ g C m}^{-2} \text{ d}^{-1}$ ) with an associated  $r^2$  of 67%.

Although previous EUE models (e.g.. Running et al., 2004; Yi et al., 2013) have relied solely on VPD to represent water related constraints to GPP, our approach also considers soil moisture to better account for the sensitivity of bryophytes and shallow rooted vegetation to surface drying (Wu et al., 2013). Including this additional moisture constraint reduced the overall TCE model and tower GPP RMSE and MRE differences by approximately 14% and 92%. However, the model simulations continued to overestimate GPP fluxes for Siikaneva, Eena River (2003), and Kytalyk (MRE =  $-0.6 \pm 0.8$  g C m<sup>-2</sup> d<sup>-1</sup>). This residual GPP bias could be influenced by inconsistencies between the coarse-scale MERRA reanalysis inputs and local tower meteorology, as reported elsewhere (e.g., Yi et al., 2013), although systematic biases for the high-latitude regions have not been identifled. Tor instance, periods of warmer (3 to  $4^{\circ}$ C) reanalysis  $T_{\text{min}}$  inputs relative to in situ measurements at Eena River in 2003 led to seasonally higher TCE modeled GPP fluxes. In contrast, the reanalysis  $T_{\text{min}}$  at Barrow was 2 to 7 °C cooler in mid-summer than the local meteorology; this resulted in significantly lower  $(p < 0.05)$  TCF model GPP estimates relative to the tower EC records (Table 2). It is also possible that differences in the light response curve and respiration models, used when partitioning the site EC NEE fluxes into GPP and  $R_{\text{eco}}$  (i.e., Aurela et al., 2007;

**Table 2.** Tower EC CO<sub>2</sub> records and TCF modeled gross primary production (GPP), ecosystem respiration  $(R_{\text{eco}})$  and net ecosystem exchange (NEE) derived using in situ information (in parentheses) or satellite remote sensing and reanalysis inputs. The Pearson correlation coefficients (r) are significant at a 0.05 probability level, excluding Kytalyk 2009 NEE ( $r \le 0.11$ ,  $p \ge 0.17$ ) and Barrow 2007N GPP and NEE  $(r < 0.1, p > 0.16)$ .

Site	Year	Flux	$\boldsymbol{r}$	8 day r	<b>RMSE</b>	<b>MRE</b>	Site EC	TCF Model
					$g \text{ C m}^{-2} d^{-1}$		Cumulative (g C m <sup>-2</sup> )	
Siikaneva	2005	GPP $R_{\rm eco}$ <b>NEE</b>	0.84 0.96(0.96) 0.49(0.91)	0.94 0.96(0.98) 0.92(0.92)	0.8 0.4(0.3) 0.5(0.3)	$-0.2$ $-0.3(0.1)$ $0.3(-0.1)$	361.1 289.9 $-71.2$	409.4 365.6 (274.9) $-43.8(-86.2)$
Lena River	2003 2006	GPP $R_{\rm eco}$ <b>NEE</b> GPP $R_{\rm eco}$	0.74 0.77(0.87) 0.90(0.94) 0.78 0.76(0.84)	0.91 0.83(0.91) 0.93(0.97) 0.86 0.91(0.91)	0.7 1. (0.3) 0.3(0.3) 1.1 0.7(0.6)	$-0.1$ $-0.5(-0.1)$ $-0.1(0.1)$ 0.5 0.3(0.2)	72.3 56.3 $-16.0$ 247.4 193.0	131.5 103.3(62.4) $-28.2(-9.9)$ 199.3 160(176.4)
Zackenberg	2008 2009	<b>NEE</b> GPP $R_{\rm eco}$ <b>NEE</b> <b>GPP</b> $R_{\rm eco}$ <b>NEE</b>	0.57(0.76) 0.75 0.67(0.44) 0.31(0.83) 0.91 0.86(0.90) 0.89(0.89)	0.62(0.89) 0.76 0.80(0.50) 0.37(0.85) 0.96 0.93(0.96) 0.92(0.92)	0.7(0.6) 1.8 1.1(1.3) 1.7(1.3) 1.3 0.8(1) 1.2(1)	$0.2(-0.2)$ < 0.1 0.3(0.3) $-0.3(-0.3)$ 0.6 0.4(0.1) $0.2(-0.1)$	$-54.4$ 218.2 215.9 $-2.3$ 305.0 250.3 $-54.7$	$-39.3(-71.0)$ 215.4 175.5(182.6) $-39.9(-35.6)$ 234.6 183.7(238.6) $-50.9(-66.4)$
Kytalyk	2009	<b>GPP</b> $R_{\rm eco}$ <b>NEE</b>	0.41 0.49(0.60) 0.11(0.92)	0.73 0.80(0.94) 0.01(0.95)	2.2 1.6(1.3) 1.6(1.3)	$-1.5$ $-2.2(-1.5)$ 0.9(1.5)	143.2 60.8 $-82.4$	224.9 200.2 (126.9) $-24.7(-16.3)$
Barrow	2007N 2009N 2009C	GPP $R_{\rm eco}$ <b>NEE</b> <b>NEE</b> <b>NEE</b>	0.12 0.23(0.61) 0.10(0.79) $\overline{\phantom{0}}$ $\overline{\phantom{0}}$	0.32 0.64(0.82) 0.20(0.79)	1.1 0.5(0.4) 0.8(0.4) 1.6 0.5	0.2 $0.4(-0.1)$ < 0.1(0.1) 1.4 0.4	152.0 117.4 $-34.6$ $-62.1$ $-8.3$	137.0 104.3(121.6) $-32.7(-30.4)$ $-15.6$ $-3.6$

