The exchange of carbon dioxide between wet arctic tundra and the atmosphere at the Lena River Delta, Northern Siberia

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Abstract. The exchange fluxes of carbon dioxide between wet arctic polygonal tundra and the atmosphere were investigated by the micrometeorological eddy covariance method. The investigation site was situated in the centre of the Lena River Delta in Northern Siberia (72°22′ N, 126°30′ E). The study region is characterized by a polar and distinctly continental climate, very cold and ice-rich permafrost and its position at the interface between the Eurasian continent and the Arctic Ocean. The soils at the site are characterized by high organic matter content, low nutrient availability and pronounced water logging. The vegetation is dominated by sedges and mosses. The micrometeorological campaigns were performed during the periods July–October 2003 and May–July 2004 which included the period of snow and soil thaw as well as the beginning of soil refreeze. The main CO2 exchange processes, the gross photosynthesis and the ecosystem respiration, were found to be of a generally low intensity. The gross photosynthesis accumulated to −432 g m−2 over the photosynthetically active period (June–September). The contribution of mosses to the gross photosynthesis was estimated to be about 40%. The diurnal trend of the gross photosynthesis was mainly controlled by the incoming photosynthetically active radiation. During midday, the photosynthetic apparatus of the canopy was frequently near saturation and represented the limiting factor on gross photosynthesis. The synoptic weather conditions strongly affected the exchange fluxes of CO2 by changes in cloudiness, precipitation and pronounced changes of air temperature. The ecosystem respiration accumulated to +327 g m−2 over the photosynthetically active period, which corresponds to 76% of the CO2 uptake by photosynthesis. However, the ecosystem respiration continued at substantial rates during autumn when photosynthesis had ceased and the soils were still largely unfrozen. The temporal variability of the ecosystem respiration during summer was best explained by an exponential function with surface temperature, and not soil temperature, as the independent variable. This was explained by the major role of the plant respiration within the CO2 balance of the tundra ecosystem. The wet polygonal tundra of the Lena River Delta was observed to be a substantial CO2 sink with an accumulated net ecosystem CO2 exchange of −119 g m−2 over the summer and an estimated annual net ecosystem CO2 exchange of −71 g m−2.

1 Introduction

There is growing evidence that the climate system of the earth has changed significantly since the industrial revolution. The observed climate change is likely to be caused at least partially by human activity, which has substantially altered the atmospheric composition by the emission of radiatively active greenhouse gases such as carbon dioxide and methane (IPCC, 2007). The Arctic is of major interest within the context of global change because it is observed to warm more rapidly and to a greater extent than the rest of the earth surface (Maxwell, 1997; Serreze et al., 2000; Polyakov et al., 2003), and much larger changes are projected by climate model simulations (Kattenberg et al., 1996; Räisänen, 2001). Furthermore, its ecosystems are highly sensitive to climate change (Chapin et al., 1992; Oechel et al., 1997b) and play a key role in many global processes, such as the atmospheric and oceanic circulations (Stocker and Schmittner, 1997; Wood et al., 1999; Eugster et al., 2000)
or the regulation of the global budget of greenhouse gases (Gorham, 1991; Roulet et al., 1992; Tenhunen, 1996).

Most land surfaces in the Arctic are covered by tundra, treeless ecosystems whose vegetation consists primarily of grasses, sedges, small flowering herbs, low shrubs, lichens and mosses. Arctic and alpine tundra occupy 7.4×10^6 km^2 or about 7.4% of the land area of the northern hemisphere (Matthews, 1983; Loveland et al., 2000). Since the biota of the arctic ecosystems are closely adapted to their extreme environment, climatic changes will have a severe impact on the distribution, composition and functionality of plant and animal communities in the tundra (Callaghan and Jonasson, 1995; Chapin et al., 1995, 1997; Walker et al., 2001). This will cause major alterations of the energy, water and carbon balance of the Arctic land surfaces, which will feed back on the atmospheric system of the Arctic (Zhuang et al., 2006; Sturm et al., 2001b; Chapin et al., 2005). While the carbon balance changes in northern ecosystems are estimated to have only minor feedback effects on the global climate in the context of the projected anthropogenic carbon emissions (Zhuang et al., 2006), the changes of the water and energy balance by hydrological and vegetation changes in the Arctic are expected to be significant also on the global scale (Bonan et al., 1995; Lafleur and Rouse, 1995; Pielke and Vidale, 1995; Bröcker, 1997; Beringer et al., 2001; Peterson et al., 2002).

