Environmental conditions regulate the impact of plants on cloud formation

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The terrestrial vegetation emits large amounts of volatile organic compounds (VOC) into the atmosphere, which on oxidation produce secondary organic aerosol (SOA). By acting as cloud condensation nuclei (CCN), SOA influences cloud formation and climate. In a warming climate, changes in environmental factors can cause stresses to plants, inducing changes of the emitted VOC. These can modify particle size and composition. Here we report how induced emissions eventually affect CCN activity of SOA, a key parameter in cloud formation. For boreal forest tree species, insect infestation by aphids causes additional VOC emissions which modifies SOA composition thus hygroscopicity and CCN activity. Moderate heat increases the total amount of constitutive VOC, which has a minor effect on hygroscopicity, but affects CCN activity by increasing the particles’ size. The coupling of plant stresses, VOC composition and CCN activity points to an important impact of induced plant emissions on cloud formation and climate.
Volatile organic compounds (VOC) such as isoprene and terpenes emitted by plants have various biological functions, including plant growth, defense and communication\textsuperscript{13}. It is estimated that 1,000 Tg of biogenic VOC are emitted globally per year, far exceeding total VOC emissions from human activities\textsuperscript{15}. The VOC emissions from plants are closely coupled to cloud formation and climate via the formation of secondary organic aerosol (SOA)\textsuperscript{4–6} that contribute to the regional and global cloud condensation nuclei (CCN) budget\textsuperscript{7} (Fig. 1). VOC emissions are regulated by biotic and abiotic environmental factors. Heat, drought or infestation, are stress factors that cause deviations from the plants’ optimal living conditions. Environmental factors and stresses change plant emissions significantly in two ways\textsuperscript{8–10}. They can either increase or decrease the amount of constitutive VOC (Fig. 1, left path), or stimulate biochemical pathways that induce the emission of other types of VOC. The latter VOC (via homogeneous nucleation) and the CCN activity of the SOA particles was directly determined (see Methods section and Supplementary Fig. 1). It is found that insect infestation, as an example of biotic factors, caused additional VOC emissions which modified SOA composition thus hygroscopicity and CCN activity. And heat, as an example of abiotic factors, increased the total amount of constitutive VOC emissions, which had a minor effect on the hygroscopicity of SOA, but affected CCN activity by increasing the particles’ size.

**Results**

**Effect of insect infestation.** The constitutive VOC emissions of boreal trees in the absence of stresses are dominated by

![Figure 1](image-url)
monoterpenes\textsuperscript{19} (see also Supplementary Fig. 2). Under biotic stress such as insect infestation, the composition of emitted VOC changes significantly and is often dominated by induced VOC such as sesquiterpenes and green leaf volatiles\textsuperscript{2,8,18}. The CCN activity of SOA is characterized using the hygroscopicity parameter $\kappa$, wherein higher $\kappa$ indicates higher CCN activity for particles of a given size. For comparison, the highly water-soluble inorganic salt ammonium sulfate has a $\kappa$ of $\sim 0.6$ and highly water insoluble and non-wettable black carbon has a $\kappa$ of 0. $\kappa$ was $\sim 0.15$ for SOA from monoterpene-dominated VOC emissions from unstressed pines, composed of 78% monoterpenes and 8% sesquiterpenes (Fig. 2, ‘MT-dominated’ case). During the measurements with insect infested boreal trees, sesquiterpenes dominated the VOC emissions (72% of the total carbon). Concomitant with the shift of the emission composition from monoterpene-dominated to sesquiterpene-dominated, $\kappa$ of SOA decreased significantly from 0.15 $\pm$ 0.02 to 0.07 $\pm$ 0.01 (Fig. 2).