Kutzbach et al., 2007; Parmentier et al., 2011a; Tagesson et al., 2012a; Zona et al., 2012), may have contributed to differences between the TCF model simulations and tower  $CO<sub>2</sub>$ records. However, further investigation is needed to determine the expected range of GPP and *Reco* that might result from variability in the flux partitioning routines.

# **3.3** *R*<sub>eco</sub> and NEE

The in situ TCF model  $R_{\text{eco}}$  simulations accounted for 59  $\pm$  28% and 76  $\pm$  24% ( $r^2$ ) of the observed variability in the respective daily and 8 day cumulative tower EC fluxes (Fig. 4; Table 2). As with GPP, the  $r^2$  agreement increased to 89% ( $p < 0.05$ ,  $N = 6$ ) when considering the total-period cumulative fluxes (Fig. 3). The overall RMSE difference for the in situ based TCF model  $R_{\text{eco}}$  and NEE simulations was  $0.74 \pm 0.45$  g C m<sup>-2</sup> d<sup>-1</sup> when using 5 cm depth  $T_s$  inputs. A corresponding across-site MRE of  $-2.1 \pm 5.7$  g C m<sup>-2</sup> d<sup>-1</sup> indicated that the TCF model simnlations overestimated *Reco* relative to the tower records, and slightly underestimated NEE (MRE =  $0.1 \pm 0.4$  g C m<sup>-2</sup> d<sup>-1</sup>). We also conducted TCF model simulations using 8-10 cm depth in situ

 $T_s$  inputs, instead of those from  $\leq$  5 cm (as reported in Table 2), to investigate the influence of deeper soil thermal controls on site  $R_{\text{eco}}$  response; this step reduced the overall RMSE by approximately 12%.

Incorporating the TCF intemal LUE model GPP estimates increased the overall RMSE for *R*<sub>eco</sub> and NEE by 23 % relative to the in sitn based simnlations, compared to a respective 3% and 14% increase when using reanalysis  $\theta$  or  $T_s$ inputs (Fig. 5). The model-tower daily and 8 day cumulative correspondence was also lower ( $r^2 = 32$  and 56%, respectively) for  $CO<sub>2</sub>$  simulations driven using internally derived GPP, relative to those using reanalysis  $\theta$  or  $T_s$  inputs ( $r^2 = 57$ ) and  $72\%$ ) in place of the in situ records. Without the in situ inpnts, the respective RMSE and MRE difference between the reanalysis-based  $R_{\rm eco}$  (NEE) simulations and the tower EC records averaged  $0.9 \pm 0.4$  and  $-0.2 \pm 0.9$  g C m<sup>-2</sup> d<sup>-1</sup>  $(1 \pm 0.5 \text{ and } 0.3 \pm 0.05 \text{ g C m}^{-2} \text{d}^{-1}).$ 