The tundra ecosystems are underlain by permafrost. Permafrost-affected soils often have a greater content of organic carbon than soils of temperate climate zones because organic matter decomposition is inhibited by cold temperatures, a short growing season, and water saturated soils. Correspondingly, the tundra ecosystems have historically been major sinks for carbon and nutrients. At least 14% of the global soil organic carbon is stored in the tundra (Post et al., 1982; Billings, 1987). However, permafrost is very susceptible to long-term warming, and an increased level of permafrost thawing might turn the tundra from a carbon sink to a source of carbon, either in the form of CO\(_2\) or as CH\(_4\) (Oechel et al., 1993; Christensen, 1993; Zimov et al., 1997). Since CO\(_2\) and CH\(_4\) are the most effective greenhouse gases besides water vapour (Rohde, 1990), an increased release of these gases by permafrost thawing would additionally amplify global warming.

In the last decade, numerous land-atmosphere flux studies relying on the eddy covariance method have been initiated, for example within the projects NOWES (Glooschenko, 1994), ABLE 3B (Harriss et al., 1994), BOREAS (Sellers et al., 1997), NOPEX (Hallidin et al., 1999) or EUROFLUX (Valentini, 2002). Most of the eddy covariance flux studies were and are conducted in the temperate and boreal zones of North America and Europe. Flux data for the Arctic regions are limited and are biased toward Alaska and Northern Fennoscandia. However, the vast tundra landscapes of Siberia are by far not adequately represented. This study presents the results of two micrometeorological field campaigns which were conducted in the wet arctic tundra of the North-Siberian Lena River Delta in 2003 and 2004. The study shall contribute to the understanding of the physical and biogeochemical interaction processes between permafrost soils, tundra vegetation, and the atmosphere which is necessary for assessing the impact of climate change on arctic tundra ecosystems and the possible feedbacks on the climate system. The objectives of the study were to characterize the exchange fluxes of CO\(_2\) on diurnal and seasonal time scales, to quantify gross photosynthesis, ecosystem respiration and net ecosystem CO\(_2\) exchange on the landscape scale, to analyze the regulation of the exchange fluxes by climatic forcings, and to estimate the annual CO\(_2\) budget of the tundra ecosystem.

2 Materials and methods

2.1 Study site

The investigation site was located on Samoylov Island in the Lena River Delta at 72°22′ N, 126°30′ E. With an area of 32,000 km\(^2\), the Lena River Delta is the largest delta in the Arctic and one of the largest in the world (Walker, 1998). In terms of its geological genesis, the Lena River Delta can be divided into three river terraces of different age, and various floodplain levels (Grigoriev, 1993; Schwamborn et al., 2002). The youngest terraces and active flood-plains which represent modern delta landscapes (Are and Reimnitz, 2000) occupy about 65% of the total area of the delta, predominantly in the central and eastern part (Fig. 1).

The Lena River Delta is located in the zone of continuous permafrost with permafrost depths of about 500 m (Grigoriev, 1960; Frolov, 2003; Zhang et al., 1999; NSIDC, 2003). With about −12°C, the permafrost temperature is very low. Colder permafrost is only encountered on the Taymyr Peninsula to the North-West of the Lena River Delta and on the Canadian Arctic Archipelago (Natural Resources Canada, 1995; Kotlyakov and Khromova, 2002). The soils of the region thaw to a depth of only 0.3–1.0 m during the summer.

