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# Future vegetation-climate interactions in Eastern Siberia: an assessment of the competing effects of CO<sub>2</sub> and secondary organic aerosols

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#### Introduction

Warming effects on ecosystem carbon cycling in northern ecosystems (Serreze et al., 2000; Tarnocai et al., 2009), and the potential for large climate-feedbacks from losses of CO<sub>2</sub> or CH<sub>4</sub> from these carbon-dense systems have been widely discussed (Khvorostyanov et al., 2008; Schuur et al., 2009; Arneth et al., 2010). Other biogeochemical processes can also lead to feedbacks, in particular through emissions of

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biogenic volatile organic compounds (BVOC) that are important precursors for tropospheric O<sub>3</sub> formation, affect methane lifetime and also act as precursors for secondary organic aerosol (SOA). These latter interactions with SOA have a cooling effect (Arneth et al., 2010; Makkonen et al., 2012b; Paasonen et al., 2013). Condensation of monoterpenes (MT), a group of BVOC with large source strength from coniferous vegetation, on pre-existing particles increases the observed particle mass, as well as the number of particles large enough to act as cloud condensation nuclei (CCN; equivalent to particles  $> \sim 100$  nm) at boreal forest sites (Tunved et al., 2006). For present-day conditions, Spracklen et al. (2008) estimated a radiative cooling of -1.8 to -6.7 W m<sup>-2</sup> of boreal forest area from the BVOC-SOA interplay.

How future changes in MT emissions affect SOA growth and climate is very uncertain. This is partially because of the lack of process-understanding of the various steps of aerosol formation and growth, and interactions with cloud formation (Hallquist et al., 2009; Carslaw et al., 2010), and partially because the issue of how spatial patterns of changing emissions of atmospherically rapidly reactive substances translate into a changing patterns of radiative forcing, and then into a surface temperature change, has not yet been resolved (Shindell et al., 2008; Fiore et al., 2012).

The Russian boreal forest represents the largest continuous conifer region in the world. About one third of this forested area ( $\sim 730 \times 10^6$  ha) is dominated by larch (Shvidenko et al., 2007), in particular by the Larix gmelinii and L. cajanderii forests growing east of the Yenisei river on permafrost soils. Despite its vast expanse, the first seasonal measurements of MT emissions from Eastern Siberian larch have only recently been published (Kajos et al., 2013). Leaf MT emission capacities are highly speciesdependent, thus any model estimate of MT emissions from boreal larch forests that rely solely on generic BVOC emission parameterisations obtained from other conifer species will give inaccurate emission and related SOA aerosol number concentrations for this region (Spracklen et al., 2008). We therefore provide here a first assessment of MT emission rates from the Eastern Siberian larch biome, combining measured emission capacities with a process-based dynamic vegetation model and quantitatively link-

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ing MT emissions and SOA formation. We use the observations and process-models to assess climate change effects on future vegetation composition, BVOC emissions and the concentration of particles of CCN size. We discuss how the climate impact of future SOA levels from changes in BVOC emissions across Eastern Siberia compares with changes in the regional CO<sub>2</sub> balance. The chief goal of the study was not to provide a full surface climate-feedback quantification (for which today's global coupled modelling-tools are insufficient) but rather to highlight the number of potentially opposing processes that need to be covered when doing so.

#### Methods

#### 2.1 Site description, BVOC and aerosol measurements

Leaf BVOC emissions fluxes, above-canopy monoterpene concentration and aerosol particle size and number concentrations were measured during the growing season 2009 at the research station Spasskaya Pad, located ~40 km to the northeast of Yakutsk (62°15′18.4" N, 129°37′07.9" E) and centred in the Eastern Siberian larch biome (Kobak et al., 1996; Tchebakova et al., 2006). In the northern direction, no major pollution sources exist within hundreds of km, the nearest mining areas are concentrated to the south and west of Yakutsk. The predominant air flow to the site is either from southern (via Yakutsk) or northern locations. Forest fires contribute to aerosol load in summer.

An eddy covariance tower for measurements of forest-atmosphere exchange of CO<sub>2</sub>, water vapour and sensible heat was established at Spasskaya Pad in the late 1990s (Ohta et al., 2001; Dolman et al., 2004) in a L. cajanderii forest growing on permafrost soil with an understory vegetation consisting of ericaceous shrubs. The forest has an average age of ca. 185 years and canopy height is little less than 20 m. Maximum one-sided larch leaf area index in summer is around two (Ohta et al., 2001; Takeshi et al., 2008). In 2009, leaf samples for BVOC analyses were taken, accessing the up-

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per part of the canopy from a scaffolding tower located within few hundred metres of the eddy flux tower (Kajos et al., 2013). Using a custom-made Teflon branch chamber, air filtered of O<sub>3</sub> was sampled onto Tenax-TA/Carbopack-B cartridges with a flow rate of 220 mL min<sup>-1</sup>. A total of 5-12 samples were taken during the day, from two trees on south-facing branches approximately 2 m below the tree top. The cartridge samples were stored at 5 °C during the campaigns, transported afterwards to Helsinki and thermally desorbed and analysed using a thermal desorption instrument (Perkin-Elmer TurboMatrix 650, Waltham, USA) attached to a gas-chromatograph (Perkin-Elmer Clarus 600, Waltham, USA). For details on chamber, adsorbents and laboratory measurements see Haapanala et al. (2009), Ruuskanen et al. (2007) or Hakola et al. (2006).

Monoterpene concentrations were measured with a high-sensitive Quadrupole PTR-MS (Ionicon, Innsbruck, Austria) located in a hut at the foot of the eddy covariance tower. Sample air was drawn through a heated PFA tube using a 20 L min<sup>-1</sup> flow from the inlet located at 30.3 m above ground. While reporting here on monoterpenes only, a range of masses, corresponding to BVOCs e.g. isoprene, methanol, acetaldehyde were sampled sequentially, with typical dwell times of 0.5s and scanning sequences of around 4s. Measurement set-up and quality control followed Holst et al. (2010). It was not possible to import a gas calibration standard to Spasskaya Pad due to security and customs restrictions, and thus the PTR-MS could not be calibrated on-site. However, the instrument had been calibrated before and after the field campaign using a gas standard mixture from Ionimed (Innsbruck, Austria) using the same detector and instrument settings as during the field campaigns.