Correspondingly, the reanalysis and remote sensing-based TCF model  $R_{\text{eco}}$  (NEE) simulations accounted for  $51 \pm 29$  $(45 \pm 34)$ % and  $71 \pm 17$   $(62 \pm 34)$ % of the observed  $r^2$  variability in the respective daily and 8 day tower EC records. The mean  $r^2$  values exclude TCF model results for Barrow



Fig. 4. TCF model CO<sub>2</sub> simulations driven using in situ (solid lines) or remote sensing and reanalysis inputs (MDMR; dashed lines), as compared with tower EC records (circles) for  $R_{\text{eco}}$  and NEE. For BA 2009, in situ  $R_{\text{eco}}$  was not available and NEE measurements from the northern (central) tower are shown in black (gray). The TCF model  $R_{\text{eco}}$  results for SM 2006 (2007) are displayed in light (dark) red and NEE is indicated in light (dark) blue.

and Kytalyk, which did not show significant ( $r \le 0.20$ ;  $p \ge 0.16$ ) agreement with the site EC records (Table 2). For Barrow, it is likely that the water table manipulations at this site led to local temperature and moisture variability that was not refiected in the coarse reanalysis and remote sensing inputs. The minimal agreement at Kytalyk is attributed to higher  $R_h$  losses driven by warmer reanalysis  $T_s$  inputs, and increased  $R_a$  contributions due to the overestimation of GPP relative to the tower EC records.

#### **3.4 CH4 fluxes**

The in situ TCF model CH<sub>4</sub> simulations explained  $64 \pm 11$  % and  $80 \pm 12$  % ( $r^2$ ) of the respective daily and 8 day cumulative variability observed in the tower EC records (Fig. 6; Table 3), when excluding Kytalyk ( $p = 0.1$ ). The  $r<sup>2</sup>$  correspondence increased to 98 % when considering the total periodof-record emissions across the six sites (Fig. 3;  $p < 0.05$ ,  $N = 9$ ). At Kytalyk, Parmentier et al. (2011b) reported large differences in measured half-hourly CH4 fiuxes following shifts in wind direction, and larger emissions from portions of the tower footprint containing *Carex* sp., *E. angustifolium* and inundated microsites. Although this may have contributed to the observed discrepancy between the TCE



**Fig. 5.** TCF model accuracy for  $R_{\text{eco}}$  relative to  $CO_2$  records from five tower EC sites. The TCF model simulations include those determined from in situ measurement inputs; reanalysis soil moisture  $(\theta)$ , soil temperature  $(T_s)$  or TCF LUE model simulated GPP inputs; TCF simulations derived entirely from remote sensing and reanalysis (MDMR) inputs. Measures of comparison include RMSE, MRE,  $r$  values for daily and 8 day cumulative fluxes. The BA 2009 results represent the local spatial mean determined from north, central and southern Barrow tower locations.

model estimates and tower EC record, attempts to systematically screen the CH4 observations based on wind direction, or to use daily EC medians instead of mean values, did not substantially improve the model results.