The climate in the Lena River Delta is characterized by very low temperatures and low precipitation. The mean annual air temperature during 1961–1999, measured by the meteorological station on Samoylov Island from the period 1999-2005 showed a mean annual air temperature of −13.6°C, the mean annual precipitation in the same period was 319 mm (ROSHYDROMET, 2004). Data from the meteorological station on Samoylov Island from the period 1999–2005 showed a mean annual air temperature of −14.7°C and a highly variable total summer precipitation (rain) between 72 and 208 mm (mean 137 mm) (Boike et al., 2007\(^1\)). Polar day begins at 7 May and ends at 7 August.

and polar night lasts from 15 November to 28 January. The summer growing season lasts about three months, from the middle of June to the middle of September. The synoptic weather conditions in the Lena River Delta are characterized by its position at the border between the Arctic Ocean and the Siberian mainland. During summer, the central delta region experiences strongly varying weather conditions due to the change between the advection of cold and moist air masses from the Arctic Ocean, and warm and dry air masses from the Siberian mainland.

Samoylov Island is situated in the southern-central part of the Lena River Delta and is considered to be representative for the modern delta landscapes. It has a size of 7.5 km² and is composed of two geomorphological units. The western part (3.4 km²) represents a modern floodplain which is flooded annually during the spring flood. The investigation site was located on the eastern part of Samoylov (4.1 km²), which is composed of the sediments of the Late-Holocene river terrace. Its elevation ranges from 10 to 16 m a.s.l., and it is not flooded annually. Its surface is characterized by wet polygonal tundra. The macro-relief of the eastern part of Samoylov Island is level with slope gradients less than 0.2%. Only along the shorelines of the larger lakes, elevation differences of up to 2.5 m occur. However, the surface of the terrace is structured by a regular micro-relief with elevation differences of up to 0.5 m within a few meters, which is caused by the genesis of low-centred ice wedge polygons (Washburn, 1979; French, 1996; Meyer, 2003). The depressed centres of the ice-wedge polygons are surrounded by elevated rims, which are situated above the ice-wedges. The formation of low-centred polygons has a strong impact on the water regime and the ecological conditions of the tundra landscape. In the depressed polygon centres, drainage is impeded due to the underlying permafrost, and water-saturated soils or small ponds occur. In contrast, the elevated polygon rims are characterized by a moderately moist water regime. The typical soil types are Typic Historthels in the polygon centres and Glacic or Typic Aquiturbels at the polygon rims, respectively (Soil Survey Staff, 1998). The vegetation in the polygon centres and at the edge of ponds
part of Samoylov Island. Wet polygonal tundra extended at least 600 m in all directions from the eddy tower (Fig. 2). The three-dimensional wind vector and the sonic temperature were measured with a sonic anemometer (Solent R3, Gill Instruments Ltd., UK) at a height of 3.65 m above ground level. The concentration of H$_2$O and CO$_2$ were measured with a closed-path infrared gas analyser (LI-7000, LI-COR Inc., USA). The gas analyzer was installed in a temperature-regulated case at the foot of the tower. The sample air intake was placed 15 cm below the centre of the sonic anemometer transducer array. From the intake, sample air was drawn at a rate of 20 L min$^{-1}$ through a heated Polyethylene tube of 5 m length and 6.35 mm inner diameter to the gas analyzer. Under these conditions, turbulent flow was maintained inside the tubing system. A 1 $\mu$m membrane filter (PTFE, TE37, Schleicher & Schuell, Germany) prevented dust contamination. The analogous signals from the fast response sensors were digitized at a frequency of 20 Hz by the anemometer and logged by a laptop computer running the software EdiSol (J. Massheder, University of Edinburgh, UK). Power was supplied by a diesel generator which was placed at a distance of 100 m from the tower. Wooden boardwalks connected all parts of the system to reduce disturbance of the swampy tundra soils and the vegetation. All equipment was set up in a line to the west-southwest from the tower.

