The summertime Boreal forest field measurement intensive (HUMPPA-COPEC-2010): an overview of meteorological and chemical influences

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1 Introduction

The Boreal forest ecosystem has a significant impact on the chemistry and physics of the Earth’s atmosphere. Presently it covers approximately 15 × 10\textsuperscript{6} km\textsuperscript{2} between 50 and 65° N, some 8 % of the Earth’s land surface and 27 % of the world’s forest (FAO 2010). The Boreal forest contains some 10 % of the total carbon in the terrestrial ecosystem and has no parallel in the Southern Hemisphere due to the absence of landmass in the corresponding southerly latitude band. The vegetation of the Boreal forest comprises mainly wind pollinated pine and spruce trees (Gymnosperms – e.g. *Pinus sylvestris*, *Picea abies*, and Angiosperms – e.g. *Betula sp.*, *Alnus incana*, *Populus tremula*) with low biodiversity in comparison to other forested areas such as tropical forests. These coniferous trees are known to emit significant quantities of reactive organic trace gases (e.g. monoterpenes) to the atmosphere as a function of temperature and to a lesser extent light (Rinne et al., 2009 and references therein). Rapid atmospheric oxidation of these biogenic trace gases and subsequent gas to particle conversion, have been shown to sustain particle concentrations of 1000–2000 particles/cm\textsuperscript{3} with low biodiversity in comparison to other forested areas such as tropical forests. These coniferous trees are known to emit significant quantities of reactive organic trace gases (e.g. monoterpenes) to the atmosphere as a function of temperature and to a lesser extent light (Rinne et al., 2009 and references therein). Rapid atmospheric oxidation of these biogenic trace gases and subsequent gas to particle conversion, have been shown to sustain particle concentrations of 1000–2000 particles/cm\textsuperscript{3} in the climatically relevant size range of 40–100 nm (Tunved et al., 2006, Kulmala et al., 2004). Previous studies from Boreal forest sites have indicated that some 12–50 % of aerosol mass and 50 % of cloud condensation nuclei (CCN) originates from forest sources (Tunved et al., 2008; Sihto et al., 2010) and that particle formation over the Finnish boreal forest causes a local radiative perturbation of between −5 and −14 Wm\textsuperscript{−2} (global mean −0.03 to −1.1 Wm\textsuperscript{−2}) (Kurten et al., 2003). It has been suggested that these cooling effects can offset the snow-vegetation albedo effect of the forest (Spracklen et al., 2008).

The enormous extent of the Boreal forest means that reactive organic emissions from such regions have the potential to reduce the Earth’s oxidation capacity (by the regional depletion of key oxidants OH and O\textsubscript{3}), and alter the Earth’s radiative budget by particle formation and subsequent scattering, absorption or reflection of light (either directly or via the role of aerosol particles as cloud condensation nuclei). Yet our understanding of atmospheric chemistry of the Boreal forest remains incomplete in key areas, in particular with regard to: the emission rate and speciation of volatile organic compounds; the extent of OH recycling during the oxidation process (e.g. Lelieveld et al., 2008), and in the precise formation and growth mechanisms of secondary organic aerosol, SOA (e.g. Kulmala et al., 2004; Kulmala and Kerminen, 2008). Indeed recent global SOA budgets, to which the Boreal forest is a major contributor, differ by an order of magnitude (Kanakidou et al., 2005; Goldstein and Galbally, 2007; Halquist et al., 2009). In future the Earth is predicted to warm, and Boreal regions more than any other (2–10°C) by 2100 (IPCC 2006). In order to predict the future impact of the Boreal forest on the atmosphere, and thereby to improve assessment of potential climatic feedbacks (e.g. Carslaw et al., 2010), extensive, high quality, and comprehensive datasets must be generated for detailed analysis. In this regard summertime assessments are particularly useful.

This paper provides an overview of the Boreal forest field measurement intensive HUMPPA-COPEC 2010 (Hyytiälä United Measurement of Photochemistry and Particles – Comprehensive Organic Particle and Environmental Chemistry). The measurements took place at the Boreal forest field station in Hyytiälä, Finland (Latitude 61° 51’ N; Longitude 24° 17’ E, elevation 181 m a.s.l., see Fig. 1) from 12 July–12 August 2010. To date, most intensive measurement campaigns performed at this station have been made in spring or autumn, to coincide with periods of most frequent particle nucleation events. In contrast, HUMPPA-COPEC 2010 was focused on characterizing summertime conditions when emission fluxes of volatile organic compounds (VOC), OH initiated photochemistry, and particle growth rates reach their seasonal maximum. In the course of the campaign, an international consortium quantified a comprehensive suite of gas and particle phase species (see Table 1). This was the largest campaign staged in the 100 year history of the Hyytiälä forestry station and this paper provides an overview of the campaign objectives, the instrumentation deployed, a site description, a regional vegetation analysis, a meteorological analysis, and an assessment of the main emission sources impacting the site.

2 Objectives

The general objective of the HUMPPA-COPEC 2010 campaign was to comprehensively characterize the atmospheric physics and chemistry over a Boreal forest site in summer. The more specific objectives are summarized below. Detailed analyses addressing these goals are provided in the accompanying papers of this special issue.

2.1 Volatile organic compound characterization

The first objective of the HUMPPA-COPEC 2010 campaign was to comprehensively characterize the fluxes, concentration ranges, and composition of VOCs emitted from the Boreal forest in summer. Earlier studies of specific monoterpenes at the Hyytiälä site have determined distinct seasonal variations in VOC concentration, with much higher levels in summer (Hakola et al., 2003). Boreal forest VOC emissions occur in numerous highly reactive, reduced forms such as isoprene (C\textsubscript{5}H\textsubscript{8}), monoterpenes (C\textsubscript{10}H\textsubscript{16}),
Fig. 1. This Fig. is a composite of (a) an aerial view taken during the campaign of the Hyytiälä site looking northwest, (b) a map of the bioclimatically defined Boreal region of Europe, and (c) a local map of the site showing locations of instruments and towers (see also Table 1).

and sesquiterpenes (C\textsubscript{15}H\textsubscript{24}) (Tarvainen et al., 2005; Hakola et al., 2006; Aaltonen et al., 2011). Additional non-terpenoid emissions or photochemical products (e.g. oxygenated VOCs) are generally less well characterized (Rinne et al., 2009), although some have been quantified at the Hyytiälä site (Janson et al., 1999; Janson and Serves, 2001; Rinne et al., 2007). One aim of the HUMPPA-COPEC campaign was to improve speciation of VOCs by applying new techniques (e.g. SPME – solid phase micro extraction) and more instrumentation than has been deployed before (see, Table 1). The main focus was on ambient air VOC measurements, which were made both within, and at various heights above the canopy to provide both a spatial and temporal assessment. In addition to individual VOCs, measurements of total OH reactivity were made by the comparative reactivity method (Sinha et al., 2008, 2010) in order to assess the extent to which VOCs are being quantified by the methods applied. Results are presented by Nölscher et al., 2011a within this special issue and these show average summer daytime OH activities to be ca. 12 s\(^{-1}\) with significant fractions (\(\sim 40\%\)) unassigned. Furthermore, two alternative instruments for measuring reactivity (Proton Transfer Reaction Mass Spectrometer, PTR-MS and Gas Chromatography with Photo Ionization Detector, GC-PID) were compared during the campaign (Nölscher et al., 2011b). Branch cuvettes were employed to speciate and quantify the VOC emissions from selected tree chemotypes. This was particularly important for the determination of very reactive compounds (e.g., alpha-terpinene, beta-myrcene) or sesquiterpenes which react with ozone on the order of seconds and are thus difficult to quantify in ambient air.