On average, the in situ TCE model simulations overestimated CH4 fiuxes relative to the tower EC records  $(MRE = -2.2 \text{ mg C m}^{-2} \text{ d}^{-1})$ , with RMSE differences varying from 6.7 to 42.5 mg C m<sup> $-2$ </sup> d<sup> $-1$ </sup>. Without including  $\mu_m$ in the TCF model, the resulting RMSE increased by  $> 10\%$ and the mean daily correspondence decreased to  $r^2$  < 40%. The most substantial difference was observed for Eena River, where excluding  $\mu_m$  reduced the daily and 8 day emission correspondence by over 60%. Unlike the TCF model  $R_{\text{eco}}$ results, deeper (10 cm depth)  $T_s$  measurement inputs did not improve the RMSE values, except for Barrow (2007N) where the RMSE decreased by 35%. This sensitivity to deeper  $T_s$ conditions may refiect changes in active layer depth following water table manipulations at this site (Zona et al., 2009, 2012), and associated changes in carbon substrate availability. In contrast, the RMSE for Eena River was 15 % higher when using in situ 10 cm  $T_s$  records in the TCF model simulations instead of 5 cm depth measurements. A 6 % decrease in the RMSE occurred for Zackenberg (2008) when using the warmer (3 to 5 °C) 2 cm depth  $T_s$  records, relative to model simulations using 5 cm  $T_s$  inputs. Contrary to expectations, the 2 cm depth  $T_s$  inputs did not improve RMSE differences for Zackenberg in 2009 when site moisture conditions were drier (Tagesson et al., 2012a).

Site	Year	r	$8$ day $r$	<b>RMSE</b>	<b>MRE</b>	Site EC	TCF model	
					$mg C m^{-2} d^{-1}$	Cumulative (mg C m <sup>-2</sup> )		
Siikaneva	2005	0.72(0.75)	0.90(0.90)	21.8(16.9)	$-9.6(-1.2)$	5.9	7.6(6.3)	
Lena River	2003 2006	0.59(0.87) 0.53(0.69)	0.88(0.97) 0.81(0.78)	9.1(7.5) 6.9(9.3)	4.7(0.5) $-1.3(-4.4)$	1.4 1.4	0.9(1.2) 1.6(1.9)	
Zackenberg	2008 2009	0.78(0.84) 0.75(0.88)	0.91(0.95) 0.84(0.95)	35.7(28.5) 28.7(21.2)	11.6(2.4) $-1.1(-6.7)$	7.6 6.3	6.1(7.3) 6.5(7.4)	
Stordalen	2006 2007	0.80(0.80) 0.80(0.79)	0.88(0.89) 0.94(0.89)	35 (33.4) 39.4 (42.5)	13.3(0.9) $12.6(-5.3)$	18.3 22.1	12.6(17.9) 17.5(23.9)	
Kytalyk	2009	0.28(0.24)	0.66(0.41)	20.1(14.9)	$-6.4(0.7)$	0.9	1.1(0.8)	
Barrow	2007N 2009N 2009C 2009S	0.51(0.78) $\overline{\phantom{m}}$	0.94(0.80)	5.8(6.7) 4.5(15.9) 4.2(10.2) 7.2(7.6)	$-1.5(-2.4)$ $-0.5(-12.6)$ $0.4(-4.7)$ $-0.2(6.3)$	0.7 0.1 0.2 0.2	0.8(0.9) 0.1(0.2) 0.3(0.3) 0.2(0.2)	

**Table 3.** Tower EC CH<sub>4</sub> records and TCF model results using in situ information (in parentheses) or satellite remote sensing and reanalysis inputs. The Pearson correlation coefficients  $(r)$  are significant at a 0.05 probability level, excluding Kytalyk 2009  $(r \le 0.28, p \ge 0.07)$ .



**Fig. 6.** TCF model CH<sub>4</sub> simulations driven using in situ (solid lines) or input remote sensing and reanalysis (dashed lines) inputs, as compared with tower EC records (circles). For BA 2009, the TCF model results are simulation means for the three Barrow tower sites; diamond shapes indicate CH<sub>4</sub> flux observations from the northern (in dark gray) and central (in light gray) towers, whereas gray circles indicate observations from the southern tower.