In addition to the fast-response eddy covariance measurements, a set of supporting slow-response meteorological measurements were conducted. The meteorological data were partly recorded at the eddy tower site and partly provided by a long-term meteorological station (Boike et al., 2007$^1$), which is located 700 m away from the eddy site in the polygonal tundra of Samoylov Island (Fig. 2). At the eddy tower were measured: air temperature and air relative humidity at 2 m height (MP103A, ROTRONIC AG, Switzerland), incoming and surface-reflected solar and infrared radiation (CNR1, Kipp & Zonen B.V., The Netherlands), and air pressure (RPT410, Druck Messtechnik GmbH, Germany). From the incoming solar radiation $S\downarrow$, the photosynthetically active radiation PAR was estimated using the equation

$$\text{PAR} = S\downarrow \cdot 0.45 \cdot 4.598 \mu\text{mol J}^{-1}\cdot \text{m}^{-2}.$$  (1)

This method follows Jacovides et al. (2003) and was shown to work very accurately by direct comparison of solar radiation and PAR sensors during a measurement campaign in the Lena River Delta in 2005 (data not shown). The long-term meteorological station recorded precipitation (tipping bucket rain gauge 52203, R.M. Young Company, USA), snow height at a polygon centre (sonic ranging sensor SR50, Campbell Scientific Inc., USA), horizontal wind speed and direction at 3 m height (propeller anemometer 05103, R.M. Young Company, USA), air temperature and relative humidity at 0.5 m and 2 m height (MP103A, ROTRONIC AG, Switzerland), net radiation (NR-Lite, Kipp & Zonen B.V., The Netherlands), and upward infrared radiation (CG1, Kipp & Zonen B.V., The Netherlands). From the upward infrared radiation.
$L \uparrow B$, the radiative surface temperature $T_{sur}$ was calculated using the formula

$$T_{sur} = \left( \frac{L \uparrow B}{\varepsilon \times 5.67 \times 10^{-8}} \right)^{\frac{1}{4}},$$

where the emissivity $\varepsilon$ was assumed to be 0.98. Profiles of soil temperature and soil volumetric water content were measured at the elevated rim and the depressed centre of a polygon (temperature probe 107 and TDR100/CS605, respectively, Campbell Scientific Inc., USA). Water level depth was measured manually in intervals of 1 to 3 days in perforated plastic pipes, which were installed in the soil active layer at three soil survey sites in the vicinity of the eddy tower. Thaw depth was measured in intervals of 3 to 7 days at 150 points arranged in a regular grid by driving a steel rod into the unfrozen soil until the hard frozen permafrost table was encountered (Kutzbach et al., 2004).

### 2.3 Processing of eddy covariance data

Turbulent fluxes were calculated over 30 min averaging intervals using the software EdiRe (R. Clement, University of Edinburgh, UK). Three coordinate rotations were applied to the wind vector components so that the mean transverse velocity, the mean vertical velocity, and the covariance of the transverse and vertical velocities were reduced to zero for each 30 min interval (McMillen, 1988; Aubinet et al., 2000). The time series of the CO$_2$ gas concentration was detrended using a recursive high-pass filter with a filter constant of 300 s (McMillen, 1988). The time lag between the signals of CO$_2$ concentration and vertical wind velocity was determined and removed for each 30 min interval. After the calculation of the CO$_2$ flux, transfer functions were applied to account for the spectral response of the gas analyzer, the separation of the sensors, the damping effect of the gas sampling tube and the detrending filter (Moore, 1986; Moncrieff et al., 1997). On average, the percentage of this correction compared to the uncorrected CO$_2$ flux was 8.5%. Additionally, the CO$_2$ flux was corrected for effects of concurrent water vapour fluxes on the density of air (Webb et al., 1980).