Continuous, on-line measurements of monoterpenes and oxygenated VOCs from dynamic branch and soil enclosures were conducted with PTR-MS. The comprehensive measurement system at SMEAR II enables comparison with other tree physiological parameters, such as photosynthesis and transpiration, and analyzing connections between the physiological status of plant and its VOC emissions. The on-line measurements from soil enclosures provide information on additional sources for VOCs from biotic or abiotic soil processes.

2.2 Radical chemistry and recycling

Many of the VOCs emitted from the Boreal forest can react rapidly with atmospheric oxidants (OH, O\textsubscript{3}, NO\textsubscript{3}). Determining which oxidizing species was most important for initiating reactions with Boreal forest VOC, and the extent of any chemical recycling was the second objective of this campaign. Recent field studies in tropical rainforest regions (Lelieveld et al., 2008; Hewitt et al., 2010) have indicated that the primary atmospheric oxidant (OH) is not simply consumed by reaction with VOCs but rather recycled to some degree. Plausible OH recycling mechanisms have also been derived from theoretical and laboratory studies of the molecule isoprene (Peeters et al., 2009; Paulot et al., 2009). This recycling is thought to be important in maintaining the
Table 1. The measurements made during the HUMPPA-COPEC intensive are tabulated alongside, the instrument type, time resolution, detection limit, responsible institute and location of the inlet (see also Fig. 1).

| Gas Phase Measurement | Time resolution | Detection Limit* | Technique         | Institution | Inlet Location (Fig. 1) |
|-----------------------|-----------------|------------------|-------------------|-------------|-------------------------|
| m28, m31, m33, m45, m59, m69, m71, m79, m93 | 5 min | \( \sim 50 \text{ ppt} \) | PTR-MS with cold trap | MPI | HUMPPA tower (24 m) |
| Total reactivity to OH | 1 min | 3 s⁻¹ | Reactivity (PTR-MS) | MPI | HUMPPA tower (24 m & 18 m) |
| Total reactivity to OH | 2 min | 3 s⁻¹ | Reactivity (GC-PID) | MPI | HUMPPA tower (24 m & 18 m) tower |
| Chiral monoterpenes + isoprene | 60 min | 1 ppt | GC-MS | MPI | HUMPPA tower (24 m) |
| Chiral emissions C_{10}H_{16} & C_{15}H_{24} | 60 min | 1 ppt | Cuvette – SPME - GC-MS | MPI | VOC tower (16 m) |
| Alkanes and alkenes | 5.5 min | \( \sim 1 \text{ ppt} \) | GC-MS | MPI | Mast (16 m/4 m) |
| Methane | 70 s | 0.01 ppm | GC-FID | MPI | Mast (4 m) |
| Speciated PANs | 3.5 s | 30 ppt⁶ | CIMS | MPI | HUMPPA tower (24 m) |
| NO₃ | 5 s | 2–4 ppt⁷ | CRD | MPI | HUMPPA tower (24 m) |
| N₂O₅ | 5 s | 5–10 ppt⁷ | CRD | MPI | HUMPPA tower (24 m) |
| NO₂ | 5 s | 20–100 ppt⁷ | CRD | MPI | HUMPPA tower (24 m) |
| \( \Sigma \text{PANs} \) | 4 s | 50–100 ppt⁷ | CRD | MPI | HUMPPA tower (24 m) |
| OH and HO₂ | LiF | MPI | White container (UH) & HUMPPA (24 m) tower |
| NO₂ | | | White container (UH) & HUMPPA (24 m) tower |
| NO, NO₂ | | | HUMPPA tower (24 m) |
| O₃ | 3 s | \( \sim 1 \text{ ppb} \) | UV | MPI | HUMPPA tower (24 m) |
| CO | 1 s | \( \sim 1 \text{ ppb} \) | UV | MPI | HUMPPA tower (24 m) |
| CH₂O | | | Hantzch | MPI | HUMPPA tower (24 m) |
| H₂O₂ and Organic Peroxides | | | Dual enzyme | MPI | HUMPPA tower (24 m) |
| \( J_{\text{NO}_2}, J_{\text{O}_1\text{D}} \) | | | Radiometer | MPI | HUMPPA tower (24 m) |
| Speciated peroxy radicals | | | LC-MS | NUCR/MPI | HUMPPA tower (24 m) |
| HONO (tower) | | | LOPAP | MPI | HUMPPA tower (24 m) |
| HONO (forest floor) | | | LOPAP | MPI | Ground – Forest floor |
| Total carbon | | | Oven – Licorr | MPI | HUMPPA tower (24 m) |
| terpenes | 30 min | \( \sim 10 \text{ ppt} \) | Cartridge – GC-MS | MPI | HUMPPA tower (24 m) |
| CO₂ | 6 min | \( \pm 3 \text{ ppmv} \) | IRGA⁸ | UH | Mast (4.2, 8.4, 16.8, 33.6, 50.4, 67.2 m) |
| H₂SO₄ | 30 s | \( 5 \times 10^4 \text{ cm}^{-3} \) | CIMS | UH | White container UH |
| OH | 30 s | \( 5 \times 10^4 \text{ cm}^{-3} \) | CIMS | UH | White container UH |

oxidation efficiency over the tropical rainforest where isoprene emissions dominate. However, over the Boreal forest where isoprene emissions are low and monoterpenes more abundant than isoprene, the recycling efficiency remains unknown. Simultaneous measurements of OH production from the photolysis rates \( J_{O_1D}, J_{HCHO}, \) and \( J_{HONO} \), in situ OH,
and OH reactivity (the total OH sink) were made to examine the budget. Whilst OH is generally considered to be the dominant oxidant of many VOCs during the day, NO$_3$ and O$_3$ persist through the night when they contribute substantially to, or even dominate VOC removal. For comparison, NO$_3$, and O$_3$ measurements were made at the same location (see Fig. 1 and Table 1).

![Image](image_url)

Table 1. Continued.