Effects of heat and drought. In contrast to biotic factors, abiotic factors such as mild heat (up to 35 °C) did not significantly change the relative contributions of the different VOC classes (monoterpenes, sesquiterpenes and others, as described in Method) for both constitutive emissions (Supplementary Fig. 2) and induced emissions (Supplementary Fig. 3). Within each class, the contribution of some individual compounds changed, most distinct in the ‘others’ class, because specific compounds respond to temperature changes differently (cf. Supplementary Fig. 4). However, even mild heat increased the total VOC emissions for both types of emissions substantially (Fig. 3), consistent with previous studies\textsuperscript{2,29}. Accordingly, heat led to larger SOA particles under similar photooxidation conditions, but the $\kappa$ values of the SOA remained relatively invariant as temperature changed, with a seemingly small decreasing trend with increasing plant temperature. For the
SOA from constitutive emissions, the difference of κ between 25 and 35°C are not statistically significant (t-test, P = 0.09) and for the SOA from induced emissions, the difference in κ is significant only for the entire temperature interval of 22–34°C (t-test, P = 0.017, Fig. 3). The latter may be partly due to the detailed changes in the ‘others’ class or minor variations in the contributions of specific VOC classes. In this study, the particles were formed by homogeneous nucleation which means that particle number and particle size increased with increasing emissions. In the presence of pre-existing particles, only the size of the particles will be affected.

The CCN activity of particles is determined by their size and hygroscopicity7,11,30. The effect of elevated plant temperature on particle size was dramatic due to the substantial increase in the amount of VOC emitted and thus their oxidation products, which enhanced particle growth by condensation. However, the effect of elevated plant temperature on κ itself was small, which is attributed to the overall little change in the emission composition. κ determines the critical diameter for CCN activation. Larger aerosol size at constant κ implies that more particles will be activated and more cloud droplets will be formed (see also Supplementary Fig. 5). Moreover, the effect of mild heat on particle size and κ is independent whether the emissions are constitutive or induced. The effect of heat in the case of induced emissions by biotic stress represent the effect of co-occurring stressors, a common situation for plants under natural conditions31.

In addition to heat, we also investigated the effect of water shortage (drought) on plant VOC emissions. Similar to the effect of heat, we found that with monoterpenes dominating the total emissions (>80%), the general emission composition of a pine did not change much with drought (Supplementary Fig. 6a). However, drought decreased the total amount of emissions (by up to ~30%, Supplementary Fig. 6b). The magnitude of the response of VOC emissions to drought was relatively insensitive compared with heat. We conclude that decreasing VOC emissions by drought should affect the CCN activity of SOA in the opposite way as heat, and the overall effect would be a smaller activated fraction and less cloud droplets.

Discussion

Biotic and abiotic stressors affect VOC emissions of plants and therefore the resulting SOA. Increased temperatures increase VOC emissions9,29 promoting particle growth leading to a higher number of particles larger than 100 nm, that is, higher fractions of CCN active particles in the boundary layer12,32–34. Biotic stresses enhances the SOA mass concentration16–18 and a model study assuming an increase of monoterpenes emissions indicates higher aerosol and CCN concentrations in forests influenced by insect outbreaks17. The results, presented here for the first time, directly show how environmental conditions of plants can affect the CCN activity of biogenic SOA via the plants’ emissions: biotic stresses by causing induced VOC emissions, modifying hygroscopicity, or abiotic stresses by changing the amount of constitutive VOC emissions, affecting particle size distribution.

Our findings provide important information on the potential impact of vegetation on the CCN activity of biogenic SOA, and thus on cloud properties. They are one puzzle piece for understanding the complex coupling between terrestrial plants and climate change. Climate change induces both phenological changes, evolutionary and genetic changes and migration12. Plant species with long lifetime such as trees have limited capability to adapt13 and if climate zones move faster than vegetation zones, as currently observed15, more trees will experience stresses more often. Furthermore, with climate change, the frequency of unfavourable environmental conditions such as heat, droughts, and infestation are projected to increase in many regions5. Our study mainly considered mild abiotic stresses, excluding extreme heat waves and severe droughts. In this case, heat enhances the CCN activity of particles by forming larger SOA particles, which also scatter more solar radiation13. Insect infestation of plants is expected to become more prevalent since herbivorous insects survive better in warming climate13, adding to the fraction of forest being already affected by infestation at present16. These biotic stresses lead to induced VOC emissions which in case of sesquiterpenes decrease CCN activity. However, these changes in κ can be compensated by higher SOA yields of induced VOC18.