The calculated flux data was screened thoroughly. Data points were rejected when sensor outputs were out of range and when instruments were being repaired or calibrated. Turbulence was considered insufficient when the mean friction velocity was below 0.1 m s$^{-1}$. Data gathered during periods with wind directions in the sector 230°–270° were discarded because of the possible disturbance by the generator. Altogether, 31% and 26% of the calculated CO$_2$ fluxes were rejected in 2003 and 2004, respectively, which is comparable to other studies (Falge et al., 2001). The turbulent fluxes calculated over 30 min intervals were averaged over 60 min for compatibility with the meteorological data of the long-term meteorological station. The gaps in the data series produced by the screening procedure were filled by means of models based on empirical relationships between the turbulent fluxes and meteorological variables. The model approach is described below.

A footprint analysis following Schuepp et al. (1990) assessed the 80% cumulative footprint, i.e. the upwind distance from which 80% of the observed flux originated, to be on average 457 m during the snow-free periods, and 781 m during periods when snow covered the surface.

### 2.4 Modelling of carbon dioxide fluxes

The calculated carbon dioxide flux $F_{CO2}$ equals the net ecosystem CO$_2$ exchange NEE which is the sum of gross photosynthesis $P_{gross}$ and ecosystem respiration $R_{eco}$. The ecosystem respiration is composed of the respiration from soil microbes $R_{soil}$, roots $R_{roots}$, and above-ground biomass $R_{above}$ (Greco and Baldocchi, 1996):

$$F_{CO2} = NEE = P_{gross} + R_{eco}$$

$$= P_{gross} + (R_{soil} + R_{roots} + R_{above})$$

The quantities $F_{CO2}$, NEE, $R_{eco}$ and $P_{gross}$ indicate throughout this study fluxes of CO$_2$ (and not of carbon) and have the unit g h$^{-1}$ m$^{-2}$. This unit would equal in this context the unit g CO$_2$ h$^{-1}$ m$^{-2}$ which has often been used in other carbon flux publications. The ecosystem respiration is well known to depend on the temperature of the soils and the above-ground biomass. For the modelling of the ecosystem respiration, the time series of air temperature, radiative surface temperature, and soil temperature at various depths were studied with respect to their ability to describe the CO$_2$ fluxes observed during dark periods (PAR<20 µmol m$^{-2}$ s$^{-1}$), when photosynthesis was assumed to be negligible. The CO$_2$ flux data and temperature data was fitted using the exponential function

$$R_{eco} = p_1 e^{p_2 T_i},$$

where $T_i$ is air, surface or soil temperature, and $p_1$ and $p_2$ are the fit parameters. The best fit ($R^2=0.79, N=611$) was achieved when the radiative ground surface temperature $T_{sur}$ was used as independent variable (Fig. 6). Consequently, the relationship

$$R_{eco} = 0.0666 \text{ g m}^{-2} \text{h}^{-1} \exp \left(0.0785 \text{ °C}^{-1} T_{sur}\right)$$

was used for the estimation of the ecosystem respiration. The resulting model for night time $R_{eco}$ was extrapolated for the estimation of $R_{eco}$ during daytime. It was assumed that $R_{above}$ is of similar magnitude during day and night for the investigated tundra canopy. This simplification has been followed successfully by many other studies of carbon exchange of northern wetlands (Alm et al., 1997; Lloyd, 2001a; Harazono et al., 2003). Because there were no dark nights during the campaign in 2004, the model relies on 2003 data only. Ecosystem respiration in 2004 was modelled using the 2003 model, which showed to produce reasonable results.
Examples of the modelling of $P_{\text{gross}}$ and values of $\text{PAR}_n$–$\text{sat}$ are given in Fig. 3.

The NEE time series were gap-filled by combining the empirical models for $R_{\text{eco}}$ and $P_{\text{gross}}$.