The reanalysis driven TCF model  $CH_4$  simulations (Fig. 6; Table 3) accounted for  $48 \pm 16\%$  and  $79 \pm 8\%$  ( $r^2$ ) of the respective daily and 8 day variability in the tower EC records when excluding the less favorable results for Kytalyk ( $r^2 = 8$  and 44%, respectively). Although slightly

lower than the in situ TCF model CH<sub>4</sub> estimates, the coarser reanalysis and remote sensing driven simulations explained 96%  $(r^2)$  of the total period-of-record emissions at these sites (Fig. 3). The corresponding model RMSE was  $18.2 \pm 13.6$  mg C m<sup>-2</sup> d<sup>-1</sup>, with an associated MRE

difference of  $1.8 \pm 7.3$  mg C m<sup>-2</sup> d<sup>-1</sup> that indicated the slight model underestimation of daily  $CH<sub>4</sub>$  emissions. The model RMSE differences increased by approximately 15 % when using reanalysis  $\mu_m$  records or internal GPP estimates in place of the in situ inputs, and by  $10\%$  when incorporating reanalysis  $T_s$  and  $\theta$  inputs (Fig. 7).

#### **3.5 Estimates of annual carbon budgets**

The reanalysis and remote sensing driven TCF model simulations indicated a net  $CO<sub>2</sub>$  sink  $(NEE = -34.5 \pm 18.5 \text{ g C m}^{-2} \text{ yr}^{-1})$  for the tower sites, excluding Barrow in 2009 (NEE =  $7.3 \text{ g C m}^{-2} \text{ yr}^{-1}$ ) where the estimated *R*<sub>eco</sub> emissions exceeded annual GPP (Fig. 8). Other studies near Barrow have also reported NEE losses from wet tundra communities, resulting from drier microscale surface conditions and warming within the hummocky landscape (Huemmrich et al., 2010b; Sturtevant and Oechel, 2013) that can strongly influence  $R_{\text{eco}}$ . The corresponding TCF model *Rqco* estimates ranged from 133 (Zackenberg in 2009) to  $494 \text{ g C m}^{-2} \text{ yr}^{-1}$  (Stordalen in 2006) with lower CO<sub>2</sub> emissions occurring in the colder, more northern tundra sites. The strongest NEE carbon sink indicated by the model simulations was observed for the peat-rich Siikaneva site  $(-70.3 \text{ g C m}^{-2} \text{ yr}^{-1})$  due to high annual GPP  $(462.5 \text{ g C m}^{-2} \text{ yr}^{-1})$  relative to the other tower locations. Although tower EC  $CO<sub>2</sub>$  records were not available for Stordalen to verify the TCF model NEE results (—50.8 and  $-65.8 \text{ g C m}^{-2} \text{ yr}^{-1}$ , respectively), the estimates are slightly smaller ( $\sim$  30 g C m<sup>-2</sup> d<sup>-1</sup>) than other NEE approximations over the same time period (Christensen et al., 2012), but are similar to observations reported for other years at this site (Olefeldt et al., 2012; Mamshchak et al., 2013).

The annual TCF model CH<sub>4</sub> estimates determined using the reanalysis inputs averaged  $6.9 \times (5.5)$ gC m<sup>-2</sup> yr<sup>-1</sup> for the six tower sites. The highest  $CH_4$  emissions were observed for Stordalen and Siikaneva ( $\geq 11.8 \text{ g C m}^{-2} \text{ yr}^{-1}$ ) due to higher model-defined CH<sub>4</sub> production rates and summer reanalysis  $T_s$  records that were often 5 °C warmer than the other sites. In contrast, model CH4 emissions were lowest for Barrow  $(1.8 \text{ g C m}^{-2} \text{ yr}^{-1})$  due to smaller GPP estimates and colder summer reanalysis  $T_s$  records that did not reflect the unusually warm site conditions in 2007 (Shiklomanov et al., 2010). The annual TCF model  $CH_4$  emissions for Lena River were relatively small (2.3 g C  $m^{-2}$  $yr^{-1}$ , on average), but are similar in magnitude to site CH4 estimates determined using more complex coupled biogeochemical and permafrost models (i.e., Zhang et al., 2012b). Although the TCF modeled CH4 fluxes contributed only 1-5% of annual carbon emissions  $(R_{\text{eco}} + \text{CH}_4)$  at the tower sites, which is similar to previous reports (Schneider von Deimling et al., 2012), these  $CH_4$  emissions reduced the NECB  $(-23.3 \pm 19.6 \text{ g C m}^{-2} \text{ yr}^{-1})$  by approximately 23 % relative to NEE. The annual model estimates indicated that the site  $CO<sub>2</sub>$  and CH<sub>4</sub> fluxes, excluding Bar-