This study shows that environmental factors have an important impact on ambient CCN concentrations and properties of SOA. The effects of induced emissions should be considered in models that simulate the CCN concentrations and the impact on climate. Neglecting such effects can lead to significant biases. For example, models often simulate the CCN concentrations assuming a constant κ value of ~0.1 for all organic aerosols regardless of the source. However, a study showed that a change of κ of SOA by 50% (from 0.14 to 0.07 and 0.21) affects the CCN number concentration by ~40% (ref. 36). To demonstrate the potential effect of κ change obtained here, we applied a decrease of κ due to biotic stresses to a typical particle size distribution dominated by organic components as observed over a boreal forest in Finland as shown in Fig. 4. Our calculations show that if the plant emissions were dominated by sesquiterpenes instead of monoterpenes, the CCN number

Figure 4 | Impact of hygroscopicity parameter κ changes on cloud condensation nuclei concentrations. Measured ambient particle number size distribution (black dotted line, left axis) and the derived accumulated cloud condensation nuclei (CCN) number size distribution (right axis) are shown. The particle size distribution was measured in a boreal forest near Jämiäjärvi, Finland in May 2013 at a period when organics dominated the total aerosol mass (>80%). The CCN concentration was obtained considering 0.2% supersaturation, a typical supersaturation in clouds, using κ of 0.15 (blue line) and 0.07 (red line) corresponding to the value for SOA from monoterpenes-dominated and sesquiterpene-dominated emissions obtained in our plant chamber study shown above. Note the log x axis.
concentration would reduce by ~60% at 0.2% supersaturation. This suggests that biotic stresses can have significant influence on the CCN concentration in areas where biogenic SOA components dominate the particle composition. Currently, there are no direct field measurements reporting the effects of biotic stress on the CCN activity of biogenic SOA and CCN number concentrations, to our knowledge. Future field measurements of CCN in periods when biotic stresses induced emissions are dominant will help to assess the impact of biotic stresses on CCN activity and concentration.

The specific impacts on the ambient CCN number concentration and cloud formation depends on parameters such as ambient particle size distribution, aerosol composition, mixing state and supersaturation in clouds. Moreover, more biotic and abiotic particle size distribution, aerosol composition, mixing state and formation and cloud formation depends on parameters such as ambient concentration.

assess the impact of biotic stresses on CCN activity and when biotic stresses induced emissions are dominant will help to CCN activity of biogenic SOA and CCN number concentrations, field measurements reporting the effects of biotic stress on the This suggests that biotic stresses can have significant influence on

Nevertheless, we provide a general scheme derived from our Supplementary Fig. 7). Despite the complexity, the effects of biotic stress plus heat caused a 93% increase in the CCN number concentration, respectively. The biotic stress alone caused a 47% increase while for the constitutive emissions, heat and drought cause a 27% increase or a 37% decrease in the CCN number concentration, thus the composition and CCN activity of SOA. For example, mechanical stress due to a passing storm can induce higher monoterpene emissions from pine38,39, which should affect CCN activity similar to heat but in a more quick and episodic way. We investigated young boreal tree species. It is possible that different plant species with different ages may exhibit different responses in VOC emissions to stresses and thus further change SOA composition and properties. For example, simulated herbivory on different tree species has been shown to cause different responses in VOC emissions, which alters the SOA composition with some variability.40,41. Because various plant species with different ages as well as various environmental factors are involved and multiple factors can have synergistic or counteracting effects, the overall impacts are complex and cannot be assessed quantitatively here. Nevertheless, we provide a general scheme derived from our findings of the effects of biotic stresses, heat and drought on particle size, κ and CCN number concentrations in Supplementary Fig. 7. Our results show the potential importance of induced emissions for SOA acting as CCN. Here for the constitutive emissions, heat and drought cause a 27% increase or a 37% decrease in the CCN number concentration, respectively. The biotic stress alone caused a 47% increase while biotic stress plus heat caused a 93% increase in the CCN number concentration compared with the reference case of constitutive emissions at room temperature (right lower white dot in Supplementary Fig. 7). Despite the complexity, the effects of various stresses, especially understudied biotic stresses, should be further investigated in laboratory and field studies as well as integrated into comprehensive models to better represent the feedbacks between terrestrial plants and climate.