| Gas Phase Measurement | Time resolution | Detection Limit* | Technique | Institution | Inlet Location (Fig. 1) |
|-----------------------|-----------------|------------------|-----------|-------------|-------------------------|
| m$_{31}$, m$_{33}$, m$_{42}$, m$_{45}$, m$_{47}$, m$_{59}$, m$_{61}$, m$_{69}$, m$_{71}$, m$_{73}$, m$_{79}$, m$_{81}$, m$_{83}$, m$_{85}$, m$_{87}$, m$_{93}$, m$_{99}$, m$_{101}$, m$_{103}$, m$_{113}$, m$_{137}$, m$_{141}$, m$_{153}$, m$_{155}$, m$_{169}$, m$_{205}$, m$_{263}$ | 6 min | ~ 50 ppt | PTR-MS Profile | UH | Mast (4.2, 8.4, 16.8, 33.6, 50.4, 67.2 m) |
| Total reactivity | data very limited | PTR-MS Flux | UH | Radiation tower (20 m) |
| m$_{31}$, m$_{33}$, m$_{37}$, m$_{45}$, m$_{59}$, m$_{69}$, m$_{81}$, m$_{87}$, m$_{99}$, m$_{101}$, m$_{113}$, m$_{137}$ | ~ 10 s | ~ 50 ppt | GC-MS | FMI | Cuvette near sawmill |
| Organic amines | data very limited | Impregnated filter- LC-MS IRGA$^c$ | FMI | Cabin (2.5 m), close to sawmill container near sawmill |
| H$_2$O | 6 min | ±0.3 ppt hv | UV | Mast (4.2, 8.4, 16.8, 33.6, 50.4, 67.2 m) |
| SO$_2$ | 6 min | 0.1 ppb | UV | Mast (4.2, 8.4, 16.8, 33.6, 50.4, 67.2 m) |
| O$_3$ | 6 min | 1 ppb | UV | Mast (4.2, 8.4, 16.8, 33.6, 50.4, 67.2 m) |
| NO | 6 min | 0.1 ppb | CLD | Mast (4.2, 8.4, 16.8, 33.6, 50.4, 67.2 m) |
| NO$_x$ | 6 min | 0.1 ppb | CLD | Mast (4.2, 8.4, 16.8, 33.6, 50.4, 67.2 m) |
| CO | 6 min | 20 ppb | IRGA | Mast (4.2, 8.4, 16.8, 33.6, 50.4, 67.2 m) |
| O$_3$ flux | 30 min | Eddy covariance – CLD | UH | Radiation tower (23 m) |
| CO$_2$ flux | 30 min | Eddy covariance – IRGA | UH | Mast (23 m) |
| H$_2$O flux | 30 min | Eddy covariance – IRGA | UH | Mast (23 m) |
| CO$_2$ flux pine branches & aspen leaves, pine stem, soil | Variable (15 min...3 h) | Cuvette – IRGA | UH | Photo-tower (16 m) |
| H$_2$O flux pine branches & aspen leaves, soil | Variable (15 min...3 h) | Cuvette – IRGA | UH | Photo-tower (16 m) |
| Aerosol particle mass | 3 days | PM$_{10}$, PM$_{2.5}$, PM$_1$ | Dekati cascade impactor | Aerosol cottage |
| O$_3$ flux, pine branches & aspen leaves | Variable (15 min...3 h) | Cuvette – UV | UH | Photo-tower (16 m) |
Table 1. Continued.

| Gas Phase Measurement | Time resolution | Detection Limit* | Technique | Institution | Inlet Location (Fig. 1) |
|-----------------------|----------------|------------------|-----------|-------------|------------------------|
| NO flux, pine branches & aspen leaves | Variable (15 min...3 h) | Cuvette – CLD | UH | Photo-tower (16 m) |
| NOx flux, pine branches & aspen leaves | Variable (15 min...3 h) | Cuvette – CLD | UH | Photo-tower (16 m) |
| m33, m45, m59, m69, m79, m81, m99, m101, m137, m153 flux (pine branches & aspen leaves) | Variable (15 min...3 h) | ~ 50 ppt | Cuvette - PTR-MS | UH | Photo-tower (16 m) |
| m33, m45, m59, m69, m79, m81, m99, m101, m137, m153 flux (soil) | Variable (15 min...3 h) | ~ 50 ppt | Cuvette – PTR-MS | UH | watershed, ground |
| mono & sesquiterpenes (herbivores; pine branches) | 2–3 times | 10–80 ng m$^{-3}$ | Cuvette – Tenax - GC-MS | UH | VOC tower (16 m) |

**Aerosol Phase Measurement**

| Total particle number concentration | Time resolution | Detection Limit/size limit | Technique | Institution | Inlet Location |
|------------------------------------|----------------|---------------------------|-----------|-------------|---------------|
| 1 min                              | 10 nm          | CPC 3022                  | UH        | Aerosol cottage |

| Aerosol number size distribution | 10 min | 3–1000 nm | Twin-DMPS | UH | Aerosol cottage |
| Aerosol number size distribution | 10 min | 0.5–10 µm | TSI-3320 APS | UH | Aerosol cottage |

| Particle concentration + size info | 30 s | 1.5 nm | PH-CPC | UH | Aerosol cottage |
| Total particle concentration >1.3 nm | 1s | 1.05 nm | PSM | UH | White container UH/Aerosol cottage |
| Aerosol particle number concentration | 10 min | 0.03–10 µm particle sizes, 12 channels | Dekati ELPI | UH | Aerosol cottage |
| Pollen | 1 h | 1 pollen grain | Spore trap | Univ. Huelva | HUMPPA tower |
| Organic acids | 1 min | N/A | (−)-APCI-MS | Univ. Mainz | Main SMEAR cottage, new side |
| Gly/Mgly | 8–24 h | 0.05–0.16 ng m$^{-3}$ | Filter-LC-MS | Univ. Mainz | Radiation tower near REA cottage |

**Chemical composition**

| Particle chemical composition | 5 min | ~0.01 µg m$^{-3}$* | AMS | UH | Main SMEAR cottage, new side |
| Particle chemical composition | 50 pg * | Laser AMS | UH | Main SMEAR cottage, new side |

### 2.3 In situ comparison of OH measurement techniques

Two main methods of determining ambient OH concentrations on-line are in use today. One is based on Laser Induced Fluorescence (LIF) (e.g. Martinez et al., 2010), and the second uses Chemical Ionization Mass Spectrometry (CIMS) (e.g. Petäjä, et al., 2009). The former technique has the advantage of simultaneously measuring HO$_2$ which is itself an important photochemical species, whereas the latter also measures gaseous H$_2$SO$_4$, which is a key species in particle production. Although these two techniques have been previously compared for measuring OH in chamber studies (Schlosser et al., 2009; Fuchs et al., 2010), direct field comparisons are rare. Prior to the campaign, daytime OH measurements from the LIF and CIMS techniques were expected to be in good agreement, while interferences to the LIF measurement were expected to cause deviation only at night (Ren et al., 2003; Schlosser et al., 2009). When anomalies...
Table 1. Continued.

| Aerosol phase measurement | Time resolution | Detection Limit* | Technique               | Institution | Inlet Location (Fig. 1) |
|---------------------------|-----------------|------------------|-------------------------|-------------|------------------------|
| Organic acids             | 8–24 h          | N/A              | Filter-LC-MS            | UMainz      | Radiation tower near REA |
| Submicron organic mass and functional group composition | 8 h | 0.1.0.4 µgm⁻³ | Filter-FTIR            | UCSD        | White container UH (4 m) |
| K,Br,S, Al,V,Fe,Cu,Zn,Se | 8 h             | 1–100 ngm⁻³      | Filter XRF              | UCSD        | White container UH (4 m) |
| black carbon              | 10 min          | 5 ng m⁻³         | Magee Scientific Aethalometer | UH         | Aerosol cottage         |
| Ion and total size distribution 0.8–40 nm | 2 min | ~100/cm⁻³ * | ANAIS                   | UH          | White container UH (4 m) |
| Elemental and organic carbon | 4 h (2 m³) | EC 0.001 µgm⁻³ | Sunset EC/OC analyzer | UH          | white container         |
| Cl, NO₃, SO₄, Na, NH₄, K, Ca, Mg in PM₂.₅ and PM₁₀ | 60 min | 0.005–0.01 µg m⁻³ | MARGA               | FMI         | Cabin (2.5 m), close to sawmill |
| Biogenic organic acids    | 2–3 days        | Filter-LC-MS     | FMI                     | container close to sawmill |
| Aerosol properties        |                 |                  |                         |             |                        |
| Cloud condensation nuclei concentration total + size segregated | 1 s/10 min | Droplet Measurement Technologies | CCN-100 | UH | Aerosol cottage |
| Particle volatility       | 15 min          | > 20 nm          | VDMPS                   | UH          | Aerosol cottage         |
| Particle volatility and hygroscopicity | 20 min | N/A               | VHTDMA                  | UH          | Aerosol cottage         |
| Total particle concentration > 3 nm, solubility to water and butanol | 10 s | 3 nm              | CPCB                    | UH          | Aerosol cottage         |
| Optical properties        |                 |                  |                         |             |                        |
| Absorption coefficient    | 1 s             | VPSAP            | UH                      | Aerosol cottage |
| Scattering coefficients   | 2 Hz            | Vneph            | UH                      | Aerosol cottage |
| Aerosol scattering Particle volatility | 15 min 10 min | N/A               | TSI 3563 Nephelometer VDMPS | UH | Aerosol cottage |
| Atmospheric ions          |                 |                  |                         |             |                        |
| Ion mass spectrum         | 5 s             | 1/cm³ per 5 min  | API-ToF 1               | UH          | White container UH (4 m) |
| Ion mass spectrum         | 5 s             | 1/cm³ per 5 min  | API-ToF 2               | UH          | White container UH (4 m) |
| Total ion concentration   | 15 s            | ~100/cm⁻³ *     | Gerdien                 | UH          | White container UH (4 m) |
| Radon progeny             | 1 h             | filter sampling & beta counting | FMI | aerosol cottage |
| ambient radiation         | 10 min          | Ionization chamber | FMI | aerosol cottage |

