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ExchanGE processes in mountainous Regions (EGER) – overview of design, methods, and first results

T. Foken^{1,g}, F. X. Meixner^{2,4}, E. Falge², C. Zetzsch^{3,g}, A. Serafimovich^{1,g}, A. Bargsten², T. Behrendt², T. Biermann¹, C. Breuninger², S. Dix^{1,a}, T. Gerken¹, M. Hunner^{1,a}, L. Lehmann-Pape², K. Hens², G. Jocher^{1,b}, J. Kesselmeier², J. Lüers^{1,g}, J.-C. Mayer², A. Moravek², D. Plake², M. Riederer¹, F. Rütz¹, M. Scheibe^{2,c}, L. Siebicke^{1,d}, M. Sörgel³, K. Staudt^{1,e}, I. Trebs², A. Tsokankunku², M. Welling², V. Wolff^{2,e}, and Z. Zhu^{2,f}

¹University of Bayreuth, Department of Micrometeorology, 95440 Bayreuth, Germany

²Max-Planck-Institute of Chemistry, Biogeochemistry Department, P.O. Box 3060, 55020 Mainz, Germany

³University of Bayreuth, Atmospheric Chemistry Research Laboratory, 95440 Bayreuth, Germany

⁴University of Zimbabwe, Department of Physics, P.O. Box MP 167, Mount Pleasant, Harare, Zimbabwe

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^anow at: TÜV Süd Industrie Service GmbH Wind Cert Services, Ludwig-Eckert-Straße 10, 93049 Regensburg, Germany

^bnow at: Alfred-Wegener Institute for Polar and Marine Research, Telegrafenberg A43, 14473 Potsdam, Germany

^cnow at: German Aerospace Center (DLR), Institute of Atmospheric Physics, Münchner Straße 20, 82234 Oberpfaffenhofen-Wessling, Germany

^dnow at: Institut national de la recherche agronomique (INRA), BP 709, 97387 Cedex Kourou, French Guiana

^enow at: Agroscope ART Research Station, Reckenholzstrasse 191, 8046 Zürich, Switzerland

^fnow at: Institute for Geographic Sciences and Natural Resources Research, Chinese Academy of Sciences (CAS), A11 Datun Road, Anwai, Beijing 100101, China

^gMember of Bayreuth Center of Ecology and Environmental Research (BayCEER), University of Bayreuth, 95440 Bayreuth, Germany

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Correspondence to: T. Foken (thomas.foken@uni-bayreuth.de)

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water vapour, and trace compounds at the *Waldstein-Weidenbrunnen* site. Our previous investigations on atmosphere-forest ecosystem coupling at the *Waldstein-Weidenbrunnen* site have been based on the mixing layer theory (Finnigan, 2000; Raupach et al., 1996). Our first approach defined coupling in relation to characteristic length scales of the distance between coherent eddies and the wind shearing (Wichura et al., 2004; Wichura, 2009). Later, a wavelet transform based technique (Thomas and Foken, 2005, 2007b) has been used, which had been developed for the WALDATEM-2003 experiment (Thomas and Foken, 2005, 2007b). There, the mean temporal scales of coherent structures were estimated via the fitting of a normal Gaussian distribution function to the probability density function of the results from the individual 30-min intervals. In 2003, Thomas et al. (2006) extended our research at the *Waldstein-Weidenbrunnen* site by the first investigation of the boundary layer structure over the forest with particular emphasis on coherent structures.

The rationale for both the experimental and modelling design of the EGER studies was based on the analysis of the spatial and temporal scales of the relevant exchange processes in the forest atmosphere, plants and soil (see Sect. 2.4.1 for details). As a first consequence, we limited the overall spatial scale of our experiment to some 100 m × 100 m (“stand/plot scale”). The constraint on the temporal scales to be considered originates from the Damköhler number (Damköhler, 1940), a ratio of two particular time scales, namely the characteristic turbulent transport time over the characteristic chemical reaction time. A Damköhler number (DA) less than unity indicates that turbulent transport is faster than chemical reaction; for $DA \leq 0.1$ reactive trace compounds are considered to be treated as (quasi-)passive scalars.

While characteristic chemical reaction times simply result from ambient concentrations, reaction kinetics, (photo-) chemical constants ($\text{NO-NO}_2\text{-O}_3$), and thermodynamic equilibrium considerations ($\text{NH}_3\text{-HNO}_3\text{-NH}_4\text{NO}_3$), determination of the corresponding spatial scale is more difficult. The characteristic turbulent transport time is a measure of the intensity of turbulent transport. In the surface layer and under near neutral conditions (cf. Wyngaard, 1982), it may be defined by roughness

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length, height above ground, and the ratio of the vertical velocity variance and the friction velocity (Vilà-Guerau de Arellano, 2003). Mayer et al. (2011) present an alternative, stability-dependent definition of the turbulent transport, based on the gradient approach. The questions of both definition and determination of characteristic transport time scales (i.e. available volumes for chemical reactions) within the forest canopy remain open; here, we attempt to tackle the problem through investigation of coherent structures and their effect on coupling of different layers within the canopy with the surface layer above the canopy.

Concentrations of a variety of trace substances can only be measured by slow-response analyzers, like the compounds of the $\text{NH}_3\text{-HNO}_3\text{-NH}_4\text{NO}_3$ triad (Trebs et al., 2004). Consequently, application of all kinds of gradient approaches (e.g. aerodynamic gradient method) is necessary to determine corresponding vertical turbulent fluxes (cf. Thomas et al., 2009). However, over (aerodynamically rough) forest vegetation, the application of these methods is constrained to heights of about twice the canopy height, because below this height the similarity laws may be considerably disturbed due to enhanced surface roughness of the forest. This layer is the so-called roughness sublayer (Garratt, 1978, 1980; Raupach and Legg, 1984; Raupach et al., 1980). In this layer weaker vertical gradients were found, but the turbulent transport is very effective due to coherent structures in the mixed layer (Finnigan, 2000; Raupach et al., 1996). To account for the impact of enhanced roughness, flux-gradient formulations are extended by including a further function, the so-called enhancement factor (Raupach and Legg, 1984; Simpson et al., 1998); explicit functions were given by Garratt (1992) and by Cellier and Brunet (1992). Recently, Harman and Finnigan (2007, 2008) presented a more sophisticated method to consider coherent structures in the mixing layer for the determination of momentum and scalar fluxes, featuring dependence on the mixing layer length scale. In our study, particular emphasis in this direction is given to the application of the aerodynamic method to vertical gradients of the $\text{NH}_3\text{-HNO}_3\text{-NH}_4\text{NO}_3$ triad. As already mentioned above, turbulent fluxes of carbon dioxide, water vapour and the $\text{NO-NO}_2\text{-O}_3$ triad have been measured by the

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eddy covariance technique, accompanied by careful analysis of all radiation and energy fluxes. To investigate horizontal in-canopy structures (advection), special setups of carbon dioxide measurements have been designed.

In order to elucidate formation pathways of nitrous acid (HONO), an important precursor of OH radicals in the lower troposphere, we included in- and above canopy measurements of HONO mixing ratio in our study. There is agreement that in the absence of light the heterogeneous HONO formation (net reaction: $2\text{NO}_2 + \text{H}_2\text{O} \rightarrow \text{HONO} + \text{HNO}_3$) is the most important formation pathway, but the mechanisms have not yet been clarified. The reduction of NO_2 by organic species might also be important under atmospheric conditions (Ammann et al., 2005; Gutzwiller et al., 2002). During daytime the main HONO sink is the photolysis $\text{HONO} + h\nu \rightarrow \text{NO} + \text{OH}$. However, recent measurements showed that gas-phase formation and the parameterized dark formation are too slow to explain measured HONO concentrations during daytime, which are significantly higher than those calculated from the photostationary state (Kleffmann et al., 2005). An additional daytime source was postulated and described by a recent overview (Kleffmann, 2007). The forest canopy provides a large surface for heterogeneous reactions as well as shading of below canopy space, resulting in different source and sink strengths above and below canopy. These differences, mirrored in concentration differences, were analysed with respect to the coupling of the forest and the air layer above the forest by coherent structures.

In-canopy models FLAME (Berger et al., 2004; Inclan et al., 1996) and CACHE (Stockwell and Forkel, 2002; Forkel et al., 2006) have already been tested at the *Waldstein-Weidenbrunnen* site. Here, the modelling concept involves a 3-D model with a nested structure (STANDFLUX, Falge et al., 1997, 2000) and a 1-D canopy-surface layer model (ACASA, Pyles et al., 2000). The application of a higher order closure model like ACASA is necessary to overcome the counter-gradient problem and also to model the probable influence of coherent structures.

We will present an overview of the EGER project, its design, the applied methods, and particularly of first results from both Intensive Observation Periods, namely

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September/October 2007 (IOP-1) and June/July 2008 (IOP-2). Concerning the latter, we focussed our analysis on the coupling between the (above canopy) surface layer, the canopy and the sub-canopy and its consequences for (a) the structure of the turbulent exchange, (b) above canopy fluxes, as well as (c) concentration gradients of non-reactive and reactive trace gases between the surface layer and the forest floor.

2 Material and methods

2.1 Site description

As the research area for this study the *Waldstein-Weidenbrunnen* site ($50^\circ 08' 31''$ N, $11^\circ 52' 01''$, 775 m a.s.l.) in the *Fichtelgebirge Mountains* (Germany) was selected. It is located in the Lehstenbach catchment in NE Bavaria (Germany), a research area of the Bayreuth Center of Ecology and Environmental Research (BayCEER). It is a FLUXNET (Balocchi et al., 2001) site (DE-Bay) with carbon-dioxide flux measurement above the spruce forest since 1996 and intensive ecological and meteorological studies in this area (Matzner, 2004). The site is located NW of the upper EGER river valley (Fig. 1).

The site rates as one of the best of the European FLUXNET sites in terms of data quality. Regarding footprint classification, it is one of the good sites with more than 85 % of the flux originating from the target area "spruce", with the remainder comprising areas of the clearing and some of the quarry (Göckede et al., 2008). It is an extensively investigated location with the good logistical conditions. At *Waldstein-Weidenbrunnen*, CarboEurope quality control tools have been developed. This included the use of *Waldstein-Weidenbrunnen* data sets for the comparison of eddy-covariance software (Mauder et al., 2008) and the test of the footprint classification scheme (Göckede et al., 2004, 2006, 2008; Rebmann et al., 2005). Many experimental efforts of the University of Bayreuth have been realized at our site, particularly a comprehensive micrometeorological experiment in 2003 (Thomas and Foken, 2007a), as well as the BEWA 2001 and 2002 field experiments (Klemm et al., 2006). The measurements for

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rain (in mg m^{-2}), integrated over the entire time period of IOP-1 and IOP-2, was found to be typical for the region. Highest wet deposition rates were observed in the rain fall just after the “Golden Days” of IOP-1 and during the heavy rainfall on 3 July 2008 (IOP-2), respectively. Average O_3 , NO, NO_2 , and SO_2 concentrations, observed at 5 31 m (a.gr.) at the main tower were within the typical range of the long-term data (monitored at the *Waldstein-Pflanzgarten* site (3 m a.gr.). During IOP-2, maximum SO_2 and NO_2 concentrations (10.8 ppb and 15.1 ppb, respectively) occurred on 9 June 2008, when easterly air masses have reached our site from the industrialized regions of the Czech Republic.

10 Within the scope of this overview paper, we have selected for each IOP a so-called “Golden Days” period to present some of our results: 20 to 24 September 2007 (IOP-1) and 28 June to 2 July 2008 (IOP-2), respectively. These periods were characterized by high radiation, no precipitation and hardly any clouds (to emphasize photochemical aspects of our studies). The “Golden Days” period of IOP-2 was warmer and drier than 15 that of IOP-1, with higher air and soil temperatures, higher maximum water vapour pressure deficits and lower soil moistures. Wind speeds were moderate and comparable for both “Golden Days” periods. While the year 2007 was the warmest in the region since permanent weather observations have started in Bayreuth (1850), the summer 2007 was warm but not extreme. IOP-1, starting just at the end of summer 2007, 20 was always under influence of cyclonic conditions and it was only during the “Golden Days” that the anticyclone “Katrin” dominated Central Europe. From 18 June 2008 onward, IOP-2 (in the beginning cold and cyclonic) experienced warmer air, which was bordering colder air masses to the north. Under these warm and cyclonic conditions, thunderstorms passed through the *Fichtelgebirge* region including our site on 25 June 25 (8.5 mm, 15:10–16:50 CET). The IOP-2 “Golden Days” period, starting on 28 June, was characterized by dry summer weather, not dominated by an anticyclone, and ended on 3 July with the cyclone “Renate” (44.1 mm of rain). Up to the end of IOP-2 cyclonic conditions (with some showers) returned.

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2.3 Measurements

2.3.1 General set-up of towers and instrumentation

The so-called “main tower” (31 m, walk-up type) of the *Waldstein-Weidenbrunnen* site (see Fig. 6a, Table 4) served for standard meteorological measurements (e.g. vertical 5 profiles of wind velocity, dry and wet bulb temperature) as well as for the measurements of vertical profiles of trace gas concentrations and trace gas fluxes. An additional, 35 m tall and slim tower for turbulence measurements was set up approx. 60 m south-east of the “main tower” (see Fig. 6b). This action was aimed to increase the data quality of micrometeorological in- and above canopy flux measurements in the footprint area 10 of the *Waldstein-Weidenbrunnen* spruce forest (canopy height 23–25 m). 60 m north-west of the “main tower” another walk-up tower (“bio-tower”) for plant physiological measurements was installed (see Fig. 6c). Along the NE-SW slope (approx. 3°) of the site, where catabatic flows could be expected, five 2 m high masts, equipped with sonic anemometers (not all masts) and intakes for CO_2 concentration measurements, were 15 set up, while another six masts (identical instrumentation) were placed between the “bio-tower” and the “turbulence tower” perpendicular to the slope and along a small trail (for more details see Serafimovich et al., 2008a, b). During IOP-1, turbulent fluxes were measured at “main” and “turbulence” towers only; during IOP-2 a Modified Bowen-ratio system (Liu and Foken, 2001) was installed at the clearing north-east of the “turbulence 20 tower” (50°08′30.32″ N, 11°52′11.0″ E, 780 m a.s.l.), close to the position of the mini-sodar, see Sect. 2.3.2).

2.3.2 Remote sensing measurements of the atmospheric boundary layer

In complex terrain like the *Waldstein-Weidenbrunnen* site, monitoring of the atmospheric boundary layer’s structure became an important tool for interpretation of the 25 turbulent exchange data. For this monitoring a composite of several remote sensing instruments was used: a sodar/RASS and a mini-sodar system at the field site, a

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482 MHz-windprofiler of the German Meteorological Service about 25 km south of the field site, and a 2D sonic anemometer at the top of the “main tower” (Table 5). Data from the two sodar systems and the sonic anemometers at the “turbulence tower” were also used to investigate coherent structures in both the canopy and the atmospheric boundary layer (Thomas et al., 2006).

2.3.3 Measurements of turbulent fluxes

As shown in Table 6, six levels of the “turbulence tower” and up to four levels of the “main tower” have been equipped with 3D sonic anemometers, most of them also with fast-response open-path CO₂ and H₂O analyzers, some of them with fast-response O₃ analyzers, and one level (IOP-2) with the intake for fast-response measurements of NO and NO₂ concentrations. The EUROFLUX methodology (Aubinet et al., 2000) and recent updates of the TK2 software (Mauder and Foken, 2004; Mauder et al., 2008) provided the basis of all calculations for the turbulent fluxes of momentum, sensible heat, latent heat, CO₂, O₃, NO, and NO₂. Furthermore, all flux measurements for the *Waldstein-Weidenbrunnen* site were footprint controlled (Foken and Leclerc, 2004; Göckede et al., 2004, 2006) and data quality checked (Foken and Wichura, 1996; Foken et al., 2004). The coordinate system for all 3-D wind measurements was chosen parallel to the stream lines using the planar-fit method (Wilczak et al., 2001). This was also the basis for determining advection through adaptation of the concept by Aubinet et al. (2003).