**Fig. 7.** TCF model accuracy relative to CH<sub>4</sub> records from six tower EC sites. Model simulations include those derived from in situ measurements; reanalysis soil moisture  $(\theta)$ , soil temperature  $(T_s)$ , surface wind velocity  $(\mu_m)$  or TCF LUE model simulated GPP inputs; TCF simulations derived solely from remote sensing and reanalysis (MDMR) inputs. Measures of comparison include RMSE, MRE, *r* values for daily and 8 day cumulative fluxes. Results for BA 2009 are means for north, central and southern Barrow tower locations.

row and Eena River, contributed to a net global warming potential (GWP) of  $188 \pm 68$  gCO<sub>2</sub> eq m<sup>-2</sup> yr<sup>-1</sup> over a 100 year time horizon (Boucher et al., 2009) with total GWP influences by CH<sub>4</sub> at approximately 9 to  $44\%$  that of *Rqco-* Similarly, the Eena River and Barrow sites mitigated GWP at a mean rate of  $-40 \text{ g } CO_2$ eq m<sup>-2</sup> yr<sup>-1</sup> in 2006 and 2007, but were net GWP contributors in 2003 and 2009 (25 and  $160 \text{ g } CO_2$ eq m<sup>-2</sup> yr<sup>-1</sup>, respectively). Although site  $CO<sub>2</sub>$  contributions from methantrophy during plant transport and soil diffusion were estimated to range from 3.8 to 58.3 g C m<sup> $-2$ </sup> yr<sup> $-1$ </sup>, these contributions represented < 14 % of total TCF model derived  $R_{\text{eco}}$ .

#### *4* **Discussion and conclusions**

The level of complexity in biophysical process models has increased considerably in recent years but there remain large differences in carbon flux estimates for northem highlatitude ecosystems (McGuire et al., 2012; Wania et al., 2013). An integrated TCF model  $CO<sub>2</sub>$  and  $CH<sub>4</sub>$  framework was developed to improve carbon model compatibility with remote sensing retrievals that can be used to inform changes in surface conditions across northem peatland and tundra regions. Although the TCF model lacks the biophysical and hydrologic complexity found in more sophisticated process models (e.g., Zhuang et al., 2004; Wania et al., 2010), it avoids the need for extensive parameterization by instead employing generalized surface vegetation growth, temperature, and moisture constraints on ecosystem CO2 and CH4 fluxes. Despite the relatively simple



Fig. 8. The TCF model simulation results for cumulative annual GPP,  $R_{\text{eco}}$ , NEE and CH<sub>4</sub> fluxes determined using satellite remote sensing and reanalysis inputs. For NEE, all sites are net  $CO<sub>2</sub>$  sinks except for BA 2009, which is a carbon source (in black).

model approach and landscape heterogeneity at the tower sites, the TCE model simulations derived from local tower inputs captured the overall seasonality and magnitude of *Rqco* and CH4 fluxes observed in the tower EC records. Overall the *R*<sub>eco</sub>, NEE and CH<sub>4</sub> emission simulations determined using local site inputs showed strong mean correspondence (8 day  $r > 0.80$ ;  $p < 0.05$ ) with tower EC records, but the strength of agreement varied considerably for the daily fluxes due to temporal lags between changing environmental conditions and carbon emissions (Zhang et al., 2012b), and larger EC measurement uncertainty at the daily time step (Baldocchi et al., 2008; Yi et al., 2013). The respective RMSE differences from the in situ TCE model CO<sub>2</sub> and CH<sub>4</sub> simulations averaged  $0.7 \pm 0.4$  gC m<sup>-2</sup> d<sup>-1</sup> and  $17.9 \pm 11.5$  mg C m<sup>-2</sup> d<sup>-1</sup>, which is comparable to other site-based model results (e.g., Marushchak et al., 2013; Sturtevant and Oechel, 2013).