3 Results

3.1 Meteorological conditions

In 2003, measurements started on 19 July during the final phase of the polar day and lasted until 21 October, when the sun rose for only seven hours above the horizon (Fig. 4). Generally, the year 2003 was characterized by comparatively high temperatures; the average temperatures of July (11°C) and September (3°C) were 4 K and 2 K higher than the long-term averages. The months July, August, and September experienced extended periods of high air temperatures which were caused by advection of warm air from the Siberian mainland. At 168 mm, the precipitation (rain) during the measurement period was exceptionally high. A large part of the rainfall was recorded during one week at the end of July (94 mm in six days). The precipitation pattern was reflected by the water table levels: Maximum water table heights were observed with 8 cm above soil surface in the polygon center and 8 cm below the soil surface at the polygon rim on 25 July, when the strongest rainfall event occurred. Afterwards, the water table heights decreased gradually and reached their minimum at the end of September with 0 cm above soil surface in the centre and 13 cm below the soil surface at the rim. The volumetric water content in the top soil (5 cm depth) in the centre and at the rim of the polygon was on average 97% (saturation: 98%) and 30% (saturation: 47%) during the period of unfrozen soils. During the course of the measurement campaign, the thaw depth increased from 28 cm to a maximum of 48 cm on 12 September. The soil temperatures followed the variations of the air temperature; however, the fluctuations were damped and phase-lagged dependent on depth below the soil surface and the position within the micro-relief. A strong temperature gradient existed in the unfrozen soil layer during the summer. Due to the relatively high temperatures in September, freezing of the soils from the top began late at around 29 September. At the end of the campaign on 21 October, the top soil layers were frozen, but underneath the frozen layer, unfrozen soil zones of several decimetres depth persisted. A snow cover built up starting 9 October. At the end of the campaign, the snow cover reached a thickness of about 15–25 cm in the polygon centres and just a few centimetres at the rims.

In 2004, the entire campaign was conducted under polar day conditions (Fig. 4). The year 2004 was characterized by a very cold winter, comparatively low temperatures during spring, and a late start of the growing season. The average temperatures of April (−20°C), May (−8°C) and June (1°C) were 2 K, 1 K and 1 K lower compared to the long-term average.
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Fig. 4. Meteorological and soil data during the measurements campaigns 2003 and 2004. (A) air temperature at 2 m height, (B) daily sum of rainfall, (C) height of water table above soil surface at the centre (squares) and the rim (triangles) of a polygon, depth of frozen ground table (circles), and snow height (line without symbol), (D) soil volumetric water content at 5 cm depth at the centre (black) and the rim (grey) of a polygon, (E) soil temperature at the polygon centre at depths of 1 cm (black line), 10 cm (grey line), 20 cm (dotted line), and 30 cm (dashed line).

means. At the beginning of the measurement campaign, the ground around the eddy tower was completely covered with snow and the soil temperatures were below −10°C throughout the soil profile. The snow height was about 40 cm in the polygon centre and about 8 cm at the polygon rim. Before snow melt, sublimation of snow occurred throughout and snow-free patches appeared at the polygon rims on 29 May. However, frequent snowfall and snow drift kept the area size of the snow free patches small. The snow melt started on 8 June. The snow height decreased rapidly, and the polygon rims were largely free of snow after two days. Snow melt in the polygon centres continued until 18 June. The soils started to thaw around 13 June. The highest temperature in the upper soil layers was measured on 9 July, when the thaw depth was about 20 cm. The thaw depth increased nearly linearly and reached a value of 26 cm at the end of the campaign. The total rainfall up to 21 July was 60 mm. A large part of the rain fell on 8–9 July. The water table was highest during snow melt (15 June) with 12 cm and 0 cm above the soil surface in the centre and at the rim of the polygon, respectively. Afterwards, the water table fell quickly, but due to the strong rainfall events of 8–9 July stayed close below or above the soil surface in the polygon centre until the end of the campaign. After thawing of the top 10 cm of the soils, the volumetric liquid water content in the top soil layer was on average 97% and 31% at the polygon centre and rim, respectively.