During IOP-1, only the turbulent O₃ flux was measured, but during IOP-2 turbulent fluxes of NO, NO₂, and O₃ were measured using eddy-covariance systems. In case of NO and NO₂, high frequency (5 Hz) time series of NO and NO₂ concentrations were recorded with a specialized, fast-response, state-of-the-art, high precision 2-channel NO-NO₂ chemiluminescence analyzer (model CLD 790SR-2, Ecophysics, Switzerland). While one channel was used for the direct measurement of the NO concentration, the measurement of the NO₂ concentration was indirect, following high efficiency conversion of NO₂ to NO (through two up-stream solid state blue light converters

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in series) and consecutive NO detection in the second channel of the chemiluminescence analyzer. To maintain continuous time series of the fast-response NO and NO₂ measurements, NO and NO₂ signals have not been calibrated on-line; instead, they have been related to NO and NO₂ concentrations which were measured side-by-side by slow-response NO-NO₂ chemiluminescence analyzers (see Sect. 2.3.3). During IOP-1 and IOP-2, these analyzers revealed detection limits (3 σ -definition) ranging between 0.05–0.12 ppb (NO) and 0.22–0.76 ppb (NO₂), respectively; data found to be below the limit of detection have been discarded. A sonic anemometer (CSAT3, Campbell Scientific Inc., USA) was used to measure fluctuations of the 3-D wind velocities at 20 Hz. A 53 m long insulated bundle containing the Teflon inlet tubes, tube heating as well as interface cables from the data logger ran from the top of the “main tower” to the analyzer and control computers in a nearby air-conditioned shelter (container). In the case of O₃, high frequency (20 Hz) time series of O₃ concentration were recorded by fast-response solid-phase chemiluminescence analyzers (Güsten and Heinrich, 1996) designed by different manufacturers (s. Table 6). Sonic anemometers were deployed next to the O₃ analyzers (s. Table 6). Combining the signals of 3-D wind velocity fluctuations with those from the fast-response O₃-analyzers provided turbulent O₃ fluxes above (IOP-1, IOP-2) and, within (IOP-2), the *Waldstein-Weidenbrunnen* spruce forest. Solid-phase chemiluminescence analyzers provide O₃ concentration only in relative units (voltage), and the analyzers’ sensitivity is not temporally constant (due to the influence of relative humidity on the chemiluminescent dye). However, absolute O₃ concentrations were simultaneously measured side-by-side each fast-response O₃-analyzer by UV-absorption based ozone analyzers (see Sect. 2.3.3). Based on these data, fast-response O₃ signals could be converted to O₃ concentration and hence, after applying the above mentioned procedures for correction and quality check of eddy-covariance measurements, the turbulent O₃-fluxes and deposition velocities were calculated.

2.3.4 Profile measurements

Measurements of the vertical profiles of wind speed and dry and wet bulb temperatures are part of a standard and continuously running monitoring programme at the *Waldstein-Weidenbrunnen* site (www.bayceer.de). Cup anemometers, characterized by low distance constant (approx. 3 m), and Frankenberger type psychrometer (Frankenberger, 1951), all equipped with high precision sensors, have been mounted at different levels along the “main tower” (s. Table 7). During both IOPs, these sensors (system I, see Table 7) have been amended by additional wind speed sensors (A100ML, Vector Instr., UK) and home-built, thermocouple-based psychrometers (system II, see Table 7).

Along the “main tower”, vertical profiles of CO₂, H₂O, NO, NO₂, and O₃ concentrations were measured at nine (IOP-1) and eleven (IOP-2) levels, respectively (see Table 7). Due to the limited absolute accuracy (but generally much better precision) of the trace gas analyzers, concentrations from different levels have been measured sequentially by one set of analyzers in order to resolve even small vertical differences of concentrations. To keep the time period required to “scan” the full profile short (<30 min), two identical but independently operating sets of analyzers have been deployed. One set was applied to levels of the upper part of the profile (>24 m, IOP-1; >16.5 m, IOP-2), the other set for the lower levels of the profile (i.e. within canopy and particularly close to the soil–trunk space interface). Both sets of analyzers had overlapping levels (24 m, IOP-1; 16.5 m, IOP-2), where concentration measurements (from side-by-side intakes) provided a continuous data set to cross-check (and to correct) for systematic offsets. Ambient air was drawn from the different intake levels through heated and insulated PTFE-tubing (4.1 mm inner diameter; opaque; length: 55 m each) into an air-conditioned container (housing the two sets of analyzers) on the forest floor close to the “main tower”. All sampling lines were continuously purged, and the “active” sampling line was sequentially switched between the different levels (switching interval: 1.5 min). NO and NO₂ concentrations were measured by 1-channel chemiluminescence analyzers (CLD TR780, Ecophysics, Switzerland). NO was detected first,

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then NO₂, after the sampling air has passed a solid-state blue light converter (installed upstream from the analyzer for species specific conversion of NO₂ to NO). Detection limits (3σ) for NO and NO₂ during the experiments were <0.12 and <0.76 ppb, respectively. Precision was <0.6 % (10 ppb NO) and <4 % (20 ppb NO₂), respectively. For measurements of O₃ concentrations, UV absorption spectrometers (TEI 49c/I, Thermo Electron, USA) which had a precision <2 % were used. Concentrations of CO₂ and water vapour were measured by NDIR absorption analyzers (Li7000 and Li840, LI-COR Biosciences, USA); corresponding precision was <0.2 % for both CO₂ and H₂O. All analyzers were calibrated at least weekly, using certified CO₂ standards (pressurized cylinders), a dew point generator (LI-COR Biosciences, USA), and a certified NO standard (5 ± 0.09 ppm, Air Liquide, Germany) in combination with a gas-phase-titration unit (model 146c, Thermo Electron, USA) to generate suitable concentrations of NO, NO₂, and O₃.

During IOP-1, concentrations of the NH₃-HNO₃-NH₄NO₃ triad were measured at two heights above the forest canopy using a novel wet chemical instrument, the GRAEGOR (GRAdient of AErosols and Gases Online Registrator, Thomas et al., 2009). Gaseous NH₃ and HNO₃ and particulate NH₄NO₃ constitute a thermodynamic equilibrium which strongly depends on relative humidity and temperature as well as on aerosol composition, especially SO₄²⁻ concentrations (Mozurkewich, 1993; Nenes et al., 1998; Stelson and Seinfeld, 1982). GRAEGOR is capable of measuring the interacting species gaseous NH₃ and HNO₃ as well as particulate NH₄⁺, NO₃⁻, and SO₄²⁻ selectively and simultaneously at two different heights. For the first time, such highly time resolved measurements of the complete triad were performed above a forest canopy. The instrument was thoroughly characterised for its ability to resolve vertical concentration differences above the forest (and a grassland site) for a range of atmospheric conditions (Wolff et al., 2010a). From the experimentally determined errors of the concentration differences the resulting flux errors were estimated. For the *Waldstein-Weidenbrunnen* forest experiment median instrumental flux errors were: 50 % (NH₃), 38 % (HNO₃), 57 % (NH₄⁺), and 68 % (NO₃⁻), respectively (Wolff et al., 2010a).

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During both IOPs, ambient HONO mixing ratio were measured with a wet-chemical instrument, the Long Path Absorption Photometer (LOPAP, Kleffmann et al., 2002; He-
land et al., 2001). This underlying technique offers a high temporal resolution (5–7 min)
and a very low detection limit of about 0.001 ppb. During IOP-1, we had set up two
5 LOPAP instruments, one close to the forest floor (0.5 m) and the other one just above
the canopy (24.5 m). The data set provides valuable information to study potential
HONO formation processes on the different surfaces, which may be heterogeneous or
even photo-enhanced sources of HONO.

Though corresponding data are not reported here, it should be mentioned (for the
10 sake of completeness of the EGER overview) that the above-mentioned measurements
of turbulent fluxes and vertical profiles were complemented by measurements of the
soil-atmosphere exchange of CO₂, ²²⁰Rn, and ²²²Rn (static chamber), NO, NO₂, and
O₃ (dynamic chambers), measurements of the plant-atmosphere exchange of CO₂,
H₂O, NO, NO₂, and O₃ (dynamic branch/twig cuvettes), as well as by laboratory incu-
15 bation measurements of biogenic NO emission from soil samples taken from plots of
the *Waldstein-Weidenbrunnen* site covered by different understory vegetation (moss,
grass, blueberries, young spruce, see Bargsten et al., 2010).

2.4 Scale and coupling analysis

As stated in the introduction, the experimental and also the modelling design of the
20 EGER studies was based on the analysis of the spatial and temporal scales of the
relevant exchange processes in the forest, atmosphere, plants and soil. The temporal
scales were thereby compared with the characteristic chemical reaction time and the
time scales of coherent structures. Furthermore, the coupling between the atmosphere
and the canopy was taken into account. In the following the basics of these methods
25 of investigation are briefly presented.

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2.4.1 The scale problem

The physical, chemical and biological processes controlling the exchange of energy
and trace substances are characterized by time scales extending from femto-seconds
(e.g. electron transport in chloroplasts), over several hours (e.g. horizontal advection),
5 to several years (climate, Beniston, 1998), and by spatial scales ranging from a few
micrometers (e.g. reactions of heterogeneous chemistry) to the size of landscape ele-
ments (about 10 × 10 km²). Investigations of the EGER project are focussing on widely
overlapping scales (Jarvis, 1995) from contributions of the individual physical, chem-
ical and biological processes, whereas our measuring and modelling techniques are
10 related to specific scales.

The extent of the scale problem is illustrated in Fig. 7, where spatial and tempo-
ral scales of atmospheric, biological, soil and chemical processes, which control the
exchange of energy and trace substances, are schematically shown. Within the en-
tire range of atmospheric processes (light blue squares, ranging from micro scale γ
15 to meso scale α , as defined by Orlanski, 1975), the scales concerning the exchange
processes of energy, water, and trace substances comprise (i) turbulent transport in
canopies (white star), (ii) advection in canopies (white circle), (iii) turbulent transport
above canopies (white diamond), (iv) coherent structures in and above canopies (blue
double arrow), (v) footprint related turbulent fluxes (white square), and (vi) horizontal
20 advection at canopy top (white triangle). In Fig. 7, the typical scales of processes in
plants (Schoonmaker, 1998), those in soil (Vogel and Roth, 2003), in the saturated
and unsaturated zone (Blöschl and Sivapalan, 1995) are shown by the brown framed,
spotted area. While there is a broad overlapping of the spatial and temporal scales
of plant and soil processes, there is only a marginal overlapping of these with the
25 scales of atmospheric processes; with increasing time scales, atmospheric processes
separate more and more from plant and soil processes with regard to space. Not in-
cluded in Fig. 7 are spatio-temporal scales of the measuring techniques from plant
enclosures and soil chambers (a few tenths of square metres) to eddy-covariance data

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with a footprint up to some square kilometres. Remote sensing techniques have even larger areas, depending on the height from where the signal originates. Note that coherent structures have a significantly larger length scale in the vertical than in the horizontal direction, and time scales are also longer. This important issue was recently underlined by Mahrt (2010) regarding the averaging technique for eddy-covariance measurements.

Gas-phase and heterogeneous reactions of relevant reactive trace gases (see reactions 1–15 in Fig. 7) are only attributed to corresponding temporal scales ($1/e$ reaction times). These characteristic chemical time scales (see Dlugi, 1993) simply result from ambient concentrations of the reaction partners, reaction kinetics or thermodynamic equilibrium considerations, and (photo-) chemical constants or variables. However, most of the corresponding (photo-)chemical constants have been derived from laboratory experiments in well-mixed reaction chambers (see Atkinson et al., 2004). Therefore, corresponding spatial scales (most likely $\ll 1$ mm) seem not to be relevant for real atmospheric conditions, particularly not within tall canopies, where (strong) decoupling by vertical thermodynamic stratification is (at least during night) more likely than full coupling with the (above canopy) surface layer, or even complete turbulent mixing throughout trunk and canopy spaces. The latter, however, is the prerequisite of corresponding volume averaging of chemical processes, which is a volume averaging of soil and plant processes – the basis of coupling of turbulent, soil, plant, and chemical processes.

2.4.2 The Damköhler number

As already mentioned in Sect. 1, the Damköhler number (DA) is the ratio of two particular time scales, namely the characteristic turbulent transport time (τ_{turb}) over the characteristic chemical time scale (τ_{chem}):

$$DA = \frac{\tau_{\text{turb}}}{\tau_{\text{chem}}} \quad (1)$$

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Dominance of the turbulent transport over chemical reaction (hence, treating a trace substance as “non-reactive” during transport through the forest or considering a trace substance as “non-reactive” for “volume averaging”) is given, if $DA \leq 0.1$. Defined by the K-approach, the characteristic turbulent time scale, τ_{turb} , may be computed from the mean transfer velocity v_t and the thickness Δz of the layer being considered (Mayer et al., 2011)

$$\tau_{\text{turb}} = \frac{\Delta z}{v_t} \quad (2)$$

The final expression for the time scales of turbulent trace gas transport thus reads:

$$\tau_{\text{turb}} = \frac{\Delta z^2 \varphi_H(\zeta) \cdot Sc_t}{\kappa \cdot z_m \cdot u_*} \quad (3)$$

with φ_H the universal function for the heat exchange (also used for trace gases), $\zeta = z/L$ with the Obukhov-length L , the von-Kármán-constant κ , the measuring height z_m , the turbulent Schmidt number Sc_t , and the friction velocity u_* . It should be stated that Eq. (3) is only applicable under Monin-Obukhov similarity conditions.

Typical time scales for reactions of chemically reactive compounds are given in Fig. 7 (Reactions 1–15, see Dlugi, 1993). However, assuming reasonable vertical mixing and considering given (measured) concentrations of reaction partners, the characteristic chemical reaction times can be calculated using (i) kinetic constants (Atkinson et al., 2004), (ii) radiation fluxes for photolysis (e.g. for O_3 , NO_2 , and HONO), and (iii) known thermodynamic quantities, NH_3 - HNO_3 - NH_4NO_3 , (see Wolff et al., 2010b), respectively. As in Mayer et al. (2011), in the following we consider for the chemical timescale (τ_{chem}) only the NO - NO_2 - O_3 triad including the (photo-) chemical reactions $O_3 + NO \rightarrow NO_2 + O_2$, characterized by the reaction coefficient $k_1 = 1.40 \times 10^{-12} \exp^{-1310/T}$ (see Atkinson et al., 2004) and $NO_2 + h\nu \rightarrow NO + O_3$ ($\lambda < 420$ nm, characterized by the NO_2 photolysis frequency $j(NO_2)$, which has been measured directly). The chemical destruction of NO_2 is then controlled by $j(NO_2)$ only and, in turn, the chemical production of NO

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was found that momentum and sensible heat transport by coherent structures is dominant in the canopy and carbon dioxide and latent heat transport by coherent structures increases with height within the canopy and reaches a maximum at the upper canopy level. The flux contribution of the ejection and sweep phase of coherent exchange were also determined. The flux contribution of the ejection phase decreases with increasing height within the canopy and becomes dominant above the canopy level. The flux fraction transported during the downward directed sweep phase increases with height within the canopy and becomes the dominating exchange process at the upper canopy level. Close to the ground surface in the subcanopy space, ejection and sweep phase contribute equally to the flux transport. This issue is discussed in more detail by Serafimovich et al. (2011). Furthermore Fig. 9 shows the coupling classification and, for a better understanding, also selected meteorological parameters and fluxes

Conditional sampling analysis shows a domination of coherent structure signatures in vertical wind measurements (Fig. 10) with probable temporal scales in the range 10 s to 40 s. The number of coherent structures detected at the turbulence tower is lowest directly under the crown (5 m) and higher in the trunk space and above the canopy with a maximum at the canopy level. Most of the structures were found in the vertical wind velocity followed by the temperature. There were more structures found in water vapour flux than in carbon dioxide flux.