In this study, we used near-surface  $T_s$  records in the model simulations to better coincide with the soil depths represented by upcoming satellite remote sensing missions, but acknowledge that deeper  $T_s$  controls are also important for regulating high-latitude carbon emissions. This was evident in TCE model *Rqco* results where RMSE differences between the in situ based simulations and tower EC fluxes generally improved when using deeper  $10 \text{ cm } T_s$  inputs instead of those from shallower ( $\leq$  5 cm) soil depths. However, the TCF model CH<sub>4</sub> simulations were more favorable when using near-surface (2 to 5 cm)  $T_s$  inputs. The observed CH<sub>4</sub> emission sensitivity to surface soil warming may be influenced by cold temperature constraints on CH<sub>4</sub> production in the carbon-rich root zone where organic acids are more abundant (Turetsky et al., 2008; Olefeldt et al., 2013). Eightweight carbon fractions have been shown to be more susceptible to mineralization following soil thaw and temperature changes than heavier, more recalcitrant soil organic carbon pools in high-latitude environments (Glanville et al., 2012). However, the depletion of older organic carbon stocks may also become more prevalent in permafrost soils subject to thawing and physiochemical destabilization (Schuur et al., 2009; Hicks Pries et al., 2013a) in the absence of wet, anoxic conditions (Hugelius et al., 2012; Hicks Pries et al., 2013b). Seasonal changes in  $T_s$  constraints were also evident in this study, especially in the Zackenberg records where the TCE model underestimated tower  $R_{\text{eco}}$  and CH<sub>4</sub> emissions in autumn by not accounting for warmer temperatures deeper in the active layer that can sustain microbial activity following surface freezing (Aurela et al., 2002). Allowing the TCE model vegetation CUE parameter to change over the growing season instead of allocating  $R_a$  as a static fraction of GPP may also improve model and tower  $R_{\text{eco}}$  agreement. In Arctic tundra,  $R_a$  can contribute anywhere from 40 to 70% of  $R_{\text{eco}}$ , with higher maintenance and growth respiration occurring later in the growing season when root systems expand deeper into the soil active layer (Hicks Pries et al., 2013a). Representing  $R_a$  as a fixed proportion of daily GPP in the TCF model, and not accounting for the use of stored plant carbon reserves, may also have contributed to the lower *Rqco* estimates during spring and autumn transitional periods when photosynthesis is reduced.

Our estimates of peatland and tundra  $CO<sub>2</sub>$  fluxes using TCE model simulations driven by MERRA reanalysis and satellite (MODIS) remote sensing inputs showed favorable agreement relative to the tower EC observations, with relatively moderate RMSE uncertainties of  $1.3 \pm 0.5$ (GPP),  $0.9 \pm 0.4$  ( $R_{\text{eco}}$ ) and  $1 \pm 0.5$  (NEE) g C m<sup>-2</sup> d<sup>-1</sup>. These model accuracies are similar to those reported in a previous TCE model analysis for the northem regions (Yi et al., 2013), and other Arctic EUE-based GPP studies (Tagesson et al., 2012a; McCallum et al., 2013). The associated modeltower RMSE for CH<sub>4</sub> was  $18.2 \pm 13.6$  mg C m<sup>-2</sup> d<sup>-1</sup>, and is comparable to results from previous remote sensing driven CH4 analyses (Meng et al., 2012; Tagesson et al., 2013). The larger observed differences between TCE model and tower EC-based GPP results may reflect seasonal changes in nutrient availability (Eund et al., 2010), although one peatland study reported that nutrient limitations to plant productivity could be detected indirectly by MODIS NDVl retrievals (Schubert et al., 2010b). It is more likely that this reduced correspondence resulted from fluctuations in the reanalysis SW<sub>rad</sub> inputs (Yi et al., 2011) and uncertainty associated with satellite NDVl and resulting EPAR inputs stemming from residual snow cover and surface water effects on optical-IR reflectances (Delbart et al., 2005). High-latitude studies have reported difficulty in using satellite NDVI to determine the start of spring bud burst and seasonal variability in leaf development (Huemmrich et al., 2010a). Evaluating other portions of the visible spectrum, including blue and green reflectances, in addition to NDVl has helped to alleviate this problem in remote sensing applications (Mamshchak et al., 2013) and should be considered in subsequent studies. Incorporating phenological constraints into the TCF LUE model may also better characterize early season GPP, especially for plant commnnities snch as *E. vaginatum* that are sensitive to changes in active layer depth (Parmentier et al., 2011a; Natali et al., 2012). Considering  $T_s$  as an additional constraint in the TCF LUE model may also better account for autumn GPP activity nnder frozen air temperatnres if plant-available moisture is still available within the root zone (Christiansen et al., 2012). Yi et al. (2013) attempted to address this condition by incorporating satellite passive microwave-based freeze/thaw records (37 GHz) to constrain GPP according to frozen, transitional, or non-frozen surface moisture states but did not report a significant improvement, likely dne to the coarse (25 km) resolntion freeze/thaw retrievals.