PTR-MS = Proton Transfer Mass Spectrometer; CIMS = Chemical Ion Mass Spectrometer; CRD = Cavity Ring Down; GC-M = Gas Chromatography-Mass Spectrometry; GC-FID = Gas Chromatography-Flame Ionization Detector; GC-PID = Gas Chromatography-Photo Ionization Detector; API-ToF = Atmospheric Pressure Ionization Mass Spectrometer; AMS = Aerosol Mass Spectrometer; CLD = Chemiluminescence detector; FTIR = Fourier Transform Infrared Spectroscopy; IRGA = Infrared absorption; UV = Ultraviolet absorption/fluorescence; CPCB = Condensation Particle Counter Batter; PH-CPC = Pulse-Height Analysis Condensation Particle Counter; PSM = Particle Size Magnifier; VHTDMA = Volatility-Hygroscopic Tandem Differential Mobility Analyzer; VPSAP = Volatility-Particle/Sort Absorption Photometer; Vneph = Volatility-Nephelometer; VDMPS = Volatility-Differential Mobility Particle Sizer; BSMA = Balanced Scanning Mobility Analyzer; ANAIS = Airborne Neutral Cluster and Air Ion Spectrometer; Gerdien = Gerdien counter; APCI-MS = Atmospheric Pressure Chemical Ionization Mass Spectrometer; Gly/MGly = Glyoxal/Methylglyoxal; HPLC/ESI-MS = High Performance Liquid Chromatography/Electrospray Ionization Mass Spectrometer; LOPAP = Long Path Absorption Photometer; * at the time resolution in column 2. a for CH₃C(O)NO₂ varied in the course of the campaign.
between the OH measurements occurred, it was planned to exploit the comprehensive VOC and particle measurements to deduce possible causes. Thus a third campaign objective was set to perform a side-by-side comparison of OH in the VOC rich Boreal forest air and determine the identity of potential interferents. These instruments measured together at ground level (Univ. Helsinki white container, see Fig. 1) to allow for comparison and subsequently the LIF instrument was moved to the top of the HUMPPA tower. In this way vertical gradients in OH could be assessed. The OH intercomparison revealed significant differences between the LIF and CIMS OH measurement, the former being up to a factor of ten higher (Novelli et al., 2011).

2.4 Connections between VOC chemistry and Secondary Organic Aerosol formation, growth and ageing

New particle events have been shown to be a regular occurrence at Boreal forest sites (Kulmala et al., 2001; Tunved et al., 2003). The mechanism of new particle formation and initial growth is currently not fully understood but thought to involve both H$_2$SO$_4$ and possibly also low volatility VOC oxidation products in the early stages (Kulmala et al., 2004). However, also the relative contribution of BVOC oxidation products to the growth of existing particles (i.e. regardless if they are involved in nucleation or early cluster growth) is an area of considerable uncertainty. Understanding the formation and growth of new particles requires knowledge of atmospheric chemistry, aerosol dynamics and meteorology all of which were measured with novel instrumentation during the HUMPPA-COPEC 2010 campaign. Atmospheric oxidation of the VOCs emitted by the Boreal forest leads to a suite of less volatile organic products. These may condense on existing particles or under certain circumstances form new particles. The co-located, simultaneous measurement of VOC composition, OH-reactivity and physical (number, size and growth rates) and chemical (state and evolution of chemical composition) properties of aerosol particles allows potential links between BVOC oxidation and aerosol to be elucidated. Furthermore, the relationships between specific gas phase species and aerosol phase ions concentrations can be examined for causal links.

FTIR spectroscopy was used to quantify nearly 100% of the organic mass as aliphatic groups, carboxylic acid groups, organic hydroxyl groups, primary amine groups, and non-acid carbonyl groups (Russell et al., 2009). Organic functional groups in combination with elemental concentrations (measured by XRF) and other chemical measurements have been used to associate the functional group composition of the aerosol with biogenic and biomass-burning sources in other Boreal forests (Schwartz et al., 2010; Bahadur et al., 2010; Takahama, et al., 2011; Russell et al., 2011). However, the HUMPPA-COPEC campaign additionally provided the opportunity to associate these functional groups on sub-micron aerosol with specific biogenic precursors.

A comparison of vibrational sum frequency generation spectra of organic particles collected during HUMPPA-COPEC to those from the Tropics has shown marked differences (Ebben et al., 2011). In the Boreal forest the secondary organic aerosol is similar in composition to aerosol generated from α-pinene.

2.5 Identification of the crucial molecules in atmospheric nucleation

Our knowledge on the formation process for very small particles or clusters (sizes 1–2 nm) is still rather limited (see Kulmala et al., 2007; Kulmala and Kerminen, 2008). The question of which trace gases are involved in the atmospheric nucleation processes remains controversial within the aerosol community. It is likely that different nucleation mechanisms are at work under different conditions. The main candidates for nucleation in the troposphere have been thought to be: binary sulphuric acid – water (Noppel et al., 2002; Vehkamäki et al., 2002), ternary sulphuric acid – water – ammonia mixtures (Coffman and Hegg, 1995; Napari et al., 2002), ion induced nucleation of sulphuric acid and water (Lovejoy et al., 2004), activation of small clusters containing one sulphuric acid molecule (Kulmala et al., 2006), kinetic type, where critical clusters are formed by collisions of sulphuric acid molecules or other molecules containing sulphuric acid (McMurry and Friedlander, 1979) or nucleation by activation of organic molecules (Bonn et al., 2008). The comprehensive aerosol and gas-phase measurements provide the possibility to compare theoretical with measured nucleation rates (e.g. Boy et al., 2011) and to develop new parameterizations.