The analysis was extended to the lower boundary layer according to Thomas et al. (2006) using sodar measurements. To derive characteristics of coherent structures from sodar/RASS individual soundings, data preparation and the wavelet analysis were used. Due to the effect on the measurements of occasional environmental noise detection and discarding of erroneous data, gap filling and de-noising were applied (Crescenti, 1998; Miller and Rochwarger, 1970; Neff and Coulter, 1986). In a first step, the data were filtered using the error flag output by the sodar/RASS system. As a next step, a quality control described in Thomas et al. (2006) was applied. An example of the spectra of wavelet variance for different observation levels is presented in Fig. 11. The data for the lower observation heights exhibit the first maximum at ~30 sec, whereas

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the observation levels above show the stronger maximum at ~50–60 sec.

The classification of the coupling was used in the following sections for a better understanding of the chemical processes. The first analysis showed that the attribution of coupling states into the classes C and Cs as well as W and Dc is difficult. Therefore the coupling classes C/Cs and W/Dc were combined to show a more significant picture of the results. First the statistics of the coupling classes is shown in Fig. 12 for both IOPs.

The typical difference between IOP 1 and 2 is the domination of class W/Dc in the autumn IOP-1 whereas in the summer IOP 2 in the same hours the case Ds is dominating. But the cases of well coupled situations are not significantly different for both IOPs. The typical asymmetry in the stability with stronger unstable situations in the morning, already stable situations in the afternoon and the strongest stable situation in the first half of the night can easily be seen in the distribution of the coupling classes as well (see Fig. 9). Obviously, IOP 2 shows an oasis effect (necessary energy of evaporation is on the cost of sensible heat flux, see Stull, 1988) with a stronger decoupling already in the hours 15–18 which is usually expected not to occur before the hours 18–21 (e.g. IOP-1). Situations with a good coupling were found in the nights of both IOPs, probably connected with low-level jets and gravity waves (see Sect. 3.3).

The analysis of coherent structure was also used to apply a relaxed eddy accumulation conditional sampling to determine the daytime soil respiration from turbulence measurements above the canopy. The method based on quadrant analysis (Shaw et al., 1983) with regard to CO₂ and water vapour contents can be used for identifying the origin of CO₂ fluxes (Scanlon and Albertson, 2001): this method was improved for a practical application and applied for different canopies by Thomas et al. (2008). However, for the dense *Waldstein-Weidenbrunnen* spruce forest corresponding results were not very satisfactory. Therefore the method was again tested during IOP-1 with some site specific modification for the conditional sampling characteristics. For the *Waldstein-Weidenbrunnen* satisfactory results were achieved leading to an integrated five-day flux.

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In Serafimovich et al. (2011) the vertical coupling according to Fig. 9 was investigated as well as the horizontal coupling between the small towers in the trunk space used for advection investigation. This coupling concept was applied to advection investigations (Siebicke, 2011). The coupling was found to be much stronger along this trail (no understory) and not along the slope (understory up to 1 m height).

Although advection is very important for the carbon balance of a forest (Finnigan, 2008), the problem posed by measurement of the advection has still not been solved. The classical approach with a more or less steady-state flow through a volume element was, in many experiments, unsuccessful (Aubinet et al., 2010; Aubinet, 2008; Feigenwinter et al., 2008). This was also tested on the *Waldstein-Weidenbrunnen* site in 2003 and during both IOPs. Even very careful analysis of the data including a planar fit analysis (Wilczak et al., 2001) adapted to the process could not make the necessary progress. Therefore Siebicke (2011) tried to explain the measured horizontal carbon dioxide gradients along both tower lines with the complicated structure of the canopy and understory. He found significant correlations between the horizontal gradients, which were measured during IOP-2 with a high time resolution, and the duration and intensity of coherent structures.

3.3 Influences of the atmospheric boundary layer

To understand influences of the dynamics of the atmospheric boundary layer on surface exchange processes, information about the spatio-temporal behaviour of the boundary layer's temperature and wind fields must be known. The two installed sodar systems at the experimental site (see Sect. 2.3.2) showed a good agreement at those heights, which are covered by both systems. Also, upper levels of sodar and windprofiler measurements showed only very small differences. Therefore, the results of the three systems were combined to make available a composite picture of the boundary layer structure.

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The specific aim of the investigation was the flow characteristics in the roughness sublayer and potential influences of the clear-cut south of the *Waldstein-Weidenbrunnen* site on trace gas fluxes. For stronger winds, the wind direction of the entire boundary layer dominates down to the canopy level. In contrast, Fig. 13 shows the influences of the clear cut on the wind field at the site. These situations (cloudy conditions) are related to very small horizontal wind velocities above the canopy and significantly lower vertical wind velocities (not shown). In general the main wind directions – in the south-west sector – are nearly identical at 200 m (sodar) and 1050 m (windprofiler) heights. A strong SE component, coming up the Lehestenbach valley, could occur as well, but only below 100 m at night.

Furthermore, the low-level jets (LLJ) found in 2003 were investigated more carefully and found to be similar to those in other studies (Banta et al., 2002). They were selected when the wind velocity above and below the jet was at least 2 m s^{-1} lower (Stull, 1988) and the duration of the jet was longer than 90 min. An example of this is shown in Fig. 14. The wind velocity of the jet was found to be above 10 m s^{-1} for about 6 h at a height of 300 m, decreasing up to 200 m. The wind direction of the jet was SE while above the jet the wind direction moved from SE over S to W. In the jet the vertical wind was negative. During IOP-1 LLJs were found on nine nights, mainly at about 150 m height with a typical wind velocity of 9 m s^{-1} . During the summer IOP-2 LLJs were found on 11 nights but typically at 250 m height with about 10 m s^{-1} .

Low level jets have a significant influence on the night-time exchange of a forest. Karipot et al. (2008, 2005) found that low-level jets enhance turbulence and mixing close to the ground due to increased shear. CO_2 fluxes are very small during calm periods following early evening hours and large fluxes are observed corresponding to the LLJ periods. The flux increases up to $9\text{--}15 \mu\text{mol m}^{-2} \text{ s}^{-1}$, though the u_* values increased only just above 0.2 m s^{-1} . This is an indication that during intermittent turbulent periods associated with features such as sporadic jets, high fluxes are possible from the large fluctuations in the accumulated CO_2 , in phase with moderate vertical velocity fluctuations. CO_2 -rich ejections from the canopy contribute more to the positive

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flux during weak-LLJ events. For the LLJ shown in Fig. 14 similar conditions (Karipot et al., 2005, 2008) were found with an increase of the friction velocity and of the carbon dioxide flux from $4 \mu\text{mol m}^{-2} \text{s}^{-1}$ to $8 \mu\text{mol m}^{-2} \text{s}^{-1}$.

The vertical distribution of the concentrations of NO, NO₂, O₃ and CO₂ is shown in Fig. 15 for the situation with the Low-Level-Jet (Fig. 14). Due to the increased shear a better mixing was found during the period with the LLJ between 02:00 and 08:00, with a maximum at 04:00. The better mixing resulted – for all trace gases – in a reduction of the accumulated concentrations near the surface (Fig. 15). In the early morning, at approximately 04:00, when the LLJ occupied lower heights, the atmosphere close to the surface is suddenly mixed with high NO₂ concentrations, probably an outflow of the upper Eger river valley during easterly winds. However, this picture contains some other features. The most impressive is the inflow of fresh air with low NO₂ and high O₃ between 22:00 and 24:00, connected with gravity wave influences. For high NO (and NO₂) concentrations during the morning hours (06:00–12:00), see next Sect. 3.4.

3.4 Local advection of reactive trace gases

Concerning the interpretation of fluxes and vertical concentration gradients of the reactive NO-NO₂-O₃ triad, the *Waldstein-Weidenbrunnen* site shares a rather general problem with any other site located in polluted, but also in rural areas: the proximity to anthropogenic sources of NO_x (industrial, domestic and/or traffic). Within the dispersion plume close to these sources, freshly emitted NO of high concentration (several hundreds of ppb) titrates ambient O₃ and starts the formation of NO₂. However, it depends upon the day/night situation, on the actual boundary layer stability regime, and above all on the distance of the site to the source, to what extent flux and concentration gradient measurements will be biased by advection of high NO (or NO₂) concentrations. In the case of the *Waldstein-Weidenbrunnen* site, the unavoidable NO source consists of the nearby district road (HO18), characterized by a rather high traffic volume of 2100 cars per day (working days). The nearest distance of HO18 to the site is about

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1.2 km. However, since main wind sectors of the site are SW to NW, and HO18 runs west of the site from SSE to NNW, “advective disturbance” of desired homogeneous concentration fields is more than likely. Exemplarily for the entire periods of IOP-1 and IOP-2, NO concentrations, measured with 5 Hz resolution (s. Sect. 2.3.2) above the canopy (30 m a.gr.), have been averaged to 30 min means and are shown for the “Golden Days” period of IOP-2 in Fig. 16. NO concentrations are observed every day to increase strongly at 06:00, reaching maximum values around 09:00, and to decline to <0.1 ppb after noontime. The early morning increase can definitely be attributed to the emission and subsequent accumulation of NO in the still existing shallow nocturnal boundary layer over the site with the begin of the commuting traffic (to district capitals Hof and Bayreuth) which starts at 05:30–06:00. Until 09:00, high NO concentrations (30 min averages) are associated with large standard deviations, indicating high variability and consequently low stationarity of NO concentrations. Although not shown here, mean NO₂ and O₃ concentrations also reveal large standard deviations. Noticeable growth of the daytime convective boundary layer (after 10:00) leads to substantial dilution of traffic emissions and consequently to much lower NO concentrations at the site. Following our sodar/RASS measurements, the convective boundary layer at the site is still present until 18:00–19:00; most likely for that reason, enhanced NO concentrations due to the return commuting traffic (16:00–19:00) could not be observed.

When enhanced NO concentrations from the district road arrive at any position of the forest edge, the actual surface layer flow will transport the NO from there to the measurement site. This will lead, particularly for NO, to the generation of chemically induced vertical concentration gradients. The wind speed at the forest floor is approximately one order of magnitude lower than above the canopy. This, in turn, allows a tenfold longer time for O₃ to react with NO at the forest floor. The reaction product NO₂, however, will be photolyzed back to NO above the canopy, while in the shadow at the forest floor this reaction is at least strongly suppressed. Both effects lead at the measurement site, finally, to noticeable differences of NO concentrations between above canopy and forest floor. A clear example is shown in the upper panel of Fig. 15, where

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a vertical NO gradient (between 31.5 m and 0.9 m a.gr.) of ≥ 0.3 ppb was observed during 08:00–10:00 (1 July 2008).

Since in the presence of strong horizontal advection and low stationarity of concentrations the interpretation of measured fluxes and/or vertical gradients in terms of turbulent exchange is generally not meaningful, we omitted all NO, NO₂, and O₃ data measured between 06:00 and 12:00 CET from our analysis in Sects. 3.5.2, 3.5.3, and 3.6.2.

3.5 Trace gas fluxes

3.5.1 Time series

Trace gas fluxes which have been measured by eddy covariance technique are shown in Fig. 17 (IOP-1: left panels, a–c; IOP-2: right panels, a–e). They comprise fluxes of water vapor ($F_{\text{H}_2\text{O}}$), carbon dioxide (F_{CO_2}), ozone (F_{O_3}), nitrogen dioxide (F_{NO_2}), nitric oxide (F_{NO}), and the composites $F_{\text{O}_x} = F_{\text{O}_3} + F_{\text{NO}_2}$ and $F_{\text{NO}_x} = F_{\text{NO}} + F_{\text{NO}_2}$. During the selected fair-weather days, $F_{\text{H}_2\text{O}}$ clearly followed diel variations of available energy and H₂O gradients; it was close to zero during night, and up to 200 W m^{-2} flux during day (Fig. 17a). Classic daily variations were found for F_{CO_2} , featuring negative fluxes during daytime (CO₂ uptake by vegetation) and positive (respiration) fluxes during night (Fig. 17 b). As expected, F_{O_3} reflected during both IOPs the well-known O₃ deposition to forest vegetation and ground (Fig. 16c, black symbols), and F_{O_x} followed closely that of F_{O_3} (Fig. 17 c, grey symbols) for the reason explained below (see Sect. 3.5.2). In contrast, F_{NO_2} (measured during IOP-2 only) was mostly directed upwards, with large fluxes during most of the daytime and smaller fluxes during night (Fig. 17d, right panel). For NO, however, peaks of downward directed fluxes were determined during the morning hours, whereas F_{NO} was indistinguishable from zero for the rest of the day and also during night-time (Fig. 17e, right panel).

Data gaps in F_{NO} and F_{NO_2} were due to routine calibration. The flux of the chemically conservative composite NO_x showed a very similar pattern to that of F_{NO_2} (Fig. 17d,

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right panel, grey symbols), with the exception of those time periods between 06:00 and 12:00 CET, when F_{NO_x} was substantially lower than F_{NO_2} (due to noticeable $F_{\text{NO}} < 0$), which in turn is most likely the result of advection induced, positive, strong vertical gradients of NO concentration above the canopy. Otherwise, $F_{\text{NO}_2} \approx F_{\text{NO}_x}$, because the contribution of F_{NO} to F_{NO_x} is one order of magnitude lower than that of F_{NO_2} . Therefore, both F_{NO_x} and F_{O_x} (the flux of the chemically conservative composite $\text{O}_x = \text{O}_3 + \text{NO}_2$), and also the respective concentration differences, will not further be considered for the analysis of fluxes and concentration gradients vs. coupling stages (see Sects. 3.4.2 and 3.5.2).

During IOP-2, in the afternoon (12:00–18:00), turbulent NO₂ fluxes measured by eddy-covariance technique at 31.5 m(a.gr.) ranged between + 0.5 and + $\sim 1.8 \text{ nmol m}^{-2} \text{ s}^{-1}$, i.e. the *Waldstein-Weidenbrunn* spruce forest appeared as a significant NO₂ source during this time period. Without loss of generality, we may consider this flux just as a net NO₂ flux, namely as the result of a downward directed deposition flux (NO₂ uptake by spruce needles) and an upward directed flux (due to complete oxidation ($\text{NO} + \text{O}_3 \rightarrow \text{NO}_2 + \text{O}$) of the NO, which is biogenically emitted from forest soil). Ignoring potential (small) emission of NO₂ from spruce needles (Lerdau et al., 2000) and estimating the NO₂ deposition flux to approx. $-0.2 \text{ nmol m}^{-2} \text{ s}^{-1}$ (cf. Breuninger et al., 2011), the corresponding biogenic NO flux from the forest soil should then range between +0.3 and + $1.6 \text{ nmol m}^{-2} \text{ s}^{-1}$. From the results of Bargsten et al. (2010), where soil samples of the *Waldstein-Weidenbrunn* site were quantified for their net potential NO soil emission, maximum soil NO emissions of + $1.7 \text{ nmol m}^{-2} \text{ s}^{-1}$ were estimated for the conditions of IOP-2.