Aerosol particles were continuously monitored with a Scanning mobility particle sizer (SMPS) located at the foot of the eddy covariance tower, connected to a Differential mobility analyzer (DMA; Hauke type: medium; custom built; for size segregation of aerosol particles) in front of a Condensation Particle Counter (CPC; 3010, TSI Inc. USA; for determining the number of the size segregated particles). The system was identical to the one described and evaluated in Svenningsson et al. (2008). Scans

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across the size range of 6-600 nm were completed every 5 min. The SMPS data were used to determine occasions of aerosol particle nucleation. The growth rates were calculated from log-normal modes fitted to the measured particle size distribution following Hussein et al. (2005). The time evolution of the diameters at which the fitted modes peaked was inspected visually, and the growth rate was determined with linear least squares fitting to these peak diameters whenever a continuous increase in diameter was observed. In this analysis we calculated growth rates for particles from 25 to 160 nm.

The source rate for condensing vapour (Q) was determined by calculating the concentration of condensable vapour needed to produce the observed growth rate ( $C_{GR}$ ) Nieminen et al., 2010) and the condensation sink from the particle size distribution (CS, Kulmala et al., 2001). In steady state the sources and sinks for the condensing vapour are equal, and thus we determined the source rate as  $Q = C_{GR} \cdot CS$ .

#### Modelling of dynamic vegetation processes, permafrost and BVOC emissions

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We applied the dynamic global vegetation model LPJ-GUESS (Smith et al., 2001; Sitch et al., 2003), including algorithms to compute canopy BVOC emission following Niinemets et al. (1999), Arneth et al. (2007b) and Schurgers et al. (2009a), and permafrost as adopted from Wania et al. (2009). LPJ-GUESS simulates global and regional dynamics and composition of vegetation in response to changes in climate and atmospheric CO<sub>2</sub> concentration. Physiological processes like photosynthesis, autotrophic and heterotrophic respiration are calculated explicitly, a set of carbon allocation rules determines plant growth. Plant establishment, growth, mortality, and decomposition, and their response to resource availability (light, water) modulate seasonal and successional population dynamics arising from a carbon allocation trade-off (Smith et al., 2001). Fire disturbance is included in the model (Thonicke et al., 2001). Similar to other DGVMs, a number of plant functional types (PFT) are specified to represent the larger global vegetation units (Sitch et al., 2003). Model results compare well with

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observations on LAI, permafrost distribution and vegetation response to warming (see results). Total present-day modelled soil C pools over the top 2 m in Eastern Siberia are 216 Gt C, and 454 Gt C for circumpolar soils above 40° N (Table 1). A recent data-base estimate was 191, 495, and 1024 Gt C in the 0-30, 0-100 and 0-300 cm soil layer, of permafrost-affected soils, respectively (Tarnocai et al., 2009). These numbers indicate that the values calculated with LPJ-GUESS are lower than observation-based ones. most likely underestimating C-density in particular in the soil layers below few tenths of cm.

BVOC emissions models, whether these are linked to DGVMs or to a prescribed vegetation map, all rely on using emission potentials (E\*, leaf emissions at standardised environmental conditions) or some derivatives in their algorithms. In LPJ-GUESS, production and emissions of leaf and canopy isoprene and monoterpenes are linked to their photosynthetic production, specifically the electron transport rate, and the reguirements for energy and redox-equivalents to produce a unit of isoprene from triosephosphates (Niinemets et al., 1999; Arneth et al., 2007b; Schurgers et al., 2009a). A specified fraction of absorbed electrons used for isoprene (monoterpene) production ( $\varepsilon$ ) provides the link to PFT-specific  $E^*$  (Arneth et al., 2007a); in case of monoterpenes emitted from storage an additional correction is applied to account for their lightdependent production (taking place over parts of the day) and temperature-driven (taking place the entire day) emissions (Schurgers et al., 2009a).

Leaf BVOC emissions are stimulated in a future environment in response to warmer temperatures. Moreover, warmer temperatures and CO<sub>2</sub>-fertilisation of photosynthesis lead to enhanced vegetation productivity and leaf area, with additional positive effects on BVOC emissions. But higher CO<sub>2</sub> concentrations have also been shown to inhibit leaf isoprene production. Even though the underlying metabolic mechanism is not yet fully understood, this effect has been observed in a number of studies (for an overview see Fig. 6 in Arneth et al., 2011). Due to limited experimental evidence, whether or not a similar response occurs in monoterpene producing species cannot yet be confirmed, especially in species that emit from storage. The model is set-up to test this

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hypothesis (see Fig. A1). Multiple interacting processes can thus lead to enhanced global monoterpene emissions in future, or -if the " $\mathrm{CO}_2$  inhibition" is included- yield emissions that are more or less similar to present-day or even slightly smaller (Arneth et al., 2007a; Schurgers et al., 2009a) (Table 1).

Monoterpene compounds can be emitted either directly following their synthesis in the chloroplast, in an "isoprene-like" fashion, or from storage pools, resulting in an emission pattern that is independent of light availability. The observed emissions of monoterpenes by larch possibly exhibit a hybrid between emission directly after synthesis in the chloroplast and emission from storage pools, as has also been found for other coniferous species (Schurgers et al., 2009a). The needle-level measurements by Kajos et al. (2013) on larch indicated a combined light- and temperature response, even though a robust differentiation to a temperature-only model was not possible due to the limited sample size. An earlier study by Ruuskanen et al. (2007) on a 5-year old *L. sibirica* tree indicated a better performance of the temperature-only emission model for monoterpene species compared to the light and temperature approach. In the model simulations performed here, half of the produced monoterpenes were stored, whereas the other half was emitted directly (Schurgers et al., 2009a).