Methods

Experimental setup and procedure. The experiments were conducted in the Jülich Plant Atmosphere Chamber (IPAC). VOC emitted from boreal forest trees (pine (Pinus sylvestris L.), spruce (Picea abies L.), and birch (Betula pendula L.)) in a plant chamber were fed into a separate reaction chamber and were degraded by photooxidation to form SOA via homogeneous nucleation. The setup used here has been described in details by Mentel et al42.43. and was used previously to study SOA formation and properties.44.46.48. A schematic representation of the setup is shown in Supplementary Fig. 1. IPAC consists of three large glass chambers (0.164, 1.150, and 1.450 m³), each in a separate temperature-controlled housing and operated as a continuously stirred tank reactor. The 1.450 m³ chamber served as reaction chamber (RC), while the smaller chambers were used to host the plants, denoted as plant chambers (PC, PC1 for the 1.150 m³ chamber and PC2 for 0.164 m³ chamber). VOC emitted from trees in a plant chamber were fed into the reaction chamber and were degraded by photooxidation to form SOA via homogeneous nucleation. The PC were illuminated with discharge lamps (HQL 400 W/D; Osram, Munich, Germany), which simulate the solar spectrum reaching photosynthetic photon flux densities (PPFD) of up to 480 μmol m⁻² s⁻¹ (PC1) and 700 μmol m⁻² s⁻¹ (PC2) at full illumination. Switching on and off these visible-light lamps provided a day-night cycle for the plants. Purified air, which was free of particles, VOC, NOx, and ozone with around 350 ppm CO₂ added, flowed through the PC and transferred the VOC emitted by the plants to the RC. Besides the flow from the PC, two additional air streams supplied the RC with ozone (≈90 ppb) and water vapor. By controlling the humidity in this air stream, the relative humidity (RH) in the RC was held at constant 65% at constant RC temperature of 17 °C. The residence time of the chambers were ~20 min in the PC1, 5–8 min in the PC2 and approximately 63 min in the RC. Inside the RC, a UV lamp (Philips, TUV 40 W, λmax = 254 nm) was switched on for certain periods to produce OH radicals from ozone photolysis. Subsequent reaction of O (1D) radicals with the RC ambient concentration depended on the introduced VOC amount since O₂, H₂O and UV intensity, and thus OH production, were kept constant. OH concentrations, derived from an OH tracer (deuterated cyclohexane-d₆), typically ranged 2 × 10⁻¹² – 6.5 × 10⁻¹² molecules per cm³. This range is about an order of magnitude higher than that observed in the atmosphere during summer.49

The experiments were conducted by applying the following cycle: first the visible light in the PC was turned on to initiate the diurnal cycle as described below. We waited until the RC reached a steady-state regarding VOC concentrations. The steady state also included the ozonolysis reactions of VOC since ozonolysis was not switched on in the RC. After reaching the steady state, the UV lamp in the RC was switched on to generate OH radicals and to induce photochemical particle formation. Formation of particles was only observed when the UV lamp was switched on and OH radicals were produced, as observed previously.50. Experiment start is defined as the time when the UV lamp was switched on.

In this study, a mixed seedling stand of two pines (Pinus sylvestris L.), one spruce (Picea abies L.), and one birch (Betula pendula L.) (3–4 years old, about 1.1–1.3 m high) was housed in the PC1. These trees are mainly monoterpene emitters under unstressed conditions.51. They were stored outdoor under natural conditions in the Forschungszentrum Jülich campus located near a forest. These trees were wounded and the wounds were used to access the exact species. They were only slightly infested and visual check only showed very slight defoliation and foliage discoloration without any significant damages. Such insect infestation is a part of the natural conditions that plants are facing. The aphid infestation was consistent with the high sesquiterpene emission composition (Supplementary Fig. 3). Similar sesquiterpene-dominated emission composition was observed reproducibly for different types of boreal trees on different individuals (pine, spruce and birch) and in different years. Therefore, such emission composition represents one typical emission composition when trees are under insect infestation. Regular forest insect observations reveal that insect infestation is common to about 10% of boreal trees.14,16,52