The fluxes of $F_{\text{NH}_4+\text{tot}}$ and $F_{\text{NO}_3-\text{tot}}$ are shown in Fig. 17 (left panels, d and e) and were mostly directed downward to the forest canopy with pronounced diel variations. Wolff et al. (2010b) provide a detailed discussion about the magnitude of $F_{\text{NH}_4+\text{tot}}$ and $F_{\text{NO}_3-\text{tot}}$ with respect to comparable previous studies reported so far. Turbulent time scales (τ_{turb} , Eq. (2)) are also shown in Fig. 17 (panels f); they remained mostly below 50 s during both IOPs.

IOP-1 (Fig. 18d and e, left panels) showed small negative values, again with medians during Wa/Dc closer to zero than those during Ds and Cs/C regimes.

3.5.3 The Damköhler number

Turbulent and chemical time scales, as well as Damköhler numbers, have been determined for the above – canopy layer only and all data obtained between 06:00–12:00 have not been considered (s. Sect. 3.4). During daytime, turbulent time scales (τ_{turb}) showed a distinctive pattern of higher medians for the decoupled conditions (Wa/Dc), when comparing different coupling stages (Fig. 19a, b). During night time a similar pattern of higher medians during Wa/Dc conditions can be found (limited by availability of data in certain classes). During IOP-1 and IOP-2, τ_{turb} was longer at night than at day for partly and fully coupled conditions. Chemical time scales (τ_{chem}), however, were fairly constant when comparing different coupling stages separately for day or night-time (Fig. 19 c, d), yet slightly longer at night. When comparing the fall period (IOP-1, Fig. 19c) with the summer period (IOP-2, Fig. 19d), interquartile ranges of τ_{chem} were wider during fall, and τ_{chem} was also longer than during the summer period. Given the relatively constant behavior of τ_{chem} , corresponding Damköhler numbers ($DA = \tau_{\text{turb}}/\tau_{\text{chem}}$, Fig. 19e, f) reflect the daytime pattern already described for τ_{turb} , with higher medians for decoupled conditions (Wa/Dc). During night-time, medians are still highest for Wa/Dc stages, but dispersion is widest for the Cs/C stages of the summer period. Only during partly or fully coupled summer daytime conditions does part of the interquartile range of Damköhler numbers fall below $DA < 0.1$ (i.e. where reactive trace gases may also be treated as “passive tracers”). Damköhler numbers (Eq. 1), given in Fig. 17g, reflected this pattern, because chemical time scales were relatively constant during the time of our observations (data not shown).

Not surprising is that the turbulent time scales and Damköhler numbers for the coupling stages Ds and Cs/C are nearly identical. Both are determined above the canopy under coupled situation but in the case of Ds the forest floor is, of course, decoupled.

Further research is necessary to investigate whether the use of the turbulent time

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scale derived by the gradient approach (Mayer et al., 2011) is the appropriate method in cases when coherent structures dominate the flux (see Sect. 3.2). Therefore, coherent structures should also be detected below 10 s, which is the lowest indicated duration for coherent structures with software used in this paper (Thomas and Foken, 2005). Consequently their duration time and the vertical scale of the structure should be used as the turbulent time scale for the Damköhler number. With such an approach Damköhler numbers may even be significantly below 0.1.

3.6 Above-below canopy concentration differences

We have not pursued determination of in-canopy fluxes according to the classical gradient approach simply for the fact that, particularly in the case of the strong influence of coherent structures, so-called “counter gradients” of meteorological quantities and concentrations may exist throughout the canopy. Therefore, we confined ourselves to typical concentration differences between the trunk space and the top of the canopy. For that, we selected the measurement heights of 24 m (immediately above the canopy) and 1 m (above the forest floor). The latter is, for reactive trace gases, no longer influenced by the strong concentration gradients within the first tens of centimeters above the forest floor (see Fig. 15). For the investigation of vertical gradients (Sect. 3.6.2) all data obtained between 06:00–12:00 have not been considered (see Sect. 3.4).

3.6.1 Time series

During both IOPs, H_2O concentration differences ($\Delta[\text{H}_2\text{O}] = [\text{H}_2\text{O}]_{24\text{m}} - [\text{H}_2\text{O}]_{0.9\text{m}}$) between the 24 m (canopy top) and 0.9 m (above understory) levels were for most of the daytime negative, and positive during summertime nights, confirming that we have observed well-known fair weather in-canopy gradients with moister conditions below the canopy compared to those above during day, and opposite gradients during night (Fig. 20a). Most of the time, $\Delta[\text{CO}_2] = [\text{CO}_2]_{24\text{m}} - [\text{CO}_2]_{0.9\text{m}}$ was negative, however,

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positive Δm concur with positive F .

However there are two exceptions: negative daytime F_{NO} coincide with negative $\Delta[\text{NO}]$, and negative daytime F_{CO_2} coincide with negative $\Delta[\text{CO}_2]$, with the exception of Wa/Ds stages in IOP-2 (7 values only). What distinguishes daytime CO_2 and NO from the other scalars and from night-time conditions? The answer lies within the “control volume” (i.e. between $z = 0$ m and $z = 32$ m) for a flux measured at $z = 32$ m. Whereas for the sign-conforming cases (night-time conditions and daytime H_2O , O_3 , and NO_2) the control volume contains either sources or sinks, for the non-conform scalars (daytime CO_2 and NO) the control volume contains both sources as well as sinks for the respective scalar.

For daytime CO_2 , the control volume contains CO_2 sources from soil, trunk and leaf respiration and the even stronger CO_2 sink, i.e. the assimilation by green leaves in understory and canopy. Hence, the mixture of biological sources and sinks within the control volume distorts the conformity between flux and concentration differences. This is completely different during the night, where the CO_2 sink term “assimilation” is missing, only source terms remain, and signs of flux and $\Delta[\text{CO}_2]$ are consistent again.

Now, for daytime NO, which is biogenically formed in the soil and emitted from the soil surface to the forest atmosphere, there are no known biological sinks in the control volume. However, most of the emitted NO is oxidized by ozone to NO_2 in the first few tenths of a meter above the forest floor, and a part obviously also still above the first meter. The typical “back reaction”, the photolysis of NO_2 to O_3 and NO, is suppressed, since it is comparatively dark below/within the canopy or in the understory. Most of the NO_2 (the former NO) can leave the control volume at $z = 32$ m, because the uptake by the needles is much lower than for O_3 (Breuninger et al., 2011; Geßler et al., 2002). Hence, in the case of NO, the mixture of mainly biological sources and mainly chemical sinks within the control volume is responsible for the non-conformity between flux and concentration differences.

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In summary, the attempted – rather qualitative – synopsis of Figs. 17 and 21 made evident a common problem of non-reactive as well as reactive trace gases, namely that a mixture of source and sink terms in the control volume of a flux requires careful assessment of primary sinks and sources, turbulence, and photochemical interactions.

Corresponding questions are the subject of ongoing analyses.

Regarding the HONO mixing ratio, differences for Ds and Cs/C regimes are very similar at daytime (slightly negative or close to zero), which can be attributed to the photolytic sink above the forest and heterogeneous formation at the forest floor in the late afternoon. At night time no clear result is possible due to the very limited number of measurements. Sörgel et al. (2011) concluded that differences in source and sink processes in and above the forest canopy became obvious only for the decoupled situations (Wa, Dc, Ds). Otherwise mixing ratio differences were close to zero. The positive values during night could partly be attributed to advection of HONO rich air above the canopy which only partly penetrated the canopy.

A more quantitative analysis of fluxes (or concentration differences) with respect to the coupling stages is, on the one hand, limited by the small number of data points within some of the coupling stages (daytime Wa/Dc during IOP-1 and IOP-2, and night-time Ds during IOP-1). On the other hand, and even more importantly, coupling stages are subject to a typical diurnal and annual variation (see Fig. 12), and therefore superimposed with meteorological differences between the coupling stage categories (see Sect. 3.5.2). Differences in the magnitude of fluxes between the coupling stage classes are therefore mainly due to differences in available energy, atmospheric demand or available soil water (in the case of $F_{\text{H}_2\text{O}}$), and only partially influenced by the coupling stage itself. Similarly, the magnitudes of F_{CO_2} (or F_{O_3}) may be modulated by the associated coupling stages, but certainly are driven by radiation, temperature and humidity, which affect stomatal opening (or chemical conversions) and consequently deposition velocities.

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3.7 Modelling of fluxes

For EGER IOP-1 campaign, latent heat exchange was analyzed with the three dimensional microclimate and gas exchange model STANDFLUX (Falge et al., 1997, 2000), and the one dimensional model ACASA (Pyles, 2000; Pyles et al., 2000). Both models were parameterized for the *Waldstein-Weidenbrunnen* site. Parameters common to both models included information on vertical leaf area distribution, and specific sets of physiological parameters for top, middle, and bottom canopy leaf gas exchange. STANDFLUX employed, in addition, horizontal leaf area distribution, tree positions and tree sizes. Further information on model theory and setup is found in Staudt et al. (2011). Latent heat exchange is modelled as the energy equivalent of the sum of evaporation from interception pools and upmost soil and litter layer, and tree and understory transpiration.

Prerequisite of comparisons between measured and modeled energy fluxes is a closed energy balance (Falge et al., 2005), because physical models are based on energy balance closure. In former studies, a value of 77 % (Aubinet et al., 2000; Foken, 2008a) was found according to

$$100\% - Res = \frac{H + LE + G}{R_n} \cdot 100\% \quad (11)$$

with the net radiation R_n , the sensible heat flux H , the latent heat flux LE , the ground heat flux G , and residual Res . In this study the heat storage was neglected. During both IOPs similar values were found with approx. 80 % for both IOP's (Fig. 22).

Coupling state analysis of measured data and model predictions, based on Thomas and Foken (2007a), complement traditional analyses of data-model comparisons. Three coupling stages are distinguished: wave motion and decoupled canopy (code Wa/Dc, $n = 8$ during daytime, $n = 57$ during night-time), decoupled subcanopy (code Ds, $n = 28$ during daytime, $n = 1$ during night-time), coupled subcanopy by sweeps and entirely coupled (code Cs/C, $n = 68$ during daytime, $n = 18$ during night-time). The use of coupling states demonstrates which turbulence stages are well represented and

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which are poorly represented by the model, and thus can serve as a diagnostic tool to improve turbulence parameterizations. By visualizing the data with respect to coupling stages, data-model mismatch at the different stages can be identified (see Fig. 23).

Figure 23 shows that for the daytime decoupled cases (Wa/Dc), both models largely overestimated eddy-covariance measurements, probably resulting from the difficulties in performing eddy-covariance measurements under decoupled conditions. During daytime, STANDFLUX underestimates the measured evapotranspiration, especially for the Cs/C stage (Fig. 23a), when comparing the model results to the eddy covariance data from the turbulence tower at 36 m (Staudt et al., 2011). The performance of STANDFLUX improves when compared against eddy covariance data from the main tower at 32 m (data not shown), because the eddy covariance fluxes measured at the main tower are lower than those measured at the turbulence tower ($LE_{32m} = 0.92 LE_{36m} - 8.8$, $r^2 = 0.88$, for 20–24 September 2007, and quality flags 1–6), potentially because the two towers sample different footprints. ACASA gave slightly better results compared to the measured evapotranspiration at Ds, and Cs/C stage (Fig. 23b). However, ACASA underestimates canopy transpiration (Staudt et al., 2011); an overestimation of evapotranspiration implies therefore an overestimation of the evaporation sources (soil and interception pool) by the model. For night-time conditions, both models underestimated eddy-covariance measurements during all coupling stages. Reasons for this underestimation might either originate in the models or the eddy-covariance measurements which might be underestimated themselves due to the lack of energy balance closure, exacerbating the underestimation of the models even more. Then again, the eddy-covariance estimates might be overestimating the uniform forest below the tower due to the contribution of clearings to the flux footprint. These clearings were found to act as a source of moisture and thus increase eddy-covariance estimates. The reasons for the modeled underestimation of eddy-covariance transpiration measurements are different for the two models ACASA and STANDFLUX. Ecosystem evapotranspiration modeled with the ACASA model agreed better with eddy-covariance measurements (Fig. 23b), but partly for the wrong reason: an overestimation of soil and

understory evapotranspiration with a constant offset of about 10 W m^{-2} , compensated the modeled canopy transpiration which was too low. In contrast, the STANDFLUX model underestimated both, canopy transpiration and soil and understory evapotranspiration. However, it should be noted that, even though relative errors of the models were large, absolute errors were only 8 W m^{-2} for ACASA and 10 W m^{-2} for STANDFLUX during decoupled conditions, and 15 W m^{-2} for both models for coupled conditions.

High variances are expected because the dynamic footprint always adds to the flux variance at halfhourly time scales, that is, the eddy covariance method does not sample the same patch all the time, whereas the model setup is static. In addition, the relative errors might show larger variances, because the models do not account for sweeps and ejections or for the conditions of the W stage, where neither the assumptions for the model nor for eddy covariance theory are fulfilled. In the stable canopy space stimulation of gravity waves occurs at times (Cava et al., 2004; Lee et al., 1997). Under certain conditions, such waves become nonlinear and lead to ejections from the canopy (Fitzjarrald and Moore, 1990). Because of difficulties in resolving the Reynolds mean, effects of such singular events may be improperly calculated in a halfhourly eddy flux record.

4 Conclusions and outlook

It was the aim of this study (i) to introduce the research project EGER (Exchange processes in mountainous Regions), (ii) to present an overview of its conceptual and instrumental design, (iii) to describe the applied methods and multi-scale methodologies, (iv) to investigate site-specific meteorological and micrometeorological conditions (i.e. forest roughness layer, coupling between surface layer and canopy, influence of boundary layer features), and (v) to present first results of above canopy exchange fluxes and concentration differences (above canopy forest floor) of non-reactive and reactive trace gases. The main results and findings are concluded as follows:

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- For the investigation of the in- and above-canopy exchange of energy, non-reactive and reactive trace gases, the consistent application of the scale concept (Fig. 7) is necessary, since the underlying soil, biological, chemical and (turbulent) transport processes form a complex multi-scale problem.
- While the individual processes widely overlap in the temporal domain, there is a larger (chemical) or smaller (soil, biological) separation from the turbulent processes regarding the spatial domain. Since for the exchange over tall vegetation (forests) the concept of a flux through an interface has to be replaced by a flux through a volume element, not only 3-D advection became a substantial problem. Volume averaging, necessary to bridge the spatial scale gap between chemical and turbulent processes, is immediately confronted with the reactivity of trace gases (e.g. is the (turbulent) transport through the canopy fast enough to allow soil/plant emitted reactive traces gases to reach the surface layer unchanged?). For the surface layer above the canopy, the ratio of the turbulent time scale over the characteristic chemical time scale, the Damköhler number, seems to be the suitable measure to distinguish trace gases between “passive tracers” and those compounds which will change their concentration significantly during turbulent transport.
- According to Fig. 7, coherent structures do not always follow the typical scaling system and this might increase turbulent mixing (Mahrt, 2010), which would consequently reduce the influence of fast chemical reactions to measured fluxes and concentration differences. First analysis of EGER data sets has already shown that the coupling concept (Thomas and Foken, 2007a) is without doubt a new and substantial tool for the interpretation of above-canopy fluxes and concentration differences (above-canopy versus forest floor). But it seems meaningful that the more point-related approach of the turbulent time scale (necessary for the calculation of the Damköhler number above the canopy) should be replaced by the spatio-temporal scale of coherent structures, which is based on the coupling

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concept. At least, this is definitely necessary when the contribution of coherent structures to vertical transport processes dominates (night time, see Figs. 18, 19 and 21).