LPJ-GUESS is a second generation DGVM (Fisher et al., 2010) and includes plant demography, such that forest successional dynamics and competition for water and light between individual age-cohorts are treated explicitly (Smith et al., 2001). The forest growth dynamics thus differentiate between early successional, short-lived species that invest in rapid growth and shade-tolerant trees with resource allocation aimed towards longer-lived growth strategies. As a result, the model's PFTs can be mapped to tree-species when required information for model parameterisation is available. This feature provides a distinct advantage when applying the necessary BVOC emission capacities that are based on species (rather than functional-type) average values (Arneth et al., 2008; Schurgers et al., 2009b; Niinemets et al., 2010). Larch, in this model setup would be represented by the shade-intolerant boreal needle-leaf summer-

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green (BNS) PFT, and for other PFTs the global parameterisation was used (Schurgers et al., 2009a).

LPJ-GUESS was recently expanded with a permafrost module following Wania et al. (2009) and Miller and Smith (2012) in which a numerical solution of the heat diffusion equation was introduced. The soil column in LPJ-GUESS now consists of a snow layer of variable thickness, a litter layer of fixed thickness (5 cm), and a soil column of depth 2 m (with sublayers of thickness 0.1 m) from which plants can extract non-frozen water above the wilting point. A "padding" column of depth 48 m (with thicker sublayers) is also present beneath these three layers to aid in the accurate simulation of temperatures in the overlying compartments (Wania et al., 2009). Soil temperatures throughout the soil column are calculated daily, and change in response to changing surface air temperature and precipitation input, as well as the insulating effects of the snow layer and phase changes in the soil's water.

Here we run the model with 0.5° spatial resolution, using climate and atmospheric 15 CO<sub>2</sub> as driving variables as described in the literature (Smith et al., 2001). Values for the BNS "larch" PFT were adopted from previous studies (Sitch et al., 2003; Hickler et al., 2012; Miller and Smith, 2012), but with the degree-day cumulative temperature requirements on a five-degree basis (GDD5) to attain full leaf cover reduced from 200 to 100 (Moser et al., 2012). Minimum GDD5 to allow establishment was set to 350 resulting in establishment of seedlings in very cold locations. Soil thermal conductivity was 2W m<sup>-1</sup> K<sup>-1</sup>. The modelled distribution of larch in LPJ-GUESS (Fig. 1) compares well with observation-based maps (Wagner, 1997). The model was spun up for 500 years to 1900 values using CO<sub>2</sub> concentration from the year 1900 and repeating de-trended climate from 1901-1930 from CRU (Mitchell and Jones, 2005). Historical (20th century) simulations used observed CO<sub>2</sub> concentrations and were based on variable CRU climate. Simulations for the 21st century were based on ECHAM climate, using RCP8.5 emissions (Riahi et al., 2007). The model requires daily radiation, precipitation and maximum and minimum air temperatures as input (Arneth et al., 2007b). The generated GCM climate was interpolated to the CRU half-degree grid, and monthly

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values interpolated to daily ones (see Ahlström et al. (2012) and references therein). These daily fields were then bias-corrected using the years 1961–1990 as reference period, as in Ahlström et al. (2012). CO<sub>2</sub> inhibition of BVOC emissions were switched on and off in separate simulations to assess the sensitivity of our results to this process. Totals across Siberia were calculated for a grid-box that ranged from 46 to 71° N and 76 to 164° E. Simulated changes in total carbon uptake or losses were translated into radiative forcing following IPCC (2007), assuming a 50 % uptake in oceans in case of a net loss to the atmosphere Sitch et al., 2007).

#### 2.3 Modelling aerosols and CCN

To model the effect of BVOCs on CCN concentrations, we use the global aerosolclimate model ECHAM5.5-HAM2 (Zhang et al., 2012). ECHAM5.5-HAM2 includes aerosol components black carbon, organic carbon, dust, sea salt and sulfate, and describes the aerosol size distribution with seven log-normal modes. The microphysics module M7 (Vignati et al., 2004) includes nucleation, coagulation and condensation. In this study, we use the ECHAM5.5-HAM2 version with activation-type as described in Makkonen et al. (2012). For simulating secondary organic aerosol, we use the recently developed SOA module (Jokinen et al., 2015). The SOA module explicitly accounts for gas-phase formation of extremely low volatility organic compounds (ELVOCs) from monoterpene oxidation. The module implements a hybrid mechanism for SOA formation: ELVOCs are assumed to condense to the aerosol population according to the Fuchs-corrected condensation sink, while semi-volatile organic compounds (SVOCs) are partitioned according to organic aerosol mass. While simulated ELVOCs are able to partition more effectively to nucleation and Aitken mode, hence providing growth for nucleated particles to CCN size, SVOCs primarily add organic mass to accumulation and coarse aerosol modes. A total SOA yield of 15% from monoterpenes is assumed (Dentener et al., 2006). While similar assumption on total SOA yield is applied by most aerosol-climate models, the simulated SOA is likely to be underestimated (e.g. Tsigaridis et al., 2014).

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Whether or not BVOCs can increase the availability of cloud condensation nuclei (CCN) depends on the availability of sub-CCN sized particles (O'Donnell et al., 2011). Anthropogenic primary emissions are introduced to the model as 60 nm particles, hence condensation of sulfuric acid and organic vapours is generally needed in 5 order to grow these particles to CCN sizes. In Siberia, the modelled primary particle emissions are dominated by wildfires, which are assumed to inject large particles with 150 nm diameter. The model is using T63 spectral resolution with 31 vertical hybrid sigma levels.