To simulate the diurnal cycle for the trees, at 02:00 UTC the lamps were turned on sequentially creating an artificial dawn of 1 h. After 15 h of full illumination, the lamps were turned off in the reverse fashion. This led to a dark period of 7 h for the plants. The temperature in the PC was varied between 22 and 34 °C to investigate the influence of temperature on VOC emissions and strengths of volatile and minor monoterpene emissions in the dark. After setting a new temperature in the PC, the plants had 8–18 h to adjust to the new conditions. In addition, two experiments were conducted in which the lamps in the PC remained off during the day and mainly VOC out of storage pools of the plants were emitted.53. For these emissions, fractions of monoterpene and sesquiterpene were between those of monoterpene-dominated and sesquiterpene-dominated cases (denotes as ‘intermediate’ case) because light dependent sesquiterpene emissions decreased more than monoterpene emissions in the dark. All other experimental conditions except otherwise mentioned were kept the same to make the experiments comparable.

In absence of biotic stress, these emissions were studied in a series of experiments. A stand of eight pine trees situated in the PC1, a single spruce and a single birch situated in the PC2 were investigated separately for VOC emissions (Supplementary Fig. 2) and particle formation.55. By raising the plant temperature, VOC emissions were increased and the CCN activity of particles produced from these tree emissions were investigated. In this series of experiments, the CCN data from the experiments with spruce and birch emissions are limited because particle sizes were small and particle numbers were low for CCN activity studies due to low VOC concentrations. Therefore mainly CCN data from pine are discussed. Similar to the stressed trees case, the temperature in the PC was varied between 20 and 35 °C to investigate effects of mild heat.

In addition, to investigate the effects of drought on pine VOC emissions, a single pine tree (unstressed) was placed in the PC2 and watering was stopped and restarted in the same way as described by Wu et al.56. Briefly, in these experiments, a three to four year old Scots pine was exposed to a diurnal rhythm of 11 h illumination (06:00-17:00) and 11 h darkness, and a simulation of twilight of 1 h each in the morning and evening, respectively. The pine was exposed to several drought/watering cycles. The emission rates at the stable period during the day were used. Heat and drought are used as two examples of the various abiotic factors affecting plant VOC emissions57,59-65.

Instrumentation. The VOC were monitored with two Gas Chromatography-Mass Spectrometry (GC-MS, Agilent) systems and with a Proton Transfer Reaction—Mass Spectrometer (PTR-MS, Ionicon) with the details described in Mentel et al.42, and Kleist et al. Briefly, GC-MS systems measured VOC at the outlet of the PC. One GC-MS system was optimized to measure VOC from C5 to C20 including monoterpenes and sesquiterpenes as well as monoterpenes plus oxygenated sesquiterpene activity similar to heat but in a more quick and episodic way. We

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oxygenated VOC from methanol up to C10 compounds. Calibration of both systems was conducted as described by Heiden et al. The PTR-MS was used to determine the concentrations of VOC and oxidation products at a time resolution of 10 min and was switched continuously between the outlet of the PC and the outlet of the RC.

The number concentration and size distributions of particles were measured with an Ultrafine Condensation Particle Counter (CPC, TSI, Model 3025A) and a Scanning Mobility Particle Sizer (SMPS, TSI, Model DMA 3071 and CPC 3022A).