- It was important to complete the experimental EGER studies by boundary-layer profiling measurements, because boundary-layer phenomena also influence transport processes at the site scale. One of the most investigated phenomena in this respect is that of low-level jets, which may re-distribute vertical concentration profiles (s. Fig. 15) and increase surface fluxes due to strong vertical shear. Similar effects are caused by gravity waves. However, both phenomena are very much related to site-specific conditions.
- The analysis of the coherent structures and the coupling can be applied to an even wider range than shown above. The method of the analysis of coherent structures even offers the selection of sweeps and ejections in and above the canopy. Serafimovich et al. (2011) have shown that coupling is not only a vertical exchange effect but also a horizontal. This horizontal coupling depends substantially on the canopy structure, which may be represented by subcanopy density and the PAI. Therefore a careful analysis of both quantities is essential. This was applied by Siebicke (2011) to give a more general explanation of the night-time in-canopy CO₂-advection problem.
- By comparison the sign of fluxes and gradients, initially only for water vapor and carbon dioxide, the fluxes show the expected diurnal cycle and are widely similar for all days. In contrast to this the gradients only follow these diurnal cycles for some selected periods and show an irregular pattern. This is a strong indication that the concentration measurements near the forest floor are decoupled from the flux and concentration measurements above the canopy. Because the sign of the flux often disagrees with the direction of the gradient, typical counter gradient fluxes occur (Denmead and Bradley, 1985). Further analysis of such situations

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needs indicators for a deeper analysis other than the forcing parameters wind and radiation.

- The aerodynamic method of using concentration gradients (observed above the canopy) for the determination of turbulent exchange fluxes can be applied to the *Waldstein-Weidenbrunnen* site only within certain limits, simply because the trees are too high to develop a site-specific roughness-sublayer correction. Furthermore, as shown by Ruppert et al. (2006) for temperature, CO₂ and H₂O, the scalar similarity is not fulfilled for all scalars and not in all spectral ranges. Although not shown here, the application of the Modified Bowen-ratio method (Businger, 1986; Müller et al., 1993) to observed above canopy concentration gradients of the NO–NO₂–O₃ triad also failed due to insufficient significance of the (very small) concentration differences. Nevertheless, the analysis of exchange fluxes of the NH₃–HNO₃–NH₄NO₃ triad between the forest canopy and the atmosphere was successful and indicated the importance of dry deposition of total ammonium and total nitrate as an input of reactive nitrogen to the *Waldstein-Weidenbrunnen* ecosystem (Wolff et al., 2010b).
- Within EGER-IOP-2, one of the first eddy covariance data sets (over any ecosystem) was obtained which consists of high resolution measurements of the complete NO–NO₂–O₃ triad. Data have been quality assured by application of the EUROFLUX methodology (Aubinet et al., 2000) and recent updates of the TK2 (latest version: TK3) software (Mauder and Foken, 2004; Mauder et al., 2008; Mauder and Foken, 2011) and delivered significant NO, NO₂ and O₃ fluxes over the *Waldstein-Weidenbrunnen* spruce forest. However, a considerable part of flux measurements might be biased by advection of traffic related NO from the nearby district road. Nevertheless, the results confirm the picture of the exchange of the NO–NO₂–O₃ triad over tall vegetation, which is to some extent “contrary” to that observed over low vegetation (e.g. Mayer et al., 2011). While over low vegetation, usually upward directed net NO fluxes (due to biogenic NO emission from soil)

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and downward directed net fluxes of NO_2 and O_3 (due to stomata uptake) are observed, there are still downward net O_3 fluxes over tall vegetation, but upward net NO_2 fluxes and downward net NO fluxes. Through a modelling study, Jacob and Bakwin (1991) firstly provided an explanation for this behaviour, namely the oxidation of soil emitted NO by O_3 (from aloft) in the low turbulence regime of the trunk space, which was first experimentally confirmed by Meixner et al. (2003) through measurements of net O_3 and NO fluxes above and NO- NO_2 - O_3 concentration profiles within and above a Brazilian primary rainforest. As in the dense rainforest, biogenic NO from the forest floor of the *Waldstein-Weidenbrunnen* site is completely oxidized to NO_2 , most likely within the first half meter above the forest floor, and escapes in form of NO_2 to the surface layer above the canopy. First quantification of the NO_2 uptake by spruce needles (field measurements, EGER IOP-2) allowed the estimate of the soil related contribution of the net NO_2 flux over the canopy which agrees well with the soil emitted NO flux (estimated from laboratory measurements).

- Classification of the above canopy net NO_2 fluxes with respect to coupling stages revealed 10-fold higher fluxes for fairly and fully coupled conditions (Ds, Cs/C), compared to the de-coupled conditions (Wa/Dc). This is a very clear indication that deep canopy mixing was necessary to export the soil emitted NO from the forest floor into the surface layer above the canopy.
- The diurnal evolution of HONO mixing ratio differences in and above the forest canopy could be well explained by the different source and sink processes (e.g. photolysis, heterogeneous formation and advection) in combination with the coupling regimes (Sörgel et al., 2011). During periods of complete coupling of the forest to the atmosphere (C and Cs), mixing ratio differences were close to zero, despite large differences (factor 10 to 25) in the photolytic sink during daytime. This was less obvious in the beginning of the dry period since HONO values were still influenced by the previous rain period (wet surfaces, low mixing ratio levels).

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During night-time, higher values above canopy could be partly explained by advection of HONO rich air, whereas higher values below canopy in the afternoon when the forest was decoupled (mostly Ds and later afternoon Dc and Wa) from the atmosphere pointed to a local source at the forest floor. In summary, the magnitude of the differences in mixing ratios above and below canopy were determined by the amount of vertical mixing, whereas the sign was determined by different source or sink processes.

- Eddy-covariance measurements are, in combination with footprint and data quality analysis, a well established tool for flux investigation, not only above the forest but also – in the case of developed turbulence – within the forest. This was shown not only with the measurements themselves but also in comparison with model results (Staudt et al., 2011).
- Keeping the uncertainties of the measurements in mind, the performance of the one-dimensional ACASA and the three-dimensional STANDFLUX models for water exchange of the forest depends on the time of the day and the coupling conditions. Both models fall short of describing night-time evapotranspiration measurements, yet there is a better performance of the third-order closure model ACASA with an advanced representation of turbulence at night. Regardless of the individual model setup, i.e. three-dimensional versus one-dimensional representation of the stand, both model performances improved considerably during daytime, particularly for coupled and partly coupled situations.

Summarizing the first results of EGER, the combination of a scale (micrometeorological) concept, based on detailed investigations of the structure of atmospheric turbulence, and measurements of reactive and non-reactive trace gases showed interesting and new results which could only be briefly covered in this study. Classification of (i) exchange fluxes observed above the canopy, (ii) turbulent and chemical time scales, and (iii) concentration differences (above-canopy versus forest floor) according to coupling stages, provides a powerful tool for qualitative (and most likely also quantitative)

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analysis of exchange processes of non-reactive, and particularly of reactive, trace compounds, and hence allows a better understanding than by interpretation of time series alone. There is certainly more effort and research necessary to make this tool a standard interpretive tool.

5 Finally, it should be emphasized that the investigation of the surface exchange of reactive trace gases, particularly those which are related to nearby anthropogenic (traffic) sources, could be strongly biased (if not endangered) by horizontal advection of tempo-
10 rally highly variable concentrations. Furthermore, there are different phenomena (LLJ, gravity waves) in the atmospheric boundary layer which might have drastic effects on the vertical distribution of non-reactive and reactive trace compounds and consequently
15 on the observed exchange fluxes. Future field experiments should be designed such that these phenomena could be quantitatively considered, which may have decisive effects on above and in-canopy concentration profiles as well as exchange fluxes.

However, the potential influence of the nearby clear-cut with convection and advec-
20 tion events, and probably a circulation system at times – as suggested as a result of the energy balance closure – between the clear-cut and the forest has not been addressed so far. To investigate these influences a special experiment was realized in June/July
25 2011 in combination with Large-Eddy Simulations. However there are still issues which need further investigations, such as gradients and fluxes close to the ground. These
30 issues are under investigation. And finally, the large data set also offers the possibility for cooperation with other partners.

Appendix A

In-canopy turbulence structure

25 The turbulence structure can typically be described with the so-called integral turbulence characteristics (normalized standard deviations), which are nearly constant
(Panofsky and Dutton, 1984) or have a small sensitivity to stratification (for an overview

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see Foken, 2008b). While the profiles of the mean wind speed above a canopy are strongly affected by the existence of the roughness sublayer (see Sect. 3.1), the effect on integral characteristics is not well investigated and only a few investigations are
5 available (Finnigan, 2000; Raupach et al., 1996). The turbulence characteristics within a forest are spatially heterogeneous and distinct from those associated with the surface
10 boundary layer (Baldocchi and Meyers, 1998). For measurements inside the canopy ($z < h_c$) a parameterization was proposed by Rannik et al. (2003)

$$\frac{\sigma_i}{u_*} = \alpha_i \left\{ \exp \left[-\alpha_i \left(1 - \frac{z}{h_c} \right)^{\beta_i} \right] (1 - \gamma_i) + \gamma_i \right\}$$

$i = u, v, w; z < h_c$ (A1)

10 and above the canopy constant values were assumed

$$\frac{\sigma_i}{u_*} = \alpha_i$$

$i = u, v, w; z < h_c$ (A2)

The values are given in Table A1. These are based on the measurements shown in Fig. 3.

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Table 1. Climate data for *Waldstein-Pflanzgarten* (period 1971–2000, Foken, 2003) and climate trends (Foken, 2004; Seifert, 2004).

Parameter	Climate (1971–2000)	Climate trend
height a.s.l.	765 m	
climate zone *	Dc	
annual mean temperature in °C	5.3	0.33 K/10 a, 99 % sign.
annual temperature amplitude in K	18.1	winter: 0.52 K/10 a, 95 % sign.
annual sum of precipitation in mm	1162.5	19 mm/10 a, not significant
month with maximum of precipitation	December, July	
snow cover **	approx. 80 days	–10 days/10 a, 95 % sign.

* Classification by Köppen/Trewartha/Rudloff according to Hendl (1991)

* Fichtelberg-Hüttstadt, 662 m a.s.l.

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Table 2. Overview of the meteorological conditions at *Waldstein-Pflanzgarten* site during IOP-1 and IOP-2.

Parameter	IOP-1 (6 Sep to 7 Oct 2007)	IOP-2 (1 Jun to 15 Jul 2008)
mean temperature in °C relation to normal	9.9 °C Sep too cold (t1.2 K); Oct too cold (t0.6 K)	15.2 °C Jun too warm (+1.5 K); Jul normal (+0.2 K)
absolute maximum temperature in °C	21.0 °C at 24 Sep 2007	28.9 °C at 22 Jun 2008
absolute minimum temperature in °C	2.8 °C at 19 Sep 2007	3.7 °C at 14 Jun 2008
precipitation sum in mm relation to normal	113.2 mm Sep too wet, from 30 Sep no rain; Oct to dry.	112.8 mm Jun too dry, no rain from 18 Jun to 3 Jul except a thunderstorm at 25 Jun 2008; Jul normal
maximum daily sum precipitation in mm	22.1 mm at 27 Sep 2007	44.1 mm at 3 Jul 2008
mean incoming shortwave radiation in $W m^{-2}$ (24 h)	80.4 $W m^{-2}$	225.6 $W m^{-2}$
maximum daily mean incoming shortwave radiation in $W m^{-2}$	152.5 $W m^{-2}$ 15 Sep 2007	at 342.7 $W m^{-2}$ at 1 Jun 2008

26314

Table 3. Overview of trace gas concentrations (ppb) and total of wet deposition (ionic components; mg m^{-2}) during the Intensive Observation Periods of the EGER project (IOP-1, IOP-2), measured at the *Waldstein-Weidenbrunnen* site (31 m a.gr.) and *Waldstein-Pflanzgarten* site (wet deposition, 1.5 m a.gr.), respectively.

Trace gas	IOP-1 6 Sep–3 Oct 2007	IOP-2 1 Jun–11 Jul 2008
ozone (O_3)	33 ppb (average) 50 ppb (95 % quantile)	55 ppb (average) 76 ppb (95 % quantile)
maximum O_3	59 ppb (23 Sep 2007)	85 ppb (6 Jun 2008)
nitric oxide (NO)	0.3 ppb (average) 1.1 ppb (95 % quantile)	0.2 ppb (average) 0.6 ppb (95 % quantile)
maximum NO	2.5 ppb at (2 Oct 2007)	3.4 ppb (7 Jun 2008)
nitrogen dioxide (NO_2)	3.1 ppb (average) 6.7 ppb (95 % quantile)	2.5 ppb (average) 4.6 ppb (95 % quantile)
maximum NO_2	28.5 ppb (28 Sep 2007)	15.6 ppb (9 Jun 2008)
sulphur dioxide (SO_2)	N/A	0.8 ppb (mean) 2.6 ppb (95 % quantile)
maximum SO_2	N/A	10.8 ppb (9 Jun 2008)
total of wet deposition (ionic components)	IOP-1	IOP-2
Cl^-	55 mg m^{-2}	$< 22 \text{ mg m}^{-2}$
NO_2^-	$< 7 \text{ mg m}^{-2}$	$< 4 \text{ mg m}^{-2}$
NO_3^-	297 mg m^{-2}	187 mg m^{-2}
SO_4^{2-}	199 mg m^{-2}	114 mg m^{-2}
NH_4^+	107 mg m^{-2}	78 mg m^{-2}
K^+	$< 30 \text{ mg m}^{-2}$	$< 18 \text{ mg m}^{-2}$
Mg^{2+}	5 mg m^{-2}	$< 6 \text{ mg m}^{-2}$
Ca^{2+}	32 mg m^{-2}	36 mg m^{-2}

26315

Table 4. Towers at the *Waldstein-Weidenbrunnen* site during the Intensive Observation Periods of the EGER project (IOP-1: 2007; IOP-2: 2008).

	Coordinates	Height [m]	Major equipment
“main tower”	50°08'31.2" N 11°52'00.8" E 775 m a.s.l.	31	FLUXNET station Wind-, dry and wet bulb temperature profiles Profiles of trace gas concentrations and fluxes Measurements of soil related quantities and trace gas soil flux measurements (dynamic chambers), within a radius of 10–50 m
“turbulence tower”	50°08'29.9" N 11°52'03.1" E	35	Profile of turbulent energy and carbon dioxide fluxes
“bio-tower”	50°08'32.9" N 11°51'57.8" E	36	Vertical profile of sapflow, temperature, humidity, and net radiation CO_2 and H_2O exchange measurements on spruce branches (dynamic cuvette)

26316

Table 5. Remote sensing techniques used during the Intensive Observation Periods of the EGER project (IOP-1: 2007; IOP-2: 2008).

Device	Coordinates	Location
2-D sonic anemometer (Thies)	50°08'31.2" N, 11°52'00.8" E 775 m a.s.l. + 32 m	<i>Waldstein-Weidenbrunnen</i> , „main tower“
mini-sodar, SFAS (Scintec AG)	50°08'30.3" N, 11°52'11.0" E 785 m a.s.l.	clearing, NE of <i>Waldstein-Weidenbrunnen</i> site
Sodar/RASS, DSDPA90.64 (METEK GmbH)	50°08'35.3" N, 11°51'48.8" E 764 m a.s.l.	<i>Waldstein-Pflanzgarten</i>
UHF-windprofiler (Vaisala)	49.98° N, 11.68° E 514 m a.s.l.	<i>Oschenberg near Bayreuth</i>

26317

Table 6. Turbulent flux measuring instrumentation at the *Waldstein-Weidenbrunnen* site during the Intensive Observation Periods of the EGER project (IOP-1: 2007; IOP-2: 2008).