ECHAM5.5-HAM2 was run with different BVOC emission scenarios in year 2000 and 2100 simulated offline with LPJ-GUESS (see previous section). The simulations apply present-day oxidant fields as in Stier et al. (2005). The assumption of unchanging oxidant fields induces some uncertainty for future simulations and inconsistency with present-day simulations with varying biogenic emissions, since both anthropogenic and biogenic emissions are likely to modify the atmospheric oxidative capacity. All simulations are initiated with a six months spin-up, followed by 5 years of simulation for analysis. The model climate is nudged towards ERA-40 reanalysis year 2000 meteorology, an approach that is widely used in aerosol-climate assessments (K. Zhang et al., 2014). Nudging towards reanalysis meteorology establishes evaluation of BVOC-aerosol coupling with unchanged meteorological fields, but restricts the model in terms of aerosolclimate feedbacks, since e.g. nudging future climate simulations with present-day meteorological winds is based on the assumption that e.g. cloudiness, or wind direction and - speed etc. are not changing. Present-day wildfire and anthropogenic aerosol and precursor emissions are applied for all simulations (Dentener et al., 2006). One of the foci here are BVOC, comparing present-day and future BVOC emissions with  $E^* = 1.9 \,\mu\text{g C m}^{-2}$  (leaf)  $h^{-1}$ , but keeping other emissions constant. The emissions of dust and sea salt are modelled interactively (Zhang et al., 2012).

The analysis of model results includes total particle number concentration (CN) and cloud condensation nuclei at 1 % supersaturation (CCN (1 %)). The simulations are also used to assess the radiative effects of SOA. In the simulations, the aerosol con**ACPD** 

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centrations are interactively coupled to the cloud-microphysics scheme (Lohmann et al., 2007) and to the direct aerosol radiative calculation. The aerosol indirect effect is evaluated as a change in cloud radiative forcing (ΔCRF). The direct aerosol effect accounts only for clear-sky short-wave forcing (\Delta CSDRF). The radiative effects are 5 calculated as differences from two time-averaged 5-year simulations as

 $\Delta CRF = CRF(BVOC_{2100}) - CRF(BVOC_{2000})$  $\Delta$ CSDRF = CSDRF(BVOC<sub>2100</sub>) - CSDRF(BVOC<sub>2000</sub>).

#### Results

#### **Present-day BVOC emissions**

The dynamic global vegetation model LPJ-GUESS reproduces the present-day circumpolar permafrost distribution (Fig. 1; shown as circumpolar map for comparison with Tarnocai et al., 2009) and, with the exception of the Kamchatka peninsula, simulates also the expanse of the larch-dominated forests in Eastern Siberia (Fig. 1; Miller and Smith, 2012; Wagner, 1997). Maximum leaf area index (LAI) calculated by the model for the Spasskaya Pad forest (62°15′18.4" N, 129°37′07.9" E, 220 m a.s.l), where the BVOC measurements were obtained, was 2.0 (averaged over years 1981-2000; not shown), and is in good agreement with the measured values during that period (1.6; Takeshi et al., 2008). For the "larch" plant functional type in LPJ-GUESS (Schurgers et al., 2009a), an emission potential of  $E^* = 2.4 \,\mu\text{g}\,\text{C}\,\text{m}^{-2}$  (leaf)  $h^{-1}$  was adopted in previous simulations from Guenther et al. (1995), a recommendation that at that time did not include observations from any larch species.

Kajos et al. (2013) measured for the first time MT E\* from L. cajanderii. Their measurements, taken over an entire growing season at Spasskaya Pad, suggested values of  $E^*$  ranging from  $1.9 \,\mu\text{g C m}^{-2}$  (leaf)  $h^{-1}$  at the lower end, to  $9.6 \,\mu\text{g C m}^{-2}$  (leaf)  $h^{-1}$ at the upper. Applying a weighted measured-average  $E^*$  of 6.2  $\mu$ g C m<sup>-2</sup> (leaf) h<sup>-1</sup>, inDiscussion Paper

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#### 3.2 Present-day aerosols, and links to BVOC

New particle formation events (Fig. 2a) were observed regularly. The calculated volumetric source rates of condensing vapours (Q), the product of vapour concentration required for the observed particle growth rate and particle loss rate (Kulmala et al., 2005), increased exponentially with temperature (Fig. 2b). MT concentrations increased with temperature as well, with a slope relatively similar to that found for the Q vs. T relationship (Fig. 2c). Consequently, a positive relationship emerged between Q and MT concentration (Fig. 2d), which supports previous field and laboratory evidence that MT and their oxidation products are a main precursor to the observed particle formation and growth.

Figure 2d shows the connection between the BVOC concentration and the formation rate of vapours causing the growth of the aerosol particles. Even though the monoterpene concentrations were measured above and the aerosol growth rates below the canopy, the observed correlation indicates that BVOC concentration is an important contributor to the regional aerosol growth and supports the theory that the condensation of organic vapour is largely responsible for the formation of secondary organic aerosol (Hallquist et al., 2009; Carslaw et al., 2010). Substantial within-canopy chemical reactions would be expected to worsen the relationship. The correlation depicted in Fig. 2d is determined in particular by the the formation of secondary organic aerosol

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on pre-existing aerosol particles, whereas the nucleation rate of new aerosol particles seems not to be dominated by the landscape-scale emissions and surface concentrations of BVOCs. For instance, most nucleation events in a Scots pine dominated landscape in Finland have been found in spring, when measured monoterpene concentrations in the near-surface were about one tenth of the summer time maximum (~60 ppt, vs. up to 500 ppt; Haapanala et al., 2007; Lappalainen et al., 2009). We found here MT concentrations of similar magnitude to these.

By contrast to temperature and BVOC concentrations, levels of radiation, which can be considered a surrogate for the concentration of the OH radical (OH), did not affect *Q* (Fig. 2b), even though OH has been considered an important player for aerosol formation. Rohrer and Berresheim (2006) showed a strong correlation between solar ultraviolet radiation and OH concentration at the Hohenpeissenberg site in Germany. Furthermore, Hens et al. (2014) demonstrated that the day-time OH concentrations in (especially) boreal forest depend on solar radiation. Hence, the poor relation between the source rate of condensing vapour and levels of radiation (Fig. 2b) indicates that OH-radical concentration did not have a major impact on *Q*. This agrees with the findings by Ehn et al. (2014) that ozone instead of OH is an important, if not the main, atmospheric agent oxidising organic vapours into a chemical form that condenses on particle surfaces. Thus, our results indicate that factors and processes besides the concentrations of SO<sub>2</sub> and OH seem to limit aerosol production in non-polluted environments (Kulmala et al., 2005).