2.6 Chiral signatures

A sixth objective of this study was to fully investigate the chiral signature (enantiomeric ratios, distribution and temporal dependence) of the Boreal forest emissions. Typically these non-superimposable, mirror image forms of chemical species (stereoisomers such as (+)-α-pinene and (−)-α-pinene) are measured and modeled together in ambient gas phase studies as they react with the same rates with ozone and OH. However, the two enantiomers are biochemically distinct molecules which may elicit diverse responses in pollinators and predators within the biosphere. Previous studies have established that clear regional chiral signatures exist, and that ambient mixing ratios of α-pinene enantiomers over the Boreal forest always favours (+)-α-pinene (Williams et al., 2007). Furthermore, mechanical damage to the needles of the Scots pine has been shown to induce higher ratios of the (+) enantiomer (Yassaa and Williams, 2005, 2007), a finding that has proved useful in identifying emissions from a nearby sawmill (Eerdekens et al., 2009). Recently laboratory studies have shown that ozone reacting
with stereoisomers on the surface of aerosol particles coated with chiral semivolatile organic compounds may depend on stereochemistry and that the differences in chemical accessibility could lead to the enrichment of one oxidation product stereoisomer over the other in ambient air (Stokes et al., 2009). To investigate whether such reactions impacted ambient enantiomeric distributions during HUMPPA-COPEC the enantiomeric monoterpenes were examined directly from the chamber (the emission signature, see Yassaa et al., 2011), from the ambient gas phase measurements (subject to photochemistry), and on the particle phase (by vibrational sum frequency and second harmonic generation analysis, see Ebben et al., 2011).

2.7 Photochemistry and aerosol dynamics in biomass burning and city plumes

While located in a remote, background area, the SMEAR II station can also be used for analyzing anthropogenic impacts on air composition. European Boreal forest regions in summer are prone to anthropogenic influences, both from biomass burning which commonly occurs in Boreal regions at this time of year, and from pollution plumes emerging from large urban centers (e.g. Helsinki, St. Petersburg). Both sources produce NOx (NO and NO2) and VOC, and thus can produce ozone photochemically during transport. Therefore the HUMPPA-COPEC campaign aimed to identify and characterize such plumes using back trajectories, as well as aerosol particle and trace gas signatures. In particular species such as formaldehyde, peroxy acyl nitrates (PANs), and peroxyacids were targeted in this regard. Furthermore, it was hoped to assess the response of the forest to elevated anthropogenic ozone levels in such periods, in particular through analysis of ozone dry deposition flux measurements in relation to photosynthesis and micrometeorological measurements.

2.8 Dependence of diel OH and its precursors on meteorology

The HUMPPA-COPEC 2010 campaign offers a unique opportunity to investigate the impact of physical and meteorological parameters on OH and its related precursors within the boreal forest environment. By analyzing the surface characteristics (fluxes of heat, moisture, momentum and reactive species), the boundary layer dynamics, the evolution of the thermodynamic variables, UV-radiation field and the chemical pathways, the processes controlling the diurnal variability of OH can be determined. Previous studies in tropical regions (Ganzeveld et al., 2008 and Vilà-Guerau de Arellano et al., 2011) have shown the importance of the night-day transition, the onset of biogenic emissions and the exchange of the residual layer (free troposphere) with the boundary layer in controlling the maximum concentrations of NO, isoprene and OH.

In order to achieve the objectives set in Sect. 2.2 and 2.7, modeling tools were applied (box-, 1-D and large-eddy simulation models) and the capabilities of these tools in reproducing the main features of the diurnal variability of OH and biogenic compounds were assessed. The subsequent sensitivity analyses then allowed for a determination of the different roles of the boundary layer dynamics, emissions/deposition and chemistry in the OH-reactivity. The evolution and the structure of the atmospheric boundary layer and the turbulent mixing inside are important parameters controlling the chemical reactions and aerosol dynamics usually observed at ground level. The main height at which new particles form remains controversial and recent findings in atmospheric modeling suggest that reactive organic compounds emitted from the canopy or soil with a strong gradient in the entrainment zone at the top of the atmospheric mixed layer are needed to explain the observed vertical distribution (Boy et al., 2011). Both nucleation events and failed events during HUMPPA-COPEC will be examined to this end.

2.9 Atmospheric nitrogen budget for the Boreal forest

A comprehensive suite of nitrogen containing compounds in the air was measured during the HUMPPA-COPEC campaign. In the gas phase NO, NO2, N2O5, NO3, HNO3, PANs, NH3 and HONO were quantified, while in the aerosol phase nitrate (NO3-) and ammonium (NH4+) were also monitored on-line. The instruments used for each species are detailed in Table 1. How nitrogen species are processed over the Boreal forest is examined in detail in this special issue, in particular with regard to interactions with oxidants such as OH and O3.

3 Instrumentation

The Hyytiälä station (SMEAR II) is one of three stations operated throughout the year by the University of Helsinki (Hari and Kulmala, 2005). Several key instruments are run with a high degree of automation at the SMEAR II-Hyytiälä site throughout the year (e.g. O3, CO2, CO, meteorological parameters) and the data made available to the public via the SMEARsite (http://www.atm.helsinki.fi/~junninen/smartSearch), details of which are provided by Junninen et al., 2009. During the HUMPPA-COPEC 2010 campaign, the standard suite of measurements was extended between July and August. A full list of instruments operated during the HUMPPA-COPEC campaign is given in Table 1.

The location of each instrument and its inlet can be seen in Table 1 and Fig. 1. Many of the instruments associated with the intensive period positioned inlets on the HUMPPA tower (24 m), extending approximately 2–3 m above the canopy top. For photochemical studies it is essential that measurements of reactive species (e.g. OH, CH2O, and monoterpenes), and their photochemical source (JNO2, JO1D) and sink terms (OH reactivity, O3, NO3) be collocated due to
their inherently high spatial variability. Furthermore, close proximity of OH and reactive organic inlets is essential when investigating intensity of segregation effects that can effectively reduce the overall rate of isoprene oxidation due to inhomogeneous mixing (e.g. Butler, 2008; Pugh et al., 2011). These instruments were situated preferentially around the HUMPPA tower in a 3-container stack, and MPI white container. Due to space constraints most aerosol instruments were located at the main aerosol hut. Less reactive gas phase species (e.g. alkanes) and measurements with long integrating times (aerosol filters) were placed in alternative locations close by (main aerosol hut, MPI blue container) due to space constraints. All measurements were made within a radius of 300 m from the main mast, see Fig. 1. In the following analysis of regional influences on the site (see Sect. 5.0), all measurements are considered to be collocated.

The height of the forest canopy was generally 20–21 m though the land surrounding the site was slightly undulating so that the forest crown was not in one horizontal plane. From the 73 m mast (see Fig. 1), measurements of selected species (e.g. O$_3$, CO$_2$) as well as meteorological parameters were made at multiple heights (4 m, 8 m, 16 m, 33 m, 50 m, 67 m). Further vertical resolution of temperature and pressure was obtained through the regular launching of meteorological sondes to approximately 5 km (ca. 4 times per day). Several other instruments measured both above and below the canopy (e.g. OH reactivity and monoterpines.)

In addition to the in situ measurements of gas phase and aerosol species, several branch cuvette systems were employed to measure the emission fluxes of monoterpenes and sesquiterpenes from several chemotypes of pine and spruce near the VOC tower and additionally CO$_2$ uptake rates at the photo tower. Furthermore, from the main mast direct flux (disjunct eddy covariance and gradient method) measurements of several VOC species in air were performed using PTR-MS, while fluxes of species from the ground were also monitored via automated flux chamber analyses.