	Height [m]	Sonic anemometer	Trace gas	Trace gas analyzer
“main tower” (IOP-1)	32	Solent R2, Gill Instruments Ltd.	CO ₂ , H ₂ O	LI-7500, L-COR Inc.
	1****	Solent R2, Gill Instruments Ltd.	O ₃ O ₃	GEFAS GmbH*** GEFAS GmbH***
“main tower” (IOP-2)	32	USA-1, METEK GmbH	CO ₂ , H ₂ O	LI-7500, L-COR Inc.
	32	CSAT3, Campbell Sci. Inc.	NO, NO ₂ O ₃	CLD 790SR-2, Ecophysics NOAA/ATDD***
	25	Solent R2, Gill Instruments Ltd.	O ₃	Enviscope GmbH***
	17	Solent R2, Gill Instruments Ltd.	O ₃	Enviscope GmbH***
	1	Solent R2, Gill Instruments Ltd.	O ₃	GEFAS GmbH***
“turbulence tower” (IOP-1, IOP-2)	36	USA-1, METEK GmbH	CO ₂ , H ₂ O	LI-7500, L-COR Inc.
	23	CSAT3, Campbell Sci. Inc.	CO ₂ , H ₂ O	LI-7500, L-COR Inc.
	18	Solent R3-50, Gill Instrum. Ltd.	CO ₂ , H ₂ O	LI-7500, L-COR Inc.
	13	CSAT3, Campbell Sci. Inc.	CO ₂ , H ₂ O	LI-7500, L-COR Inc.**
	5.5	CSAT3, Campbell Sci. Inc.*	CO ₂ , H ₂ O	LI-7500, L-COR Inc.
	2.5	CSAT3, Campbell Sci. Inc.	CO ₂ , H ₂ O	LI-7500, L-COR Inc.

* During IOP-1 Solent R2, Gill Instruments Ltd.;

** during IOP-1 KH20 Campbell Scientific, Inc.;

*** all fast-response O₃-analyzers were based on solid-phase chemiluminescence technique (Güsten and Heinrich, 1996), but designed by different manufacturers (indicated);

**** O₃-flux measurements at 1 m have been performed one week before those at 32 m (using the same instrumentation).

26318

Table 7. Profile measurements at the “main tower” of the *Waldstein-Weidenbrunnen* site during the Intensive Observation Periods of the EGER project (IOP-1 and IOP-2).

	IOP-1 6 Sep–7 Oct 2007	IOP-2 1 Jun–15 Jul 2008
<i>measurement levels at “main tower” (in m)</i>		
wind speed I	4.6, 10.0, 16.5, 18.0, 21.0, 25.0, 31.0	4.6, 10.0, 16.5, 18.0, 21.0, 25.0, 31.0
wind speed II	7.6, 13.3, 19.8, 24.3, 26.3, 31.3	7.6, 13.3, 19.8, 24.3, 26.3, 31.3
dry & wet bulb temperature I (aspirated)	5.0, 13.0, 21.0, 31.0	5.0, 13.0, 21.0, 31.0
dry & wet bulb temperature II (aspirated)	4.9, 9.9, 15.9, 19.5, 24.4, 26.6, 30.9	4.9, 9.9, 15.9, 19.5, 24.4, 26.6, 30.9
CO ₂ , H ₂ O, O ₃ , NO, NO ₂	5.0, 10.0, 16.0, 24.2 (2x), 31.0	3.0, 10.0, 16.5 (2x), 20.5, 25.0, 31.5
NH ₄ NO ₃ , NH ₃ and HNO ₃	24.2, 30.4	N/A
HONO	24.5	N/A
NO, NO ₂ , O ₃ , CO ₂ and H ₂ O exchange measurements on spruce branches (dynamic cuvette)	13.0	N/A
<i>measurement levels at “forest floor” (in m)</i>		
wind speed I	2.0	2.0
wind speed II	0.04, 0.30, 1.0, 2.00	0.04, 0.3, 1.0, 2.0
dry & wet bulb temperature I (aspirated)	0.05, 2.0	0.05, 2.0
dry & wet bulb temperature II (aspirated)	0.09, 0.26, 1.00, 2.05	0.09, 0.26, 1.00, 2.05
air temperature (non-aspirated)	0.01, 0.02, 0.04, 0.08, 0.16, 0.32	0.01, 0.02, 0.04, 0.08, 0.16, 0.32
CO ₂ , H ₂ O, O ₃ , NO, NO ₂	0.05, 0.3, 1.0, 2.0	0.005, 0.03, 0.1, 0.3, 0.9
²²⁰ Rn/ ²²² Rn	0.04, 0.27	0.00, 0.03, 0.1, 0.3
HONO	0.5	1.0
<i>measurement levels at “bio-tower” (in m)</i>		
air temperature & rel. humidity (aspirated)	9.3, 11.7, 14.8, 17.2, 20.2, 22.6	9.3, 11.7, 14.8, 17.2, 20.2, 22.6
net radiation	11.9, 17.4, 22.8	11.9, 17.4, 22.8
CO ₂	N/A	0.03, 1.0, 16.0, 18.7, 21.4, 24.1, 29 ± 3, 36.0

26319

Table A1. Coefficients for Eq. (A1).

Reference	<i>i</i>	<i>a_i</i>	<i>α_i</i>	<i>β_i</i>	<i>γ_i</i>
Rannik et al. (2003), neutral, for <i>Hyytiälä</i> site	<i>u</i>	2.30	1.0	1.0	−0.3
	<i>v</i>	1.75	1.0	0.85	−0.2
	<i>w</i>	1.25	0.9	1.2	−0.63
<i>Waldstein-Weidenbrunnen</i> site, IOP-1	<i>u</i>	2.01	8.97	1.37	0.29
	<i>v</i>	1.60	5.18	1.11	0.34
	<i>w</i>	1.13	0.9	1.2	−0.63

26320

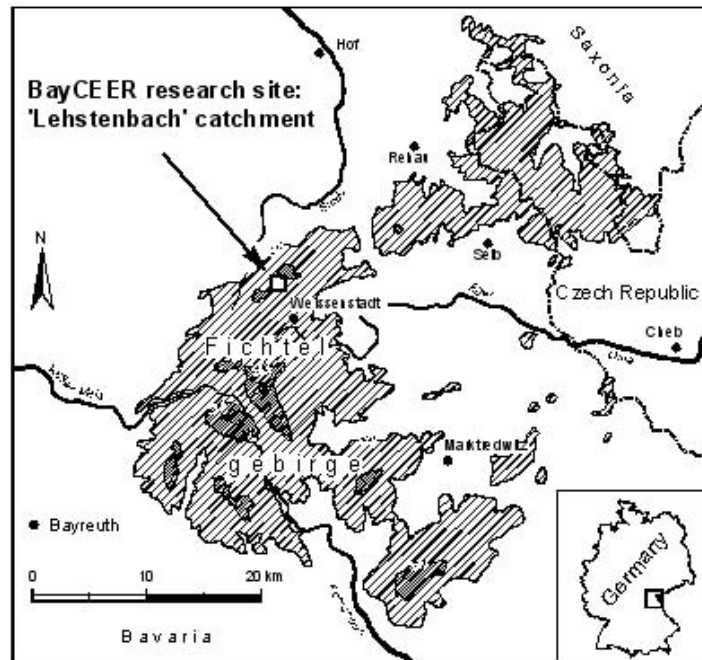


Fig. 1. Map of the “Fichtelgebirge” region (Gerstberger et al., 2004). The *Waldstein-Weidenbrunnen* site is located in the *Lehstenbach* catchment north-west of the small town *Weissenstadt* and the upper *Eger river* valley. Hatchings indicates elevations >500 m and >750 m a.s.l., respectively.

26321

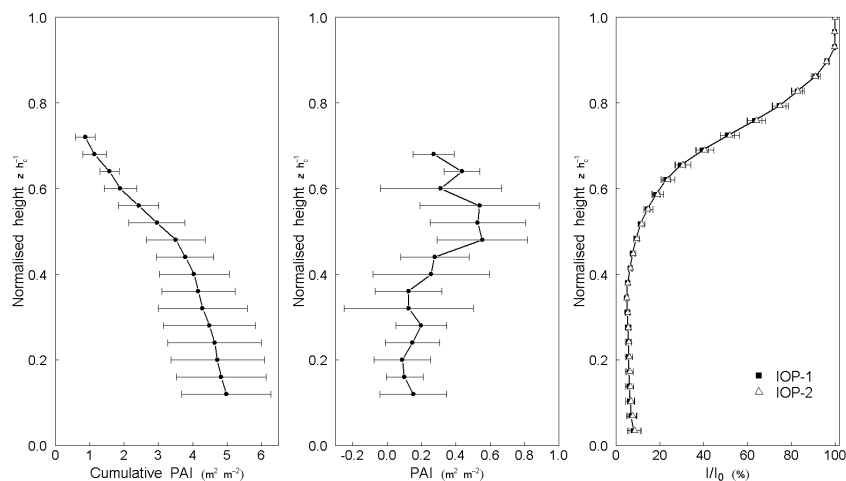


Fig. 2. Vertical profile of the (left) cumulative and (middle) absolute overstory plant area index (PAI) of the *Waldstein-Weidenbrunnen* site. Profiles are mean values of the cumulative and absolute PAI profile measurements. Furthermore, the right figure shows the vertical distribution of the normalized shortwave radiation for both IOPs. The height is normalized by a canopy height of 25 m.

26322

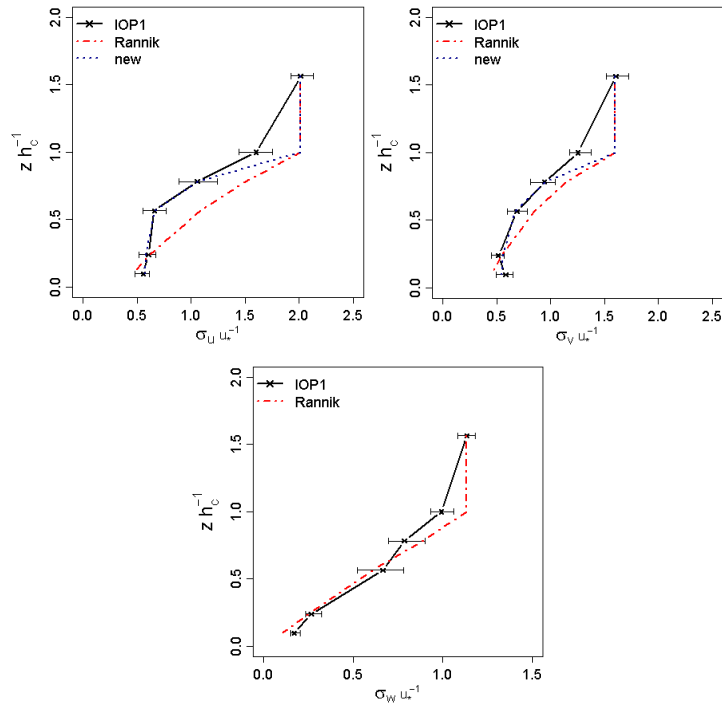


Fig. 3. Results of measurements (x, black line) and model parameterizations (red dot-dashed line) for profiles of the integral turbulence characteristics of wind velocity components u, v and w within the *Waldstein-Weidenbrunnen* forest stand and inside the lower roughness sublayer (near neutral stratification). For u and v the parameterization of Rannik et al. (2003) was modified to fit the measurements during IOP-1, for w the original coefficients were used, see Appendix.

26323

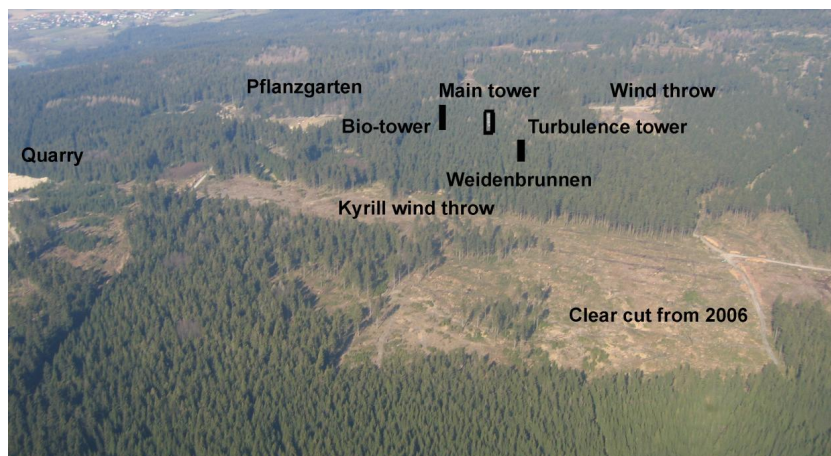


Fig. 4. The *Waldstein-Weidenbrunnen* site shortly after the “hurricane like” low pressure system “Kyrill” (view from south, photograph: Foken, T., 15 March 2007). The “main tower” can be seen within the *Waldstein-Weidenbrunnen* field site; “turbulence” and “bio” towers were set-up after 15 March 2007, but their locations are marked. The Modified Bowen-ratio system has been located at the wind throw north-east of the “turbulence tower”.

26324

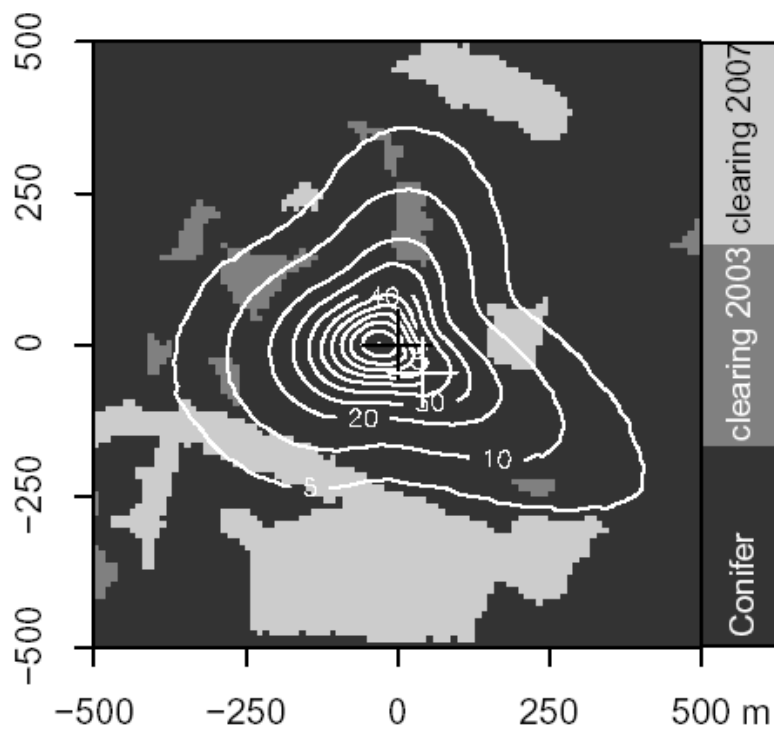


Fig. 5. Clear cuts in the planning phase of EGER (2003) and after “Kyrill” (18 January 2007). “Emma” on 1 March 2008 has affected the SW corner of the map. The black and white crosses indicate main and turbulence tower positions, respectively. White isolines indicate relative flux contribution to the footprint for 2003 data (Siebicke, 2008).