## 3.3 Future carbon pools, vegetation distribution and BVOC emissions in Siberia

In a warmer environment with higher atmospheric  $CO_2$  levels, the simulations indicated drastically reduced area of permafrost in Siberia (Fig. 1). Total net primary productivity in the simulated domain increased from an annual average of 3.5 to 5.9 Pg C a<sup>-1</sup> at the end of the 21st century. An overall C loss of 100 Pg C assumed to be in the form of  $CO_2$  (since the model does not yet include a dynamic surface hydrology which would

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be necessary to assess changing methane emissions) at the end of the 21st century was calculated from the shrinking Siberian areas of permafrost (Table 1). However, warming and higher levels of atmospheric CO<sub>2</sub> led also to increasing LAI, and to larchdominated areas showing the expected north- and north-eastwards shift (Fig. 1) compared to present-day climate (Miller and Smith, 2012). The carbon uptake in expanding vegetation into permafrost-free areas, combined with enhanced productivity across the simulation domain overcompensates for the losses from C-pools in permafrost areas (Table 1).

Future MT emissions were enhanced directly as a result of warmer leaves, and augmented by the future higher LAI of larch and evergreen conifers (Figs. 1d and A1; Table 1). Since the emissions scale with the emission factors applied, the proportional increase between present-day and future climate conditions is independent of the value of E\*. Whether or not leaf MT emissions are inhibited by increasing atmospheric CO<sub>2</sub> levels to similar degree to what was found for isoprene is difficult to assess from today's limited number of studies (e.g. Niinemets et al. (2010) and references therein). Similarities in the leaf metabolic pathways of isoprene and MT production suggest such an inhibition, but possibly this effect does not become apparent in plant species where produced MT are stored, unless the storage pools become measurably depleted by the reduced production. By contrast, species emitting MT in an "isoprene-like" fashion immediately after production should more directly reflect CO<sub>2</sub> inhibition. Evergreen conifers typically emit most MT from storage pools, although recent experiments have shown that some light-dependent emissions also contribute to total emission fluxes. Accordingly, based on the leaf-level measurements, larch could follow a hybrid pattern between emission after production and from storage (Kajos et al., 2013). Without accounting for CO<sub>2</sub> inhibition, MT emissions across the model domain more than doubled (Fig. 1; Table 1) by 2100, as a consequence of higher emissions per leaf area due to warmer temperatures, and of the larger emitting leaf area in response to higher photosynthesis.

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Boreal vegetation has been shown to respond to the recent decades' warming and increasing atmospheric CO<sub>2</sub> levels with a prolonged growing season and higher maximum LAI, similar to patterns in our simulations (Piao et al., 2006). The calculated enhanced biomass growth is in-line with experimental evidence of higher C in plant biomass in warming plots at tundra field sites (Elmendorf et al., 2012; Sistla et al., 2013). In Siberian mountain regions, an upward movement of vegetation zones has been recorded already (Soja et al., 2007), while the analysis of evergreen coniferous undergrowth abundance and age shows spread of evergreen species, especially *Pinus siberia*, into Siberian larch forest (Kharuk et al., 2007). These observations thus support the modelled shift in vegetation zones, and change in vegetation type composition and productivity. Likewise, other models with dynamic vegetation also have shown a strong expansion of broadleaved forests at the southern edge of the Siberian region in response to warming (Shuman et al., 2015).

Warming and thawing of permafrost soils is being observed at global monitoring network sites, including in Russia (Romanovsky et al., 2010). Estimates of carbon losses from northern wetland and permafrost soils in response to 21st century warming range from a few tens to a few hundreds Pg C, depending on whether processes linked to microbial heat production, thermokarst formation and surface hydrology, winter snow cover insulation, dynamic vegetation, C–N interactions, or fire are considered (Khvorostyanov et al., 2008; Schuur et al., 2009; Arneth et al., 2010; Koven et al., 2011; Schneider von Deimling et al., 2012). For instance, a modelled range of 0.07–0.23 W m<sup>-2</sup> forcing associated with a 33–114 Pg CO<sub>2</sub>-C loss from permafrost regions was found for a simulation study that was based on the RCP8.5 climate and CO<sub>2</sub> scenarios, but excluding full treatment of vegetation dynamics (Schneider von Deimling et al., 2012). In a recent literature review, Schaefer et al. (2014) found a range from cumulative 46 to 435 CO<sub>2</sub>-equivalents (accounting for CO<sub>2</sub> and CH<sub>4</sub>), or 120 ± 85 Gt C by 2100 in response to different future warming scenarios and modelling approaches. In

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our simulation, the  $CO_2$ -C loss from the decreasing Siberian permafrost region would be equivalent to a 0.13 additional Wm $^{-2}$  forcing in 2100 (see methods). Likely, this number is too low since the model does not include thermokarst processes, which can facilitate rapid thaw (Schaefer et al. (2014) and references therein). The modelled carbon loss was offset when taking into account vegetation dynamics and processes across the entire Siberian study-domain (Table 1), including a shift in PFT composition, and enhanced productivity especially in the southern regions, such that the overall carbon uptake including enhanced net primary productivity and expanding woody vegetation resulted in a small negative ( $-0.09 \, \text{W m}^{-2}$ ) effect.

The temperature-dependency of monoterpene emissions, especially those from stored pools, is a well-established response on the short-term. However, a change in concentrations and hence partial pressure in the storage pools, for instance in response to long-term warming, would affect emission capacities. Changes in measured E\* when investigated over the course of a growing season have been reported and could be related to a changing production rate (Niinemets et al., 2010). Likewise, observed profiles of  $E^*$  within tree canopies appear not only related to changes in leaf area-to-weight ratios along the canopy light and temperature gradients, but also to varying production rates (Niinemets et al., 2010). Emission capacities in Q. ilex leaves adapted to warm growth environment were notably enhanced (Staudt et al., 2003), but the experimental basis for an acclimation response of BVOC emissions to temperature remains remarkably poor (Penuelas and Staudt, 2010) and is indicative of the general lack of global modelling studies accounting for possibly acclimation of process responses to environmental changes (Arneth et al., 2012). In our simulations we aim to provide a range of a possible plastic BVOC-CO<sub>2</sub> response by switching the direct CO<sub>2</sub> inhibition on and off for both isoprene and monoterpene, but we do not account for other acclimation processes.