The average wind speed during the observation periods in 2003 and 2004 was 4.7 m s⁻¹. Very light winds occurred seldom with wind speeds <1 m s⁻¹ observed less than 2% of the observation time. During the campaign 2003, there was no single predominant wind direction, however, wind directions east-northeast, south, and south-west occurred more
Fig. 5. CO₂ budget components and meteorological controls in 2003 and 2004. (A) NEE measured by the eddy covariance method, (B) modelled ecosystem respiration $R_{eco}$, (C) modelled gross photosynthesis $P_{gross}$ (black) and gross photosynthesis near light saturation $P_{n-sat}$ (grey), (D) photosynthetically active radiation PAR; (E) surface radiative temperature $T_{sur}$. NEE, $P_{gross}$ and $R_{eco}$ are given as CO₂ fluxes.

frequently than other directions (data not shown). During the campaign 2004, there was a clear dominance of easterly winds (about 23% of the measurement period, data not shown), followed by winds from north-westerly directions. Wind directions not acceptable for flux calculations due to disturbance by the generator $230°$–$270°$) occurred 13.5% and 5.6% of the observation time in 2003 and 2004, respectively.

3.2 Carbon dioxide fluxes

During most of the measurement campaign 2003, NEE oscillated regularly between net uptake of CO₂ during daytime and net CO₂ release during night (Fig. 5). Between mid-July and mid-August, the amplitude of the diurnal oscillation was greatest, and daytime CO₂ uptake dominated. Afterwards, the amplitude decreased, and the CO₂ release during night time gained more importance relative to the daytime CO₂ uptake. After 29 September, uptake of CO₂ ceased and NEE was continuously positive. At the end of the campaign at 21 October, CO₂ release was still substantial with about 0.013 g h⁻¹ m⁻². The NEE time series was markedly affected by the synoptical weather pattern. For instance, during the strong rainfall on 25 July 2003 daytime CO₂ uptake reached only $−0.17$ g h⁻¹ m⁻² whereas it rose to more than $−0.50$ g h⁻¹ m⁻² on the partly cloudy 26 July. Also, advection events of warm air from the south were clearly visible as shifts to higher nightly CO₂ release, e.g. during 4–8 August and 2–7 September 2003.

The modelled ecosystem respiration ranged between $0.01$ g h⁻¹ m⁻² and $0.55$ g h⁻¹ m⁻² during the campaign.
The largest \( R_{\text{eco}} \) values and its largest diurnal amplitude were encountered in the first week of August, when the surface temperature was at its maximum during a pronounced event of advection of warm air from the south. The mean \( R_{\text{eco}} \) during 4–8 August was 0.23 g h\(^{-1}\) m\(^{-2}\). Afterwards, \( R_{\text{eco}} \) decreased; it averaged to 0.11 g h\(^{-1}\) m\(^{-2}\) during the second half of August, to 0.09 g h\(^{-1}\) m\(^{-2}\) during September and to 0.03 g h\(^{-1}\) m\(^{-2}\) during October.

The modelled gross photosynthesis ranged between zero and 0.70 g h\(^{-1}\) m\(^{-2}\) during the campaign 2003. Following the daily trend of PAR, the amplitude of the diurnal oscillation of \( P_{\text{gross}} \) was large. From the beginning of measurements until about 10 August, photosynthesis took place for 24 h per day, however, with only small absolute values during the night hours. The largest values of \( P_{\text{gross}} \) were observed from mid-July to the first week of August. The mean \( P_{\text{gross}} \) during this period was approximately 0.26 g h\(^{-1}\) m\(^{-2}\). Afterwards, the \( P_{\text{gross}} \) decreased; it averaged to 0.17 g h\(^{-1}\) m\(^{-2}\) during the second half of August, and to 0.07 g h\(^{-1}\) m\(^{-2}\) during September. During October, \( P_{\text{gross}} \) was zero. During the midday maxima of PAR, \( P_{\text{gross}} \) frequently reached \( P_{\text{n-sat}} \), indicating that photosynthesis was limited by the saturation of the photosynthetically active tissue of the tundra canopy. Only on very cloudy days, midday \( P_{\text{gross}} \) was limited by irradiation.