26325



Fig. 6. Towers at the *Waldstein-Weidenbrunnen* site: **(a)** permanent main tower since 1996 (height 31 m) for routine measurements, FLUXNET station and chemical measurements, **(b)** turbulence tower (height 35 m) since 2007 with second FLUXNET complex and turbulence measurements, **(c)** biological tower (height 36 m) for gas exchange measurements at needles.

26326

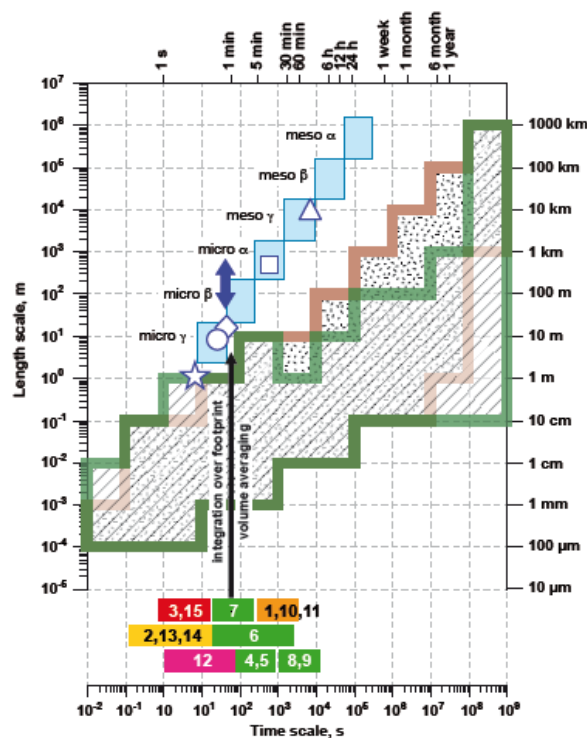


Fig. 7. Caption on next page.

26327

Fig. 7. Temporal and spatial scales of atmospheric, turbulent, plant, physiological, soil and relevant chemical processes forming the conceptual backbone of the EGER project. Atmospheric Processes (Orlanski, 1975) are given in light blue squares of one order of magnitude (from *micro γ* to *meso α*). Forest canopy related transport processes comprise turbulent transport in canopy (white star), vertical advection in canopy (white circle), transport above canopy (white diamond), coherent structures (blue double arrow), footprint averaged turbulent flux (white square), and horizontal advection at canopy top (white triangle). The scales of plant processes, relevant for energy and matter exchange with the atmosphere (Schoonmaker, 1998), those of soil processes are shown by the brown framed, spotted area (Vogel and Roth, 2003; Blöschl and Sivapalan, 1995). Time scales of relevant chemical reactions are shown according to Dlugi (1993, updated): (1) $\text{HO}_2 + \text{HO}_2 \rightarrow \text{H}_2\text{O}_2 + \text{O}_2$, (2) $\text{HNO}_3 + \text{NH}_3 \leftrightarrow \text{NH}_4\text{NO}_3$, (3) $\text{O}_3 + \text{NO} \rightarrow \text{NO}_2 + \text{O}_2$, (4) $\text{O}_3 + \text{isoprene} \rightarrow \text{reaction products (P)}$, (5) $\text{O}_3 + \text{monoterpenes} \rightarrow \text{P}$, (6) $\text{NO}_3 + \text{monoterpenes} \rightarrow \text{P}$, (7) $\text{NO}_3 + \text{isoprene} \rightarrow \text{P}$, (8) $\text{OH} + \text{isoprene} \rightarrow \text{R}$, (9) $\text{OH} + \text{monoterpenes} \rightarrow \text{R}$, (10) $\text{O}_3 + \text{olefins} \rightarrow \text{R}$, (11) $\text{O}_3 + \text{NO}_2 \rightarrow \text{NO}_3 + \text{O}_2$, (12) $\text{N}_2\text{O}_5 + \text{H}_2\text{O} \rightarrow 2\text{HNO}_3$, (13) $\text{HNO}_3 (+ \text{H}_2\text{O}) \rightarrow \text{H}^+ + \text{NO}_3^-$, (14) $\text{H}_2\text{O} + 2\text{NO}_2 (\text{het}) \rightarrow \text{HNO}_2 + \text{HNO}_3 (\text{g})$, (15) $\text{NO}_2 + h\nu \rightarrow \text{NO} + \text{O}$ (for details of concentration ranges see Dlugi, 1993).

26328

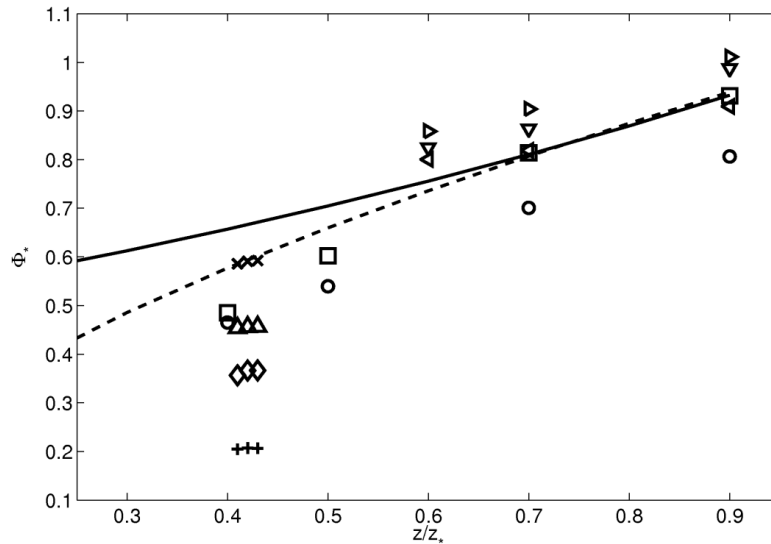


Fig. 8. The calculated ϕ_* values of the EGER data are presented by crosses, upward-pointing triangles, diamonds, and plus signs. The crosses represent u_* values smaller than 0.4 and larger than 0.2. Further the upward-pointing triangles represent $0.4 < u_* < 0.6$, the diamonds $0.6 < u_* < 0.8$ and the plus signs $0.8 < u_* < 1.2$. The calculated ϕ_* values of the COPS data for a canopy height of 1.2m are represented by squares and circles. The squares represent $0.2 < u_* < 0.3$ and the circles represent $0.3 < u_* < 0.6$. The calculated ϕ_* values of the COPS data for a canopy height of 2.9m are represented by downward-pointing, right-pointing, and left-pointing triangles. The downward-pointing triangles represent $0.2 < u_* < 0.3$, the right-pointing triangles display $0.3 < u_* < 0.4$ and left-pointing triangles $0.4 < u_* < 0.5$. Furthermore, the solid line represents the equation developed by Garratt (1992) and the dashed line represents the equation developed by Mölder et al. (1999).

26329

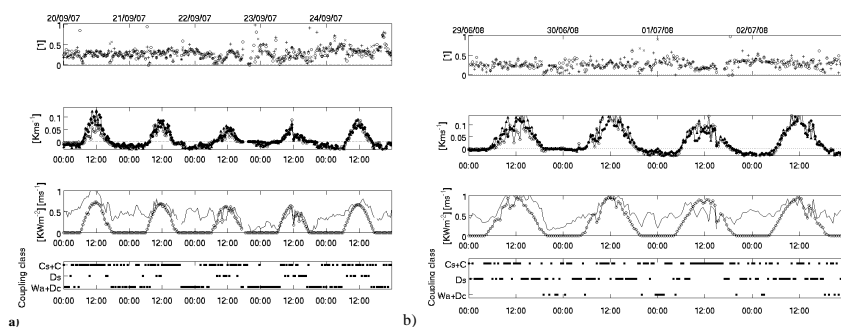


Fig. 9. Measurements and characterisation of the turbulent exchange for the period 20 to 24 September 2007 during IOP-1 (a) and 29 June to 2 July 2008 during IOP-2 (b). Upper panel: relative flux contribution of coherent structures $F_{cs}F_{tot}^{-1}$ for carbon dioxide (open circles), buoyancy (crosses) and latent heat (pluses) at $1.44h_c$; second panel: kinematic buoyancy flux of coherent structures H_{cs} at $1.44h_c$ (filled circles), $0.93h_c$ (grey triangles) and $0.72h_c$ (open circles); third panel: friction velocity u_* (solid line), incoming shortwave radiation Rg (open circles) and wind direction (filled circles) at $1.44h_c$; lowest panel: coupling situations.

26330

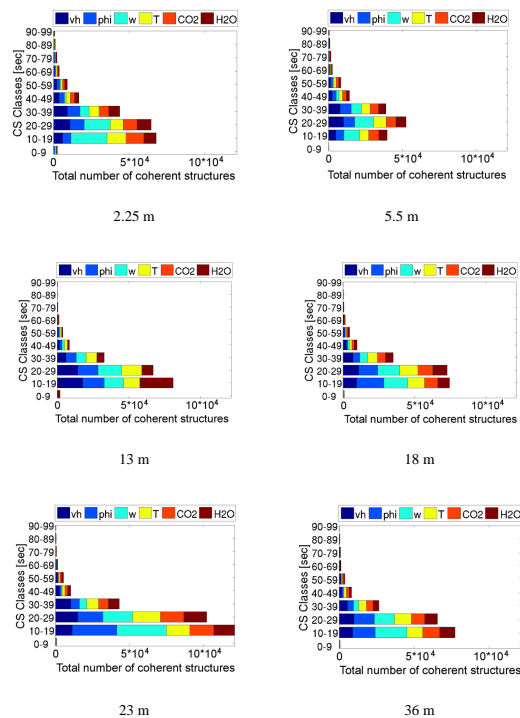


Fig. 10. Total number of coherent structures (CS) detected from 14 September 2007 until 08 October 2007 in carbon dioxide CO_2 , water vapour H_2O , wind direction ϕ , sonic temperature T_s , horizontal wind velocity v_h , and vertical wind velocity w from measurements at the turbulence tower on six levels.

26331

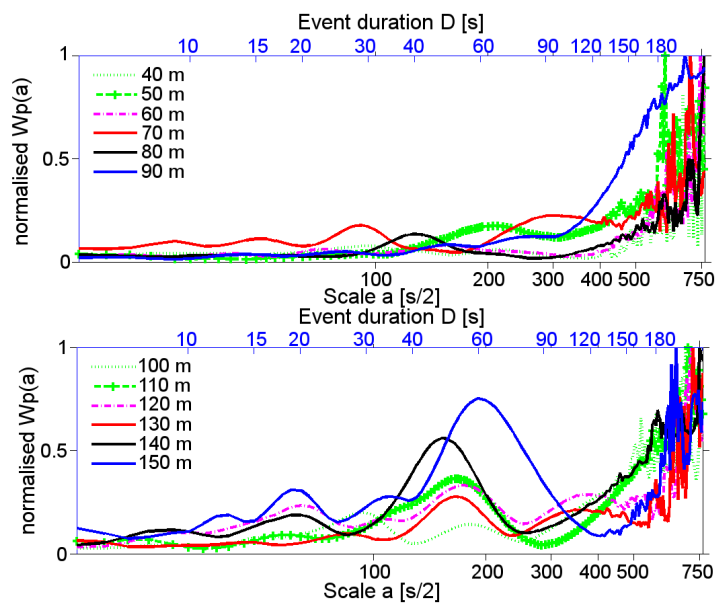


Fig. 11. Normalized spectra of the wavelet variance for different observation levels for the vertical wind during IOP-1 14 September 2007, 00:00–00:25 CET.

26332

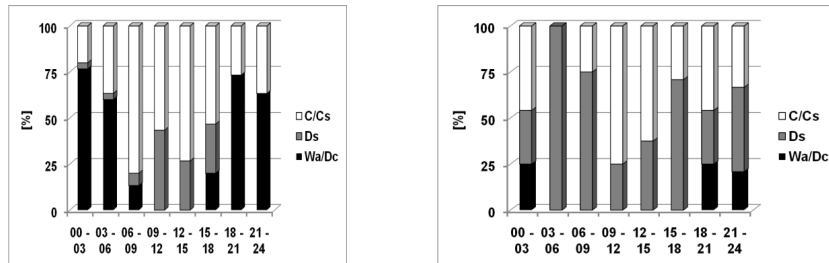


Fig. 12. Percentage of different coupling classes during the Golden Days of IOP-1 (left, 30 three-hour periods) and IOP-2 (right, 24 three-hour periods) in the daily cycle.

26333

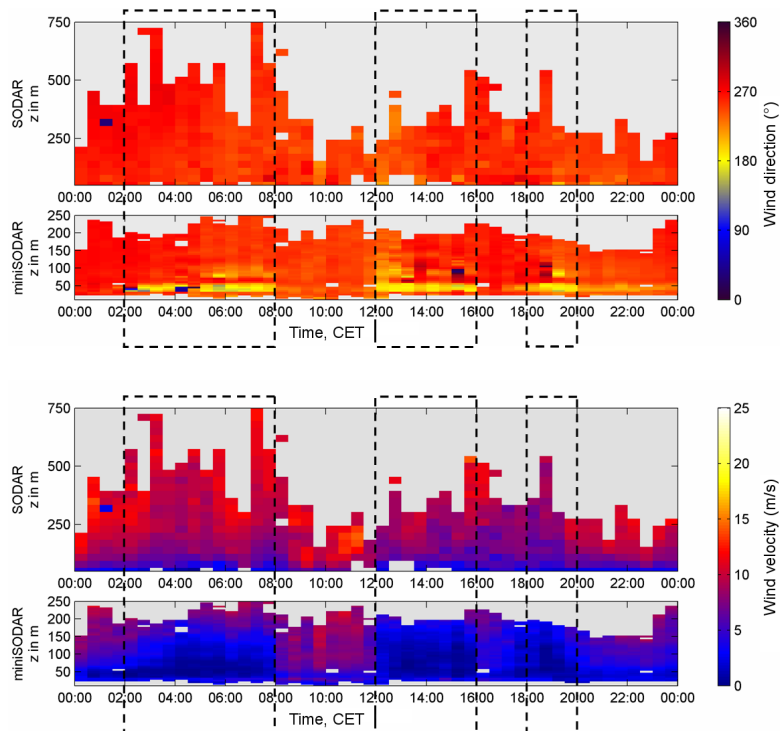


Fig. 13. Time-high profile from sodar and mini-sodar data of the wind direction (above) and the wind velocity (below) on 28 June 2008. Periods with southerly winds directly above the canopy originating from the clear-cut are highlighted.

26334

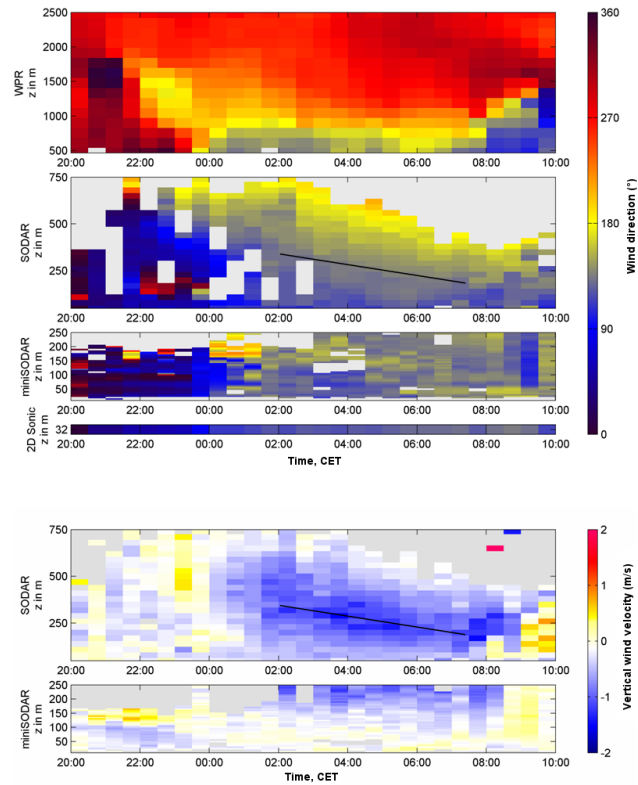


Fig. 14. Time-high profile from windprofiler, sodar, mini-sodar and sonic data of the horizontal wind velocity (above), the wind direction (middle), and vertical wind velocity (below) in the night from 30 June to 1 July 2008. The axis of the low-level jet is highlighted.