The TCF model assessment of annual NECB for the six northern tower EC sites indicate that  $CH<sub>4</sub>$  emissions reduced the terrestrial net carbon sink by 23 % relative to NEE. Althongh GPP at the Lena River and Barrow sites mitigated GWP additions from  $R_{\text{eco}}$  and CH<sub>4</sub> in two of the years examined, in most years the tower sites were GWP contributors by approximately  $165 \pm 128$  gCO<sub>2</sub>eq m<sup>-2</sup> vr<sup>-1</sup> when considering the impact of CH4 on atmospheric forcing over a 100 year time span. These resnlts are consistent with other model-based analyses of Arctic carbon fluxes (McGuire et al., 2010) and emphasize the importance of evaluating  $CO<sub>2</sub>$ and  $CH_4$  emissions simultaneously when quantifying the terrestrial carbon balance and GWP for northem peatland and tnndra ecosystems (Christensen et al., 2012; Olefeldt et al., 2012). However, ongoing efforts are needed to better inform landscape-scale spatial/temporal variability in soil moisture. temperature and vegetation controls on  $CO<sub>2</sub>$  and  $CH<sub>4</sub>$  fluxes for future model assessments using a combined network of in situ soil measurements and strategically placed EC tower sites (Sturtevant and Oechel, 2013), and regional airborne surveys. The upcoming SMAP mission may also help to determine landscape soil moisture and thermal constraints on northern carbon fluxes through relatively fine-scale (3 km resolution) and lower frequency ( $\leq$  1.4 GHz) microwave retrievals with enhanced soil sensitivity (Entekhabi et al., 2010; Kimball et al., 2012), complimented by recent improvements in Arctic-specific reanalysis data (Bromwich et al., 2010). These advances, in conjunction with a suitable model framework to quantify ecosystem NEE and  $CH<sub>4</sub>$  emissions, provide the means for regional carbon assessments and monitoring of the net ecosystem carbon bndget and nnderlying enviromnental constraints.

**Supplementary material related to this artiele is available online at <http://www.biogeoseienees.net/ll/> 1961/2014/bg-ll-1961-2014-snpplement.pdf.**

*Acknowledgements.* Financial support for this study was provided by the NASA Terrestrial Ecology and Science Terra and Aqua programs (NNX09AP52G, NNX11AD46G) and the NESSF program (NNX13AM92H) with work performed at the University of Montana under contract to the National Aeronautics and Space Administration. Additional support was also provided by the National Science Foundation ARC (1204263), and the Helmholtz Association (Helmholtz Young Investigators Group, grant VH-NG-821). J. Rinne also acknowledges financial support from the Academy of Finland (125238).

Edited by: R Stoy

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