4 Meteorology

4.1 General

For the campaign period (July–August 2010) the Hyytiälä site was influenced by a high pressure region centered to the east of Finland and a sequence of weak frontal systems. Surface pressures varied between 980–1005 hPa, and due to the northern latitude there was approximately 18 h daylight per day. The temperatures measured at the site (at 16.8 m) in this period were anomalously high ($T_{\text{max}} = 32.4$, $T_{\text{min}} = 10.1$, $T_{\text{ave}} = 20.0^\circ$C). Indeed the highest temperature recorded anywhere in Finland for 100 years ($37.2^\circ$C) was registered during this campaign, albeit not at the Hyytiälä site. The measurements during these high temperature conditions may hold clues about the forest response to future climate change. Figure 2a shows the temperatures recorded at the Hyytiälä site for the campaign period over the past 6 years. The temperatures in 2005, 2006, 2007, 2008, and 2009 show a similar spread of temperatures (6–25$^\circ$C) while in 2010 temperatures exceeded 25$^\circ$C for several periods (e.g. 10–14 July, 27–30 July). The temperatures were not, however, anomalously high for the whole campaign. Cooler periods (e.g. 23–25 July, average temperature 14$^\circ$C) were recorded, in particular associated with air from the northwest. Interestingly, the levels of ozone for 2010 were also anomalously high (see Fig. 2b), the highest levels corresponding closely to the highest temperatures. Again, this may not be untypical for future conditions. Precipitation was generally low (<2 mm day$^{-1}$) except for 15 July, 27 July and 4 August when heavier rain
events (associated with frontal passages) were experienced at the site. On several occasions these rain events and associated lightning caused power disruption at the site.

### 4.2 Air mass origin

Three day air mass back trajectories were compiled for the duration of the HUMPPA-COPEC campaign. The trajectories were calculated every 3 h starting from 00:00 UTC, from a height of 25 m (a.g.l.), between 10 July and 12 August using the National Oceanic and Atmospheric Administration (NOAA) GDAS database, which applies the Hybrid Single-
Particle Lagrangian Integrated Trajectories (HY-SPLIT 4) program (Draxler and Rolph, 2011; Rolph, 2010). The three day trajectory endpoints were found to partition into three distinct sectors relative to the Hytiälä site (NW, SW and SE). This trajectory derived air mass origin information is summarized in Fig. 3a. For most of the 2010 campaign air advected to the site originated three days previously from the SW (53.7%), although significant periods of SE (20.7%) and NW (10.3%) air were also encountered. These trajectories correspond closely to the distribution of wind directions measured at 16.8 m shown in Fig. 3b. Figure 3b also includes wind direction data from the previous 5 years for comparison and it shows that 2010 had more southerly and less northwesterly influence than in previous years. Table 2 summarizes the time periods each sector influenced the site. Conditions generally remained steady for 2–3 days at a time, which is typical of synoptic scale frontal system influences mentioned in Sect. 4.1. Also included in Table 2a are the local and regional influences (e.g. biomass burning) observed during each time period. These are discussed in more detail below in Sect. 5.

In order to determine whether the anomalously high temperatures experienced during this campaign were related to anomalous circulation patterns, a similar trajectory analysis to that presented above for 2010 was also performed for the years 2005–2009 and the results presented in Table 3. For the years 2010, 2009 and 2007 southwesterly flow was dominant in summer. However, in 2006 and 2008 the prevailing wind direction was northwesterly which has been identified as the wind sector least impacted by anthropogenic pollution and thus most representative of pristine Boreal forest (Tunved et al., 2006). The year 2010 has a significantly larger proportion of trajectories from the south than all other years analyzed (74.4%) and this is likely the reason for the higher temperatures and ozone experienced in 2010. High O$_3$ levels (>120 µg m$^{-3}$) in several other Southern Finland stations of the National Air Quality measurement network (FMI, http://www.ilmanlaatu.fi/ilmanyt/lityykset/rajaarvovilystys.php?p=O$5$;SM8(\&lyys=2010), were also reported during July–August 2010, indicating long-range transport from Central Europe.

4.3 Atmospheric boundary layer characteristics

The measurements of turbulent fluxes of heat, moisture and momentum at the surface (SMEAR II tower) combined with the profiling of the thermodynamic variables enable us to characterize the atmospheric boundary layer (ABL) dynamics during the campaign. The vertical structure of the ABL during night and day was monitored by 175 radiosondes launched throughout the campaign. In order to obtain a detailed quantification of the evolution of the ABL, we scheduled 5 intensive observation days (24 h) with radiosonde launches every 2 h. From these observations (temperature, pressure, humidity), the evolution of the boundary layer height and the dynamic characteristics of the entrainment zone can be determined. In Fig. 4a and b an example is shown of the evolution of the vertical profiles of potential temperature ($\Theta$) and specific humidity ($q$) at 06:00, 10:00, 14:00 and 18:00 local time (LT) of 6 August 2010 (SW sector, see Table 2). At 06:00 LT a ground inversion (extending up to approximately 150 m) with a residual layer is clearly observed. The profile at 10:00 LT already shows a well mixed boundary layer extending up to 1100 m and capped with a thermal inversion (~2 K) at the same height. We note the well mixed profile of specific humidity at 14:00 and 18:00 LT, indicating the effectiveness of convective turbulence in mixing the species. For example, at 14:00 LT the free convection time scale (Stull, 1988) is estimated to be around 10 min.

Figure 4c shows the boundary layer height ($h$) estimated from the vertical profiles of potential temperature. Several criteria can be used to estimate $h$, but in this case the maximum potential temperature gradient near the inversion layer has been used. Note the strong temperature gradient at 2 km is not connected to the current boundary layer, but to advection and residual layers and is therefore not considered. As the figure shows, the boundary layer height typically grows from less than 200 m in the early morning to around 1700 m at the end of the afternoon. After the ground inversion has broken up (after 08:30 LT), the boundary layer connects with the residual layer aloft and is rapidly merged. After this, the boundary layer resumes its growth by entrainment. In spite...
Table 3. The percentage distribution within the sectors (NW, SW, SE, NE) are shown for the years 2005–2010.

| Year | South-West | South-East | North-West | North-East | Trajectories Used % |
|------|------------|------------|------------|------------|---------------------|
| 2010 | 53.7       | 20.7       | 10.3       | 0          | 84.7                |
| 2009 | 53.7       | 3.7        | 0.5        | 16.2       | 74.1                |
| 2008 | 29.4       | 0          | 34.6       | 14         | 78                  |
| 2007 | 44.1       | 18.4       | 16.9       | 2.2        | 81.6                |
| 2006 | 21.3       | 27.9       | 34.6       | 14.7       | 98.5                |
| 2005 | 29.4       | 18.4       | 23.5       | 0.7        | 72                  |

Fig. 5. Land use plots extracted from the Corine (2006) database for a 50 km and 5 km radius around the Hyytiälä site.

of the large sensible heat flux at the surface (maximum value of 350 W m\(^{-2}\) near noon), the final boundary layer height was less than expected, possibly due to subsidence and differences in horizontal heat advection between the boundary layer and free troposphere, which can be induced by horizontal differences in land use conditions. Model results show that not taking this into account leads to an overestimation of 600 m of the boundary layer height at 18 LT. Governing dynamic factors (surface fluxes, role of entrainment and advection, clouds) need to be considered in the data analysis since they influence the transport and mixing of the reactant species. A complete analysis of the boundary layer dynamics and its implications to the OH reactivity in the Boreal forest is provided elsewhere in this special issue.