26335

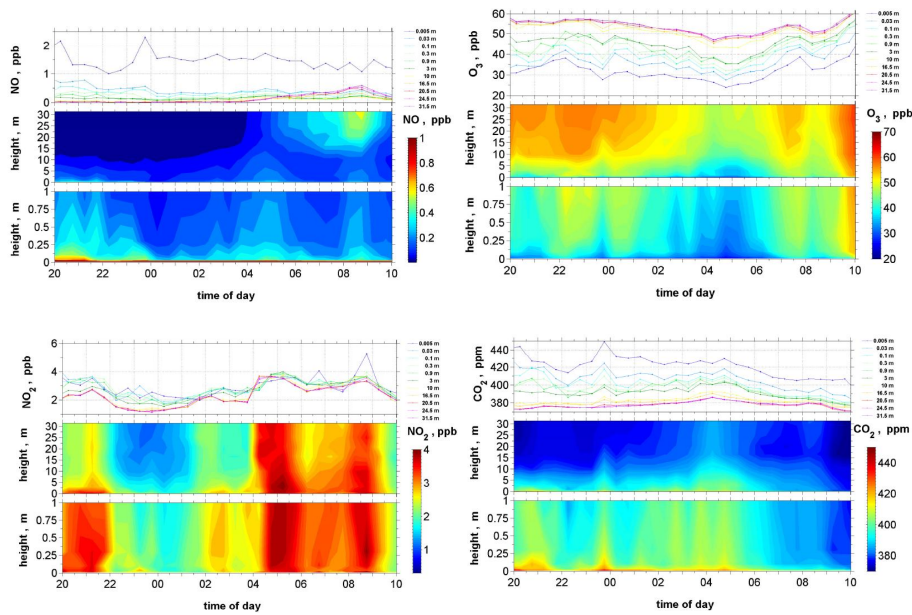


Fig. 15. Averaged vertical profiles in the night from 30 June to 1 July 2008 (compare with Fig. 14) for NO, NO₂, O₃ and CO₂ time in CET. For each trace gas the concentration at different levels, the concentration in and above the 25 m high forest and close to the forest floor are shown.

26336

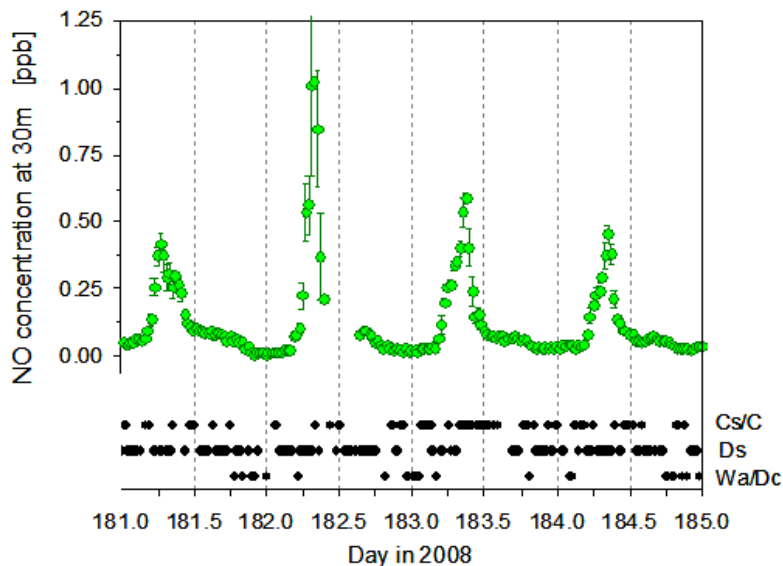


Fig. 16. Diel variation of NO concentration measured at *Waldstein-Weidenbrunnen* main tower (above canopy; 30 m a.gr.) from 29 June to 2 July 2008 (IOP-2). Original 5 Hz data have been averaged to 30 min means, error bars indicate corresponding standard deviations. Every day, between 06:00 and 12:00 (CET), when the convective boundary layer was still emerging, particularly high NO concentrations with large standard deviations were observed due to advection of freshly emitted exhaust gases from a busy (2100 cars day⁻¹) district road (HO18) running from SSE to NNW between 1 and 2 km distance west to the *Waldstein-Weidenbrunnen* site. The occurrence of canopy coupling stages (Cs/C, Ds, Wa/Dc) is given at the bottom.

26337

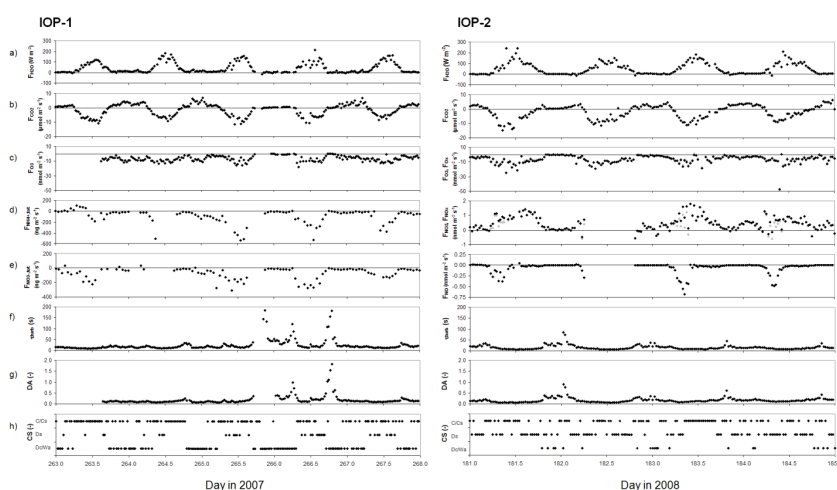


Fig. 17. Results of flux measurements of gaseous and particulate trace compounds at the “main tower” during 20–24 September 2007 (IOP-1, DOY 263–267, left panels) and for 29 June to 2 July 2008 (IOP-2, DOY 181–184, right panels). **(a)** $F_{\text{H}_2\text{O}}$, **(b)** F_{CO_2} , **(c)** F_{O_3} (black symbols) and F_{O_x} (grey symbols, IOP-2 only), **(d)** $F_{\text{NH}_4^+}$ (left panel) and F_{NO_2} and F_{NO_x} (right panel, black and grey symbols, respectively), **(e)** $F_{\text{NO}_3^-}$ (left panel) and F_{NO} (right panel), **(f)** turbulent time scale τ_{turb} (24–32 m), **(g)** Damköhler number DA (24–32 m), **(h)** coupling regimes (determined from measurements at the “turbulence tower”, see Sect. 3.2). Measurement heights were 32 m (a. gr.) for all fluxes measured by eddy covariance during IOP-1 **(a–c)**, and during IOP-2 **(a–e)**; fluxes of total ammonium and total nitrate **(d and e, IOP-1)**, determined by aerodynamic gradient technique, refer to 27.1 m (geometric mean of the heights of both intake levels).

26338

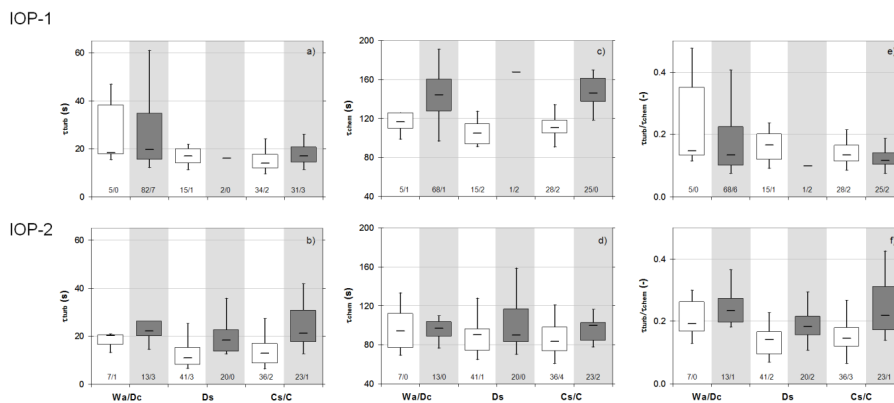


Fig. 19. Statistical dispersion and skewness of turbulent time scale τ_{turb} (**a, b**), chemical time scale τ_{chem} (**c, d**), and Damköhler number $DA (= \tau_{\text{turb}} / \tau_{\text{chem}}$; **e, f**) at the “main tower” for 20–24 September 2007 (IOP-1, DOY 263–267, top panels), and for 29 June to 2 July 2008 (IOP-2, DOY 181–184, bottom panels), classified by coupling regimes (Wa/Dc, decoupled conditions; Ds, decoupled subcanopy; Cs/C, coupled subcanopy by sweeps and fully coupled subcanopy) and day- (open box-plots) and night-time (grey shades). See remark on Fig. 18.

26341

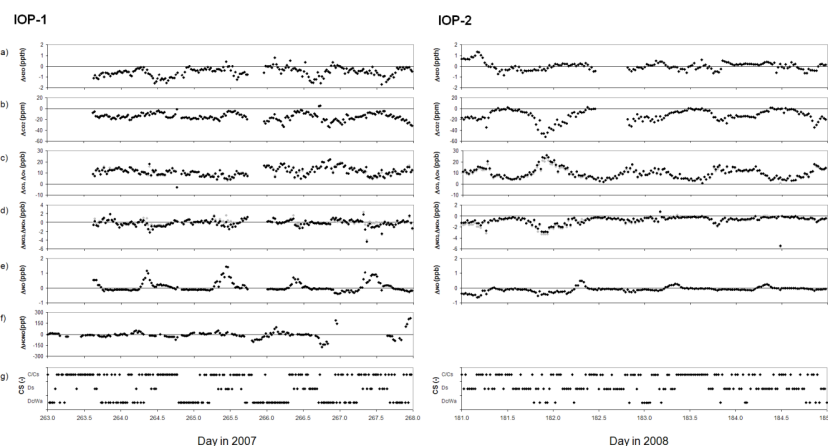


Fig. 20. Concentration differences $\Delta[C] = [C]_{24\text{m}} - [C]_{0.9\text{m}}$ between the 24 m and the 0.9 m level at the “main tower” for 20–24 September 2007 (IOP-1, DOY 263–267, left panels), and for 29 June to 2 July 2008 (IOP-2, DOY 181–184, right panels): (**a**) $\Delta[\text{H}_2\text{O}]$, (**b**) $\Delta[\text{CO}_2]$, (**c**) $\Delta[\text{O}_3]$ and $\Delta[\text{O}_x]$ (black and grey symbols, respectively), (**d**) $\Delta[\text{NO}_2]$ and $\Delta[\text{NO}_x]$ (black and grey symbols, respectively), (**e**) $\Delta[\text{NO}]$, (**f**) $\Delta[\text{HONO}]$ (IOP-1 only); (**g**) coupling regimes (determined from measurements at the “turbulence tower”, see Sect. 3.2).

26342

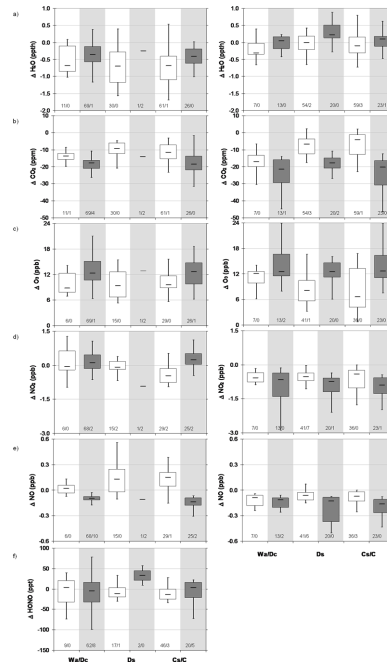


Fig. 21. Statistical dispersion and skewness of concentration differences Δ at the “main tower” for 20–24 September 2007 (IOP-1, DOY 263–267, left panels) and for 29 June to 2 July 2008 (IOP-2, DOY 181–184, right panels): $\Delta\text{H}_2\text{O}$ (a), ΔCO_2 (b), ΔO_3 (c), ΔNO_2 (d), ΔNO (e), and ΔHONO (f, IOP-1 only). Data were classified by coupling regimes (Wa/Dc, decoupled conditions; Ds, decoupled subcanopy; Cs/C, coupled subcanopy by sweeps and fully coupled subcanopy) and day- (open box-plots) and night-time (grey shades). See remark on Fig. 18.

26343

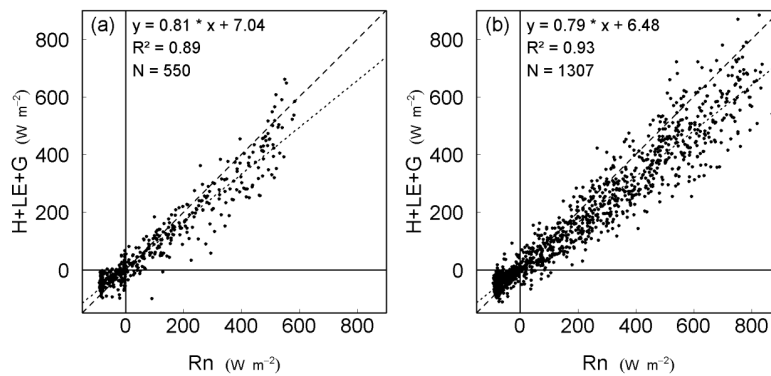


Fig. 22. Energy balance closure at the turbulence tower during IOP-1 (left) and IOP-2 (right).

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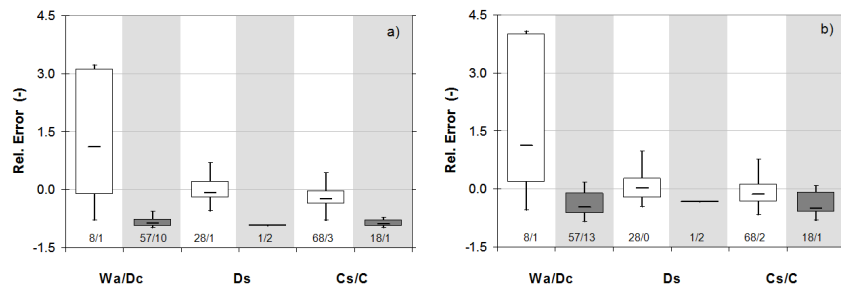


Fig. 23. Statistical dispersion and skewness of relative model error with respect to coupling stages (for definition see text) from **(a)** STANDFLUX, and **(b)** ACASA; measured and modelled evapotranspiration data ranged from 20 September through 24 September 2007 (IOP-1, DOY 263–267); measured evapotranspiration from turbulence tower 36 m level. Data were classified by coupling regimes (Wa/Dc, decoupled conditions; Ds, decoupled subcanopy; Cs/C, coupled subcanopy by sweeps and fully coupled subcanopy) and day- (open box-plots) and night-time (grey shades). See remark on Fig. 18.