**VOC emission composition and related SOA composition.** Induced VOC emissions are de-novo emissions, that is, they are directly coupled to photosynthetic activity of the plants. Although VOC induced by stress depend on tree species and specific stressors, there are mainly three groups of compounds based on their biosynthesis pathways: terpenoids (such as monoterpenes, sesquiterpenes), C3 (acrylonitrile, (LOX) products) (also known as green leaf volatiles) and aromatic products of the shikimate pathway (for example, methyl salicylate (MeSA)). Constitutive terpenoid emissions are also of de novo type. In case plants have storage pools (for example, resin ducts), they are able to store some compounds like monoterpenes. Compounds from pools are emitted by physical or chemical processes. Because monoterpenes and sesquiterpenes are the dominant components in the VOC emissions of this study, which together typically account for more than 90% of the total VOC emissions, the VOC are classified as three classes: monoterpenes, sesquiterpenes and others (including LOX products, oxygenated VOC, aromatic compounds, isoprene and so on). We use the emission composition to refer to the relative contributions of different emission classes (monoterpenes, sesquiterpenes and others) in total VOC emissions.

On the basis of the particle yield data of monoterpenes and sesquiterpenes from plant emissions in our previous studies (17% for monoterpines and 5% for sesquiterpenes), the relative fraction of SOA components from monoterpines and sesquiterpenes oxidation could be roughly estimated from the VOC emission composition. For example, assuming the total SOA mass can be predicted by a linear combination of SOA yields from each precursor, the mass ratio of the SOA components from monoterpenes to the SOA components from sesquiterpenes in the 'Intermediate' case in Fig. 2 is around 0.8.

**Dropout activation measurement.** The number concentration of activated particles was measured with a Cloud Condensation Nuclei Counter (CCNC, Droplet Measurement Technologies, CCNC-100) for supersaturations (SS = RH-100%) between 0.17 and 1.1% with the setup described previously. In parallel, the particle size distribution (15.1–399.5 nm) and total number concentration were measured with a SMPS system (TSI Model 3071 and CPC 3022A) and a water-based CPC (TSI, Model 3785), respectively. Before entering the instruments, the poly-dispersed aerosol was dried with a silica gel diffusion drier to RH <5% and then neutralized with a Kr-85 neutralizer (TSI Model 3077).

The activated fraction was calculated as the ratio of the number concentration of activated particles to the total particle number concentration. The calculated activated fraction was compared with the cumulative size distribution starting from the maximum diameter. The size, at which the cumulative particle size distribution was equal to the activated fraction, was defined as dry critical activation diameter ($D_{\text{crit}}$). It was assumed that the aerosol particles were internally mixed. The method is applicable to the period when the nucleation already stopped, i.e., we measured the CCN activity at the period when the conditions in the RC reached the steady state. Five different SS were set in the CCNC: the first step for 20 min, the others for 10 min each. To ensure stable conditions in the CCNC, only data from the last 6 min of each SS step were used to determine $D_{\text{crit}}$, which overlapped with three SMPS scans each lasting 2 min. For each set of SS, the $D_{\text{crit}}$ values derived from three size scans were averaged. The contribution of multiple charged particles was corrected with the measured size distribution assuming a natural charge distribution. Ammonium sulphate aerosol was used to calibrate the SS of the CCNC based on data sets in the literature. The CCNC data were corrected for overlap with other instruments using the Oui software.

From the $D_{\text{crit}}$ and SS data, the hygroscopicity parameter $\kappa$ was determined using the method in Petters and Kreidenweis. Petters and Kreidenweis developed a theory to parameterize CCN activity data using $\kappa$ based on Köhler theory. $\kappa$ is a measure of the hygroscopicity of particles, which is defined in the following equation:

$$\frac{1}{\Delta w} = 1 + \kappa \frac{V_i}{V_e}$$

where $\Delta w$, $V_i$ and $V_e$ are the water activity, volume of solute and volume of water in an activated droplet, respectively.

The following equation can be derived by using equation (1) in the original Köhler theory:

$$S = \left(1 + \kappa \frac{V_i}{V_e}\right)^{-1} \exp\left(\frac{M_w \sigma_s}{RT \rho_w \Delta_D}\right)$$

where $\sigma_s$ is the surface tension of droplet solution; $\rho_w$ is the density of water; $R$ is the gas constant; $T$ is the temperature. $\kappa$ is assumed to be equal to that of water. Although organics can partition to the droplet surface, using the surface tension of water to calculate $\kappa$ is a reasonable assumption for droplet at activation.