The assessment of climate effects of changes in the CO<sub>2</sub>-C balance vs. those of BVOC-SOA interactions is challenging, since the translation of regional changes in emissions of atmospherically reactive species into related radiative forcing and then

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into a response in the climate system is highly non-linear and poorly understood (Shindell et al., 2008; Fiore et al., 2012). Based on a synthesis of measured aerosol number concentrations and size distribution combined with boundary layer growth modelling Paasonen et al. (2013) estimated a growing-season indirect radiative cloud albedo feedback of  $-0.5 \,\mathrm{W\,m^{-2}\,K^{-1}}$  for the Siberian larch region. The observation-based indirect feedback factors exceeded direct ones by roughly an order of magnitude (Paasonen et al., 2013), but a simple extrapolation based on the region's growing season temperature increases of ~ 5.5 K simulated at the end of the 21st century in our study with the ECHAM GCM does not account for the important non-linearities in the system. Present-day CCN (1.0%) concentration over Siberia was estimated to vary from extremely low values of less than 50 cm<sup>-3</sup> north of 60° N to a few hundred per cc in the southern part of Siberian domain (Fig. 3). Over the larch-dominated area (Fig. 1) the sensitivity of CCN to E\* was 5-10%. In the future, a scenario of decreasing anthropogenic emissions led to a strong decrease in calculated atmospheric SO2 concentrations and also of particle nucleation (Makkonen et al., 2012a). What is more, SOA formation only partly enhances the survival of small particles by providing additional growth (Makkonen et al., 2012a), but partly also suppresses it by increasing the coagulation sink for small particles (Fig. A2, lower left panel; see also O'Donnell et al., 2011). When only BVOC emissions were changed between present day and 2100, the relatively higher emission of BVOC under ambient and future conditions leads to substantially increased aerosol growth rates (GR) over a large part of the Siberian domain. However, increased aerosol mass due to increased SOA formation led to an increase in the condensation sink and eventually to decreased particle formation rates in some regions (Fig. A2, lower right panel). These competing effects of increased growth and increased sink are essential for quantifying the importance of the cloud albedo forcing feedback. In addition, the negative effect via future BVOC emissions may also be altered through changes in aerosol background (e.g. fire), which strongly influences the indirect aerosol effect of SOA, since in large parts of Siberia, the simulated BVOC oxidation products condense on CCN-sized aerosols already present from wildfires. In the

applied future scenarios, Siberian wildfire intensity was assumed to increase (Makkonen et al., 2012a). When separated for areas of low and high wildfire emissions (Fig. 4) it becomes clear that in areas of low wildfire activity, the increase in SOA formation was proportionally high (60 %) in nucleation mode ( $d_p$  < 10 nm), and the relative increases in SOA formation in Aitken, accumulation and coarse modes were 50, 31 and 40 %, respectively. However, the distribution of BVOC oxidation products was rather different in areas of high wildfire activity. SOA formation in coarse mode was more than doubled, while SOA in nucleation mode decreased by 30 %. It is clear that the effect of increased BVOC emission on particle population has distinct effects depending on existing background aerosol distribution. Averaged over Siberian areas of low wildfire activity, the median (mean) increase of CCN (0.2 %) was calculated to be 1 % (7 %) due to BVOC emissions changes from year 2000 to year 2100, while areas of high wildfire emission lead to median (mean) increase of 0.3 % (0.5 %).

Even though the Siberian MT emissions more than double until 2100 (Table 1), the increasing wildfire emissions and decreasing new particle formation due to reductions in anthropogenic  $SO_2$  largely offset the effect of increased BVOC emissions on CCN concentration. In wildfire plumes, the simulated CCN concentrations were high even without BVOC-induced growth of smaller particles. The radiative effect due to BVOC emission change between years 2000 and 2100 was estimated from ECHAM-HAM simulations averaged over 5 years. The increase in BVOC emission leading to additional secondary organic aerosol induces a  $-0.2\,\mathrm{W\,m^{-2}}$  change in direct clear-sky aerosol forcing over the Siberian domain until the year 2100. Furthermore, the increase in CCN concentrations leads to a strengthening of the cloud radiative effect by  $-0.5\,\mathrm{W\,m^{-2}}$  (Table 2). These changes in radiative fluxes only take into account the changing BVOC emission, and the potential concurrent changes in anthropogenic and wildfire emissions might decrease the simulated radiative effect of biogenic SOA (Carslaw et al., 2013).

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Up to now, studies that investigate the role of terrestrial vegetation dynamics and carbon cycle in the climate system typically account solely for CO<sub>2</sub>, while studies that look at BVOC-climate interactions often ignore other processes, especially interactions with vegetation dynamics or the CO<sub>2</sub>-balance of ecosystems. However, for understanding the full range of interactions between atmospheric composition, climate change and terrestrial processes we need a much more integrative perspective. Our analysis seeks to provide an example of how to quantify a number of climatically relevant ecosystem processes in the large Eastern Siberian region in a consistent observational and modelling framework that accounts for the multiple interactions between emissions, vegetation and soils. It poses a challenge to combine effects of well mixed greenhouse gases and locally constrained, short-lived substances. On global-scale level, the opposing estimates in radiative effects from ecosystem-CO2 and BVOC-SOA interactions are miniscule but it is to be expected that some of the forcing effects from SOA could lead to a notable change in regional temperatures. Clearly, our numbers are uncertain but they pinpoint the necessity for assessing surface—atmosphere exchange processes comprehensively in climate feedback analyses. While doing so, we are aware of the fact that a number of additional processes are not included in our analysis. For instance, it remains to be investigated whether a similar picture would emerge when additional feedback mechanisms are taken into consideration, e.g. SOA formation from isoprene (Henze and Seinfeld, 2006) or effects of atmospheric water vapour on reaction rates and aerosol loads, or that some of the SOA might like to partition more to the gasphase in a warmer climate. Likewise, neither the albedo effect of northwards migrating vegetation (Betts, 2000; W. Zhang et al., 2014), changes in the hydrology (which affects CH<sub>4</sub> and N<sub>2</sub>O vs. CO<sub>2</sub> fluxes), nor changes in C-N interactions (Zaehle et al., 2010) are considered here, which would require a coupled ESM that combines a broad range of dynamically varying ecosystem processes with full treatment of air chemistry and

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aerosol interactions. Quantifying the full range of terrestrial climate feedbacks, either

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globally or regionally, with consistent model frameworks that account for the manifold interactions is not yet possible with today's modelling tools.