5 Local and regional influences

5.1 Land use within 50 km and 5 km of the Hyytiälä station

Potential local and regional influences on measurements at the Hyytiälä station were assessed within a 50 km and 5 km radius of the site using the Corine land cover database (EEA Technical Report, 1997). The data were edited using the ArcGis 9.2 (Esri ©) Geographic Information Systems (GIS) software. These radii were chosen based on the typical summer atmospheric lifetimes of one of the main biogenic species namely \(\alpha\)-pinene. Assuming an OH concentration of \(1 \times 10^6\) and ozone \(1 \times 10^{12}\) molecules cm\(^{-3}\) (0.04 pmol mol\(^{-1}\) and 40 nmol mol\(^{-1}\)) the atmospheric lifetime of \(\alpha\)-pinene is approximately 2 h based on laboratory measured rate coefficients (Atkinson et al., 2006). Wind speeds measured at 68 m during the campaign varied between 0.1–7.0 ms\(^{-1}\) with an average of 2 ms\(^{-1}\). Thus the distances covered at these wind speeds within one \(\alpha\)-pinene lifetime approximate to 5 km and 50 km, respectively.

Figure 5 shows the distribution of land use within 50 and 5 km of the Hyytiälä site. On both scales the majority of the surrounding area is forested and this is consistent with the analysis of Finnish map information presented by Haapanala et al., 2007. There is one significant urban centre visible on the 50km plot namely Tampere (population 211 000), which was southwest of the site and therefore within the prevailing wind sector, see Sect. 4.2. Tables 4a and b give the area covered by specific land use types within 50 km and 5 km of Hyytiälä, respectively. The data is presented as a total and
as a function of the wind sectors identified in Sect. 4. It can be seen from the 50km scale analysis that coniferous forest and mixed forest account for over 65% of the land use surrounding Hyytiälä. Small or medium sized water bodies are also significant within the region comprising some 15% of the area within 50 km, which is representative of Finland. Closer examination of the individual sector analyses for the 50 km radius shows that the Northwest sector has the highest percentage of coniferous forest. This is in agreement with Tunved et al., 2006 who identified this sector as representing the pristine boreal forest with minimum anthropogenic influence. It should be noted, however, that wind only came from the sector for circa 10% of the campaign. The south west sector (corresponding to the prevailing wind) shows the largest percentage of urban fabric (4.3%), although this is still small in comparison to the combined total of coniferous and mixed forest (59%). The influence of anthropogenic sources is discussed further in Sect. 5.4. The dominance of the forest cover also extends to the southeast sector although here a somewhat higher percentage of agriculture and arable land is evident. Within 5 km of Hyytiälä the coverage due to coniferous and mixed forest increases to approximately 80% in all sectors. Inclusion of the category “transition woodland scrub”, which was regenerated forested land in the seedling or sapling stage, increases this coverage to over 90% in all wind sectors and to 94% of the overall land use within 5 km. An inspection of the areas designated by the Corine land cover database as coniferous forest, mixed forest and woodland scrub was performed within 5km of the site. Those areas identified as coniferous forests were found to be Scots pine (Pinus sylvestris L.) and Norway spruce (Picea abies L. [Karst.]) which have been identified previously as strong terpene emitters. Example forests of these trees, located within 5 km of the Hyytiälä site, are shown in Fig. 6. The most extensive “coniferous” species over Finland as a whole is Scots pine (Pinus sylvestris), although the Norway spruce (Picea abies) is also common. The areas classified as mixed forests, see Fig. 6, were found to include conifers and deciduous trees, mainly birches such as Silver birch (Betula pendula Roth) and lesser extent Downy birch (B. pubescens Ehrh). Woodland scrub included species such as willow (Salix sp.) and aspen (Populus sp.), which are both known to be emitters of isoprene (Hakola et al., 1998; Rinne et al., 2009).

5.2 Biomass burning

A consequence of the region-wide heatwave was that numerous devastating wildfires began mid July in the Nizdni-Novgorod region of the Russian Federation, some 400 km east of Moscow. The burning continued for one month, causing over 50 casualties and leaving thousands homeless. By August 18 an estimated 5.8 million ha had burned in the region (http://www.fire.uni-freiburg.de). The fires significantly impacted regions of dry peatlands but also extended to wheat and timber. The biomass burning emissions were advected over Moscow and beyond. In Fig. 7 it can be seen that clear biomass burning signals were detected at the SMEAR II Hyytiälä station on three distinct occasions. These were: event 1–26 July, 00:00 and 28 July, 00:00; event 2–28 July, 12:00 and 30 July, 12:00; event 3–7 August, 12:00 and 9 August, 18:00, with all times in local time (UTC + 3). Acetonitrile (CH$_3$CN), a specific marker for biomass burning (Lobert et al., 1990) and CO a general marker for combustion increased significantly during these periods. Additional ensemble type back trajectories for these periods indicated that the air arriving at the Hyytiälä site had passed over the burning affected Novgorod region. These incidences are also noted in Table 2.

5.3 Sawmill influence

In the course of a previous campaign in Hyytiälä (QUEST IV) the occasional influence of nearby sawmill activity at Korkeakoski (330 000 m$^3$ wood per year, 10 km SE from Hyytiälä) was noted (Eerdekens et al., 2009). In Korkeakoski the “UPM” sawmill was in operation from 2–13 August, the “JPJ” woodmill operated between 9–12 August and the Paahtopuu pellet factory operated between 12–13 August. This influence manifested itself as unusually high concentrations of monoterpenes and as a strong enrichment of (+)-α-pinene over its (−)-α-pinene enantiomer. These events were also associated with a southeasterly wind direction corresponding to the bearing of the mill from the site, see Fig. 5. Figure 8 shows the sum concentration of both α-pinene enantiomers and their ratio for the 2010 HUMPPA-COPEC campaign. On several occasions, peak concentrations are associated with enantiomeric ratio enrichments in (+)-α-pinene. These events coincide with short episodes of southeasterly winds suggesting that sporadic influences of the sawmill were detected during the campaign. Case studies of these events, which increase for short periods to an order of magnitude higher than average values, can help elucidate the efficiency of OH radical recycling as a function of monoterpenes concentrations. Two strong events were noted in the campaign: event 1, 18 July 2010, 01:00–06:45 LT, wind direction 114–122°, (−)/(+)α-pinene ratio 0.38–0.48; and event 2, 6 August 2010, 01:40–04:00 LT, wind direction 109–200°, (−)/(+)α-pinene ratio 0.35–0.49. These incidences are also noted in Table 2. Several other short transient events are noted: event 3, 4 August 2010, 09:45 LT; event 4, 3 August 2010, 20:10 LT; event 5, 3 August 2010, 23:25 LT.