Compared with the Köhler theory to other parameterizations of water activity (namely van’t Hoff factor approach) yields

$$\kappa = \frac{M_r \rho_s}{M_w \rho_w}$$

where $\rho_s$ and $\rho_w$ are the density of solute and water, and $M_r$ and $M_w$ are the molecular weight of solute and water, respectively. i is the van’t Hoff factor. It is the actual number of molecules or ions produced per solute molecule when a substance is dissolved. Since most organics do not dissociate, i is close to 1. The variability of density of most organics in SOA is small and can be assumed to be constant. equation (3) shows that $\kappa$ is inversely proportional to the molecular weight of solute assuming other factors are relatively constant.

$\kappa$ of SOA formed from the photooxidation of VOC emitted by unstressed trees and stressed trees were compared using t-test. $\kappa$ of SOA formed at different plant temperature were also compared.

**Ambient particle measurement.** The ambient particle size distribution and chemical composition were measured in a boreal forest near Janiärv, Finland in May and June of 2013 during the PEGASOS (Pan-European Gas-AeroSol-climate interaction Study) Finland campaign using instruments on board a Zeppelin-NT airship. The particle number and size distribution were measured using a CPC (TSI model 3786) and SMPS (TSI, DMA 3081 and CPC 3786). The particle chemical composition was measured using a High-Resolution Time-of-Flight Aerosol Mass Spectrometer (HR-ToF-AMS, Aerodyne Research Inc.) adapted to airship measurement. The measured particle size distribution was used to derive the CCN number concentrations. The activation diameter ($D_{\text{act}}$) was obtained at 0.2% supersaturation. In the calculation of CCN number concentration, the activation fraction was set to 0.5 and 1, respectively, if $D_{\text{act}}$ was above, within and below a size bin of SMPS.

**Modelling of the effects of stresses on CCN concentrations.** The effects of various biotic stress, heat and drought on and $\kappa$ and particle size, thus on CCN number concentration were conceptually modelled based on the results in this study as shown in Supplementary Fig. 7. The CCN number concentration was derived at 0.2% supersaturation using a typical particle size distribution in the boreal forest in Finland (cf. Fig. 4).

While for the SOA from constitutive emissions, heat and drought have little effect on $\kappa$, heat increases particle size and drought decreases particle size due to their effects on VOC emission strengths. Similarly, for SOA from induced emissions, heat also increases particle size while having little effect on $\kappa$. The particle size changes due to heat were calculated using SOA mass changes corresponding to the exponential changes of VOC emissions with temperature as derived from Fig. 3. A temperature increase of 3.7 °C (a global mean temperature in 2100 projected by IPCC report 2013 (ref. 3) was used here. The particle size change due to drought was calculated using the SOA mass change corresponding to the total VOC changes in Supplementary Fig. 6. A cubic relationship between the particle mass and diameter was assumed in the calculations of the effects of these environmental factors on particle size. The effect of particle size (shown as median diameter) on CCN number concentration was obtained by shifting the size distribution in constant logarithmical diameter steps.

Induced emissions decrease $\kappa$ while increasing particle size due to the higher SOA yield of sesquiterpenes compared with monoterpenes. The particle size change due to biotic stress induced emissions was calculated using the SOA mass corresponding to the SOA yields of sesquiterpene-dominated emissions (17%) and monoterpenes-dominated emissions (5%) and assuming the same total VOC emissions (ppbCC).

**Data availability.** The data supporting the findings of this study are available on reasonable request to the corresponding author.
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Author contributions
D.F.Z., A.B. and T.F.M. wrote the manuscript. T.F.M., A.K.S. and J.W. organized and designed the laboratory experiments. T.F.M. organized the PEGASOS Zeppelin Campaign. D.F.Z., A.B., R.T., E.K., C.W., J.W., T.F.M. conducted the laboratory data collection and analysis. D.F.Z., R.T., F.R. conducted ambient data collection and analysis. D.F.Z., A.B., A.K.-S., Y.R., J.W., T.F.M. edited the manuscript. All authors discussed the results and commented on the paper.

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