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**Table 1.** Simulated changes in net primary productivity, BVOC emissions, and C pool size in vegetation and soils. Unless stated otherwise, values are for the simulated Siberian domain (76–164° E, 46–71° N), and represent an area of  $1.2\,\mathrm{E^7\,km^2}$ . NPP $_{\mathrm{global}}$  (given as a reference value) is global vegetation net primary productivity. BVOC in Tg C a $^{-1}$ , CO $_2$ -C fluxes in Pg C a $^{-1}$ , C pools in PgC. Simulations for monoterpene emissions for the boreal needleleaf summergreen (BNS) plant functional type were made using maximum (9.6  $\mu$ g $_{\mathrm{C}}$ g $^{-1}$ h $^{-1}$ ) and minimum (1.9  $\mu$ g $_{\mathrm{C}}$ g $^{-1}$ h $^{-1}$ ) values for  $E^*$  measured in Spasskaya Pad (see text),  $E^*$  = 6.2  $\mu$ g $_{\mathrm{C}}$ g $^{-1}$ h $^{-1}$  represents a weighted average from all observations at the Spasskaya Pad location. For BVOC, CO $_2$  inhibition was switched on and off (Arneth et al., 2007b).

	1981–2000	2031–2050	2081–2100	
NPP <sub>global</sub>	$58 \pm 15$	$66 \pm 17$	$76 \pm 14$	
NPP	$3.5 \pm 0.2$	$4.5 \pm 0.2$	$5.9 \pm 0.2$	
Carbon in	circumpolar pe	ermafrost regio	on	
Vegetation	$109 \pm 0.7$	$106 \pm 1.6$	$78 \pm 1.8$	
Litter	$81 \pm 0.5$	$68 \pm 0.3$	$44 \pm 0.3$	
Soil (0 to 2 m depth)	$454 \pm 0.03$	$392 \pm 0.4$	$255 \pm 0.5$	
Total	$644 \pm 0.4$	567 ± 1.1	$377 \pm 1.0$	
C-pools in p	ermafrost area	a of study dom	ain	
Vegetation	$41 \pm 0.6$	$38 \pm 0.6$	$35 \pm 0.7$	
Litter	$40 \pm 0.3$	$34 \pm 0.2$	$23 \pm 0.2$	
Soil (0 to 2 m depth)	$216 \pm 0.06$	$187 \pm 0.1$	$140 \pm 0.3$	
Total	$297 \pm 0.4$	$259 \pm 0.4$	$198 \pm 0.2$	
C-pools in	entire Siberia	n study domai	n	
Vegetation	45 ± 0.5	56 ± 1.5	77 ± 2.8	
Litter	$41 \pm 0.5$	$43 \pm 0.3$	$41 \pm 0.7$	
Soil (0 to 2 m depth)	$219 \pm 0.3$	$221 \pm 0.3$	$223 \pm 0.3$	
Total	$305 \pm 1.1$	$320 \pm 2.1$	$342 \pm 2.0$	
BV	OC, with CO <sub>2</sub>	inhibition		
Total_iso	4.11 ± 0.29	$4.52 \pm 0.32$	$4.80 \pm 0.24$	
BNE, MT	$1.03 \pm 0.07$	$1.06 \pm 0.06$	$1.02 \pm 0.04$	
BINE, MT	$0.23 \pm 0.01$	$0.23 \pm 0.01$	$0.18 \pm 0.01$	
BNS, MT_1.9	$0.09 \pm 0.01$	$0.10 \pm 0.02$	$0.09 \pm 0.01$	
BNS, MT_6.2	$0.28 \pm 0.04$	$0.33 \pm 0.06$	$0.29 \pm 0.04$	
BNS, MT_9.6	$0.43 \pm 0.06$	$0.52 \pm 0.09$	$0.45 \pm 0.06$	
Total_MT <sub>BNS-1.9</sub>	$1.40 \pm 0.09$	$1.44 \pm 0.10$	$1.33 \pm 0.06$	
Total_MT <sub>BNS_6,2</sub>	$1.60 \pm 0.11$	$1.68 \pm 0.14$	$1.53 \pm 0.88$	
Total_MT <sub>BNS_9.6</sub>	$1.75 \pm 0.12$	$1.86 \pm 0.16$	$1.69 \pm 0.10$	
BVOC, no CO <sub>2</sub> inhibition				
Total_iso	$3.9 \pm 0.29$	$6.0 \pm 0.48$	11.0 ± 1.06	
BNE, MT	$0.99 \pm 0.07$	$1.41 \pm 0.1$	$2.33 \pm 0.19$	
BINÉ, MT	$0.22 \pm 0.01$	$0.30 \pm 0.02$	$0.42 \pm 0.02$	
BNS, MT_1.9	$0.08 \pm 0.01$	$0.14 \pm 0.02$	$0.20 \pm 0.03$	
BNS, MT_6.2	$0.21 \pm 0.03$	$0.35 \pm 0.06$	$0.52 \pm 0.07$	
BNS, MT_9.6	$0.42 \pm 0.06$	$0.69 \pm 0.11$	1.02 ± 0.13	
Total_MT <sub>BNS-1.9</sub>	$1.34 \pm 0.09$	1.92 ± 0.13	$3.04 \pm 0.23$	
Total_MT <sub>BNS</sub> =6.2	$1.47 \pm 0.10$	2.13 ± 0.16	3.36 + 0.27	
Total_MT <sub>BNS</sub> =6.2	1.67 ± 0.13	2.47 ± 0.22	$4.90 \pm 0.47$	

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**Table 2.** Simulated changes in radiative effects due to change in BVOC emission between years 2000 and 2100, averaged over Siberian domain, Northern Hemisphere and globally. CRF: cloud radiative forcing; CSDRF: direct aerosol effect that accounts only for clear-sky short-wave forcing.