5.4 Urban/anthropogenic influence

Clear anthropogenic influences on the Hyytiälä dataset can be deduced from the SO$_2$ and n-pentane traces shown in Fig. 9. Both species are anthropogenically emitted. A strong peak of SO$_2$, in excess of 2 nmol mol$^{-1}$, can be seen on 14 July and this corresponds to ensemble back trajectory
analyses (not shown) that have crossed St. Petersburg, Russia, a large industrialized area (population >4.5 million) located to the southeast of the site. Peaks are also seen in pentane at these times. Such strong peaks in SO$_2$ and pentane are also observed coincident with the biomass burning events 26–30 July. Again in these cases the air was advected over the StPetersburg region suggesting that this is a strong source of SO$_2$ and hydrocarbons to the atmosphere. In the clean air sector from the northwest SO$_2$ was below 200 pmol mol$^{-1}$. The incidences of high SO$_2$ are also noted in Table 2. On two occasions peaks in n-pentane were observed which did not correlate with SO$_2$ (e.g. 16 July and 6 August). This

Table 4. Land use within (a) 50 km and (b) 5 km of the Hyytiäliä site expressed in areas of activity/vegetation (m$^2$), extracted from the European Corine landcover database 2006, defined in GIS nomenclature for types with significant coverage.

| CLC code | Land use                          | 5 km radius | NW % | NE % | SW % | SE % |
|----------|----------------------------------|-------------|------|------|------|------|
| (a)      |                                   |             |      |      |      |      |
| 131      | MINERAL EXTRACTION SITES          | 799736      | 1.02 |  -   |  -   |  799736 |
| 211      | NON IRRIGATED ARABLE LAND         | 979787      | 1.35 |    - |  -   | 979787  |
| 243      | AGRICULTURE AND NATURAL VEG.     | 1418096     | 1.81 | 376  | 1.91 | 1418096 |
| 312      | CONFEROUS FOREST                 | 4763427     | 60.65| 11035| 56.18| 4763427 |
| 313      | MIXED FOREST                     | 20093416    | 25.58| 5607 | 28.35| 20093416|
| 324      | TRANSITIONAL WOODLAND-SHRUB      | 5988018     | 7.62 | 1358 | 6.91 | 5988018 |
| 412      | PEAT BOGS                        | 2000000     | 0.00 |    - |  -   | 2000000 |
| 512      | WATER BODIES                     | 1624184     | 2.07 | 1265 | 6.44 | 1624184 |
| TOTAL    |                                  | 78537116    | 100.00| 19642| 100.00| 78537116|

(b)

| CLC code | Land use                          | 5 km radius | NW % | NE % | SW % | SE % |
|----------|----------------------------------|-------------|------|------|------|------|
| 112      | URBAN FABRIC                     | 15055842    | 1.92 | 8485 | 0.43 | 8485697 |
| 121      | INDUSTRIAL & COMERCIAL UNITS     | 23592576    | 0.30 | 319  | 0.02 | 3196448 |
| 122      | ROAD AND RAIL NETWORKS           | 1126126     | 0.01 |     -|  -   | 1126126 |
| 124      | AIRPORTS                         | 3149041     | 0.04 |     -|  -   | 3149041 |
| 131      | MINERAL EXTRACTION SITES         | 6341889     | 0.08 | 711  | 0.04 | 711848 |
| 132      | DUMP SITES                       | 579159      | 0.01 |     -|  -   | 579159  |
| 141      | GREEN URBAN AREAS                | 2378588     | 0.03 |     -|  -   | 2378588 |
| 142      | SPORT FACILITIES                 | 1834799     | 0.02 | 369  | 0.02 | 369541  |
| 211      | NON IRRIGATED ARABLE LAND        | 28085774    | 3.58 | 512  | 2.65 | 5121127 |
| 243      | AGRICULTURE AND NATURAL VEG.     | 45505805    | 5.57 | 8861 | 4.51 | 8862866 |
| 311      | BROAD-LEAVED FOREST             | 5106062     | 0.07 | 1371 | 0.07 | 1371976 |
| 312      | CONIFEROUS FOREST               | 29402478    | 37.44| 1050 | 53.51| 1050577 |
| 313      | MIXED FOREST                    | 22687578    | 28.89| 3656 | 18.67| 3655684 |
| 324      | TRANSITIONAL WOODLAND-SHRUB     | 466920775   | 5.95 | 9715 | 4.95 | 9715107 |
| 332      | BARE ROCKS                      | 253594      | 0.00 | 253  | 0.01 | 253594  |
| 411      | INLAND MARSHES                  | 5551665     | 0.07 | 832  | 0.04 | 832045  |
| 412      | PEAT BOGS                       | 21429312    | 0.27 | 703  | 0.36 | 703320 |
| 511      | WATER COURSES                   | 1000605     | 0.01 | 335  | 0.02 | 335384  |
| 512      | WATER BODIES                    | 121911642   | 15.52| 288  | 14.71| 288760 |
| TOTAL    |                                  | 785396079   | 100.00| 19640| 100.00| 7853960 |

Fig. 6. Densely forested areas within 5 km of the Hyytiäliä site: (a) Scots pine designated by the Corine database as “coniferous forest”, (b) Spruce, designated by the Corine database as “coniferous forest”, and (c) Birch and conifers, designated by the Corine database as “mixed forest.”
strongly suggests a separate source of the hydrocarbon influenced the site at this time.

5.5 Particles

Number concentrations for particles between 3–25 nm and 3–1000 nm are shown in Fig. 10a and b. The smaller size range represents the nucleation mode particles whereas the larger size range also includes the Aitken and accumulation modes (termed total particles in Fig. 10). Several elevated events can be seen in the nucleation mode particles. One example, on the 12 of July shows coincident peaks of nucleation mode and total particle concentrations. This is likely a pollution plume advected from southwest of the site possibly Tampere. In contrast a second event, on 23 July, began when total particles (3–1000 nm) were at a campaign minimum concentration at 06:00 local time, see Fig. 10b. Nucleation mode particles increased sharply in concentration over the following four hours reaching circa 30000 cm$^{-3}$. This was the only nucleation event seen at the site during the summer campaign. The generally high levels of total particles from 3–1000 nm observed in the summer campaign provided ample surface area for condensation of semi volatile gas phase species and thus inhibited nucleation. Several “failed events” were noted in which particles were observed to be nucleating but did not grow into the larger size ranges.

6 Conclusions

The overall aim of this campaign was to obtain a dataset characteristic of the Boreal forest in summer with which the specific campaign goals (see Sect. 2.1–2.10) can be addressed. During the HUMPPA-COPEC 2010 summer campaign the Hyytiala site was impacted by periods of extremely high temperature, advected biomass burning emissions, emissions from urban centers in the southwest and occasionally by a nearby sawmill. The latter has the effect of extending the VOC:NO$_x$ space measured in the campaign and was a valuable test of the OH reactivity measurements. Fortunately none of these various effects entirely dominated the field study period allowing contrasts between high and low temperatures from clean boreal forest emissions and polluted air masses to be made. Analysis of meteorological sonde data has shown that the boundary layer height, although variable during the campaign, increased typically from less than 200 m in the early morning to around 1700 m at the end of the afternoon. During the campaign air was advected mostly from the SW (53.7 %) of the time with smaller influences from the SE (20.7 %) and the NW (10.3 %). In comparison with the previous five years the campaign period in 2010 was more impacted by air from the south (SW and SE) leading to higher ozone and temperature values at the site. A regional...
Fig. 10. Number concentrations of particles are plotted for the size range 3–25 nm – (red) and 3–1000 nm (black). Measurements taken from the main mast at 16.8 m.

50 km wide surface cover analysis showed that the southern sectors (SW and SE) contain more anthropogenic influence than the NW sector, although coniferous forest was found to dominate in all sectors.

The unusually high temperatures experienced on this campaign are also useful to gauge the response of the Boreal forest to warmer conditions, at least in the short term. For example strong increases in highly reactive mono- and sesquiterpene emissions were observed during this campaign at higher temperatures, leading to higher OH reactivities. Temperatures in boreal regions are predicted to rise over this century, and datasets such as HUMPPA-COPEC 2010 can help verify proposed feedbacks to atmospheric properties. It should be noted, however, that over longer climate-scale periods ecological feedbacks as well as larger scale synoptic shifts may dominate.

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