	$\Delta$ CRF (W m <sup>-2</sup> )	ΔCSDRF (W m <sup>-2</sup> )
Siberia	-0.50	-0.21
Northern Hemisphere	-0.30	-0.01
Global	-0.03	-0.01

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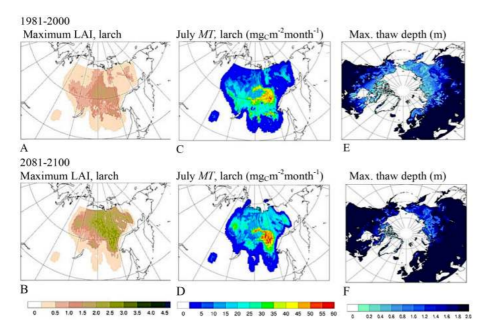


Figure 1. Simulated maximum summer leaf area index (LAI; a, b) and July emissions of monoterpenes (c, d; mg C m<sup>-2</sup> month<sup>-1</sup>) from Eastern Siberian larch. The latter were calculated applying emission factors of 6.2, obtained from the measurements at Spasskaya Pad. (e) and (f) Maximum permafrost thaw depth (August), shown here as the circumpolar map for comparison with Tarnocai et al. (2009). Values are averages for a simulation 1981-2000 (a, c, e), and for 2081-2100 (b, d, f), applying climate and CO<sub>2</sub> concentrations from ECHAM-RCP8.5. Emissions in (c) and (d) do not account for direct CO2 inhibition.

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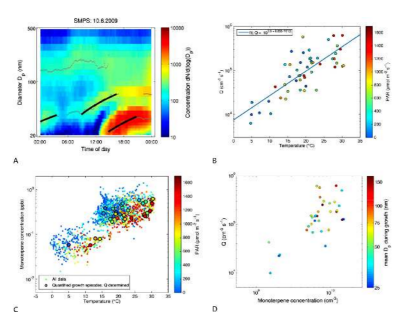
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**Figure 2.** Particle growth rates obtained from particle number size distribution ( $\bf a$ , example from day 10 June 2009). The colours indicate the measured concentrations ( $\bf dN/d\log D_p$ , cm<sup>-3</sup>) of particles with different diameters ( $\bf D_p$ , nm) over the course of a day, small circles are mean diameters of concentration modes fitted for each measurement, and the temporal change of these diameters is represented with black lines from which the growth rate is calculated. ( $\bf b$ ) shows the calculated volumetric source rates of condensing vapours ( $\bf Q$ ) as a function of air temperature (°C); data are separated by levels of photosynthetically active radiation (PAR). ( $\bf c$ ) Monoterpene concentrations (half hourly data) measured above the canopy vs. temperature measured at the same level (data separated by PAR, the data applied in ( $\bf b$ ) and ( $\bf d$ ) are indicated by encircled symbols), and relationship between volumetric source rate of condensing vapours and monoterpene concentration ( $\bf d$ ; data separated by particle diameter).

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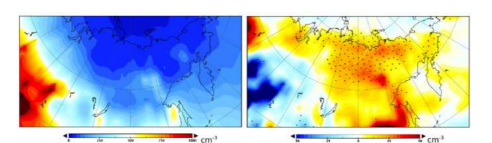
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**Figure 3.** Annual average boundary-layer CCN (1.0%) concentration (cm $^{-3}$ ) in Siberia with present-day anthropogenic and BVOC (for BNS:  $E^* = 1.9$ ) emissions (left panel), and changes in CCN (1.0%; right panel) concentration due to increase in BVOC emission between years 2000 and 2100 (simulations with CO<sub>2</sub> inhibition off). Areas with statistical significant changes in CCN are indicated.

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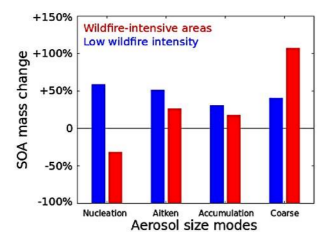




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**Figure 4.** Relative increase in SOA mass, simulated by ECHAM5-HAM in different aerosol size modes due to BVOC emissions increase from the year 2000 to 2100. The areas are averaged over Siberia, and the BVOC emissions for years 2000 to 2100 (example is for  $E^* = 1.9$ ). Areas were separated by wildfire emissions (using an emission limit of  $10^{-11}$  kg m<sup>-2</sup> s<sup>-1</sup>).

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**Figure A1.** Present-day (top panels: 1981–2000) and end of 21st century (bottom panels: 2081–2100) total monoterpene (left panels) and isoprene (right panels) emissions for the month July (mg<sub>C</sub> m<sup>-2</sup> month<sup>-1</sup>). Simulations show results with CO<sub>2</sub> inhibition switched on and off.

20 30 40 50 60 70 80 90 100 110 120

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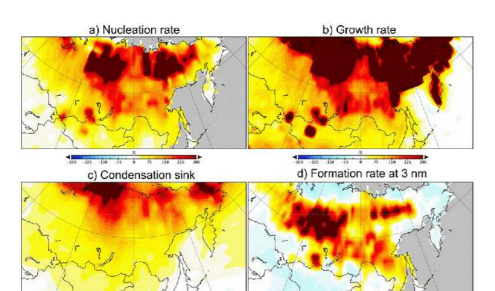
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**Figure A2.** Relative change between years 2000 and 2100 (%) nucleation rate **(a)**, growth rate **(b)**, condensation sink **(c)** and formation rate of 3 nm particles in response to altered BVOC emissions (see methods).

-150 -112 -75 -38 0 38 75 112 150

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