The pattern of the NEE time series of the campaign 2004 (Fig. 5) reflects the drastic changes of the physical conditions and the developmental stages of organisms during this period. Already at the last days of May, when the first snow-free patches appeared at some polygon rims, a weak diurnal oscillation of NEE between night time CO\(_2\) emission and daytime CO\(_2\) uptake was observed. NEE was around +0.003 g h\(^{-1}\) m\(^{-2}\) during the night of 28/29 May and about −0.002 g h\(^{-1}\) m\(^{-2}\) during midday of 29 May. The amplitude of this oscillation increased slowly during the first week of June. During the snow melt (8–18 June), the uptake of CO\(_2\) increased strongly and clearly dominated the NEE pattern. Afterwards, the midday peak values of CO\(_2\) uptake fluctuated substantially on the meso-scale but increased in general with NEE reaching a maximum of about −0.25 g h\(^{-1}\) m\(^{-2}\) on 19 July. The night time CO\(_2\) emissions varied between 0.02 g h\(^{-1}\) m\(^{-2}\) and 0.15 g h\(^{-1}\) m\(^{-2}\) (mean 0.06 g h\(^{-1}\) m\(^{-2}\)) and showed no clear trend.

Ecosystem respiration and gross photosynthesis were modelled beginning 2 June. Before this date, the models did not appropriately reflect the small-scale dynamics of the CO\(_2\) fluxes. During the period 2–20 June, the temperature of the top soil layers did not exceed 0°C and \( R_{\text{eco}} \) was low with little diurnal variation. Afterwards, \( R_{\text{eco}} \) increased strongly but also strongly depended on the synoptic weather pattern. The maximum of \( R_{\text{eco}} \) was modelled for 8 July (0.57 g h\(^{-1}\) m\(^{-2}\)), when air, surface and soils temperatures were high during the advection of warm air from the South. \( R_{\text{eco}} \) was markedly lower during the advection of cold air from the North (e.g. 28 June–5 July). \( R_{\text{eco}} \) averaged to 0.05 g h\(^{-1}\) m\(^{-2}\) and 0.12 g h\(^{-1}\) m\(^{-2}\) during the first and second half of June, respectively, and to 0.16 g h\(^{-1}\) m\(^{-2}\) during July.

Before the beginning of snow melt on 8 June, gross photosynthesis was low with midday peak values of about −0.05 g h\(^{-1}\) m\(^{-2}\). During this period, only few snow-free patches existed at some polygon rims. During the period of the snow melt (8–18 June), when more and more vegetation was released from the snow cover, \( P_{\text{gross}} \) increased strongly to midday peak values of −0.2 g h\(^{-1}\) m\(^{-2}\). During the snow-free period, \( P_{\text{gross}} \) increased further and reached midday peak values around −0.4 g h\(^{-1}\) m\(^{-2}\) in July. More distinctly than in 2003, the photosynthesis was limited by the canopy photosynthetic potential. Especially in the first phase of the campaign, \( P_{\text{gross}} \) frequently exceeded \( P_{\text{n-sat}} \) showing a high degree of light saturation of the canopy. With the development of the plants and increasing temperatures towards the end of June, \( P_{\text{n-sat}} \) increased so that the limitation of \( P_{\text{gross}} \) by the canopy photosynthetic potential lost importance.

3.3 The regulation of respiration and photosynthesis

The study of the relationship between ecosystem respiration \( R_{\text{eco}} \) and various temperatures revealed a close exponential relationship between \( R_{\text{eco}} \) and the air temperature at 0.5 m above ground as well as the radiative ground surface temperature (Fig. 6a, b). However, the relationships between \( R_{\text{eco}} \) and the soil temperatures at 1 cm depth at a polygon rim and centre site were less clear (Fig. 6c, d). The low performance of the exponential fits of \( R_{\text{eco}} \) and soil temperature data was primarily caused by the different trends of the data series during the refreeze of soils (Figs. 4 and 5). During this time, due to the release of large amounts of latent heat, the soil temperatures remained for a long time at 0°C ("zero curtain effect"), whereas \( R_{\text{eco}} \) decreased steadily.

The characteristic parameters of the tundra canopy photosynthetic activity as derived by the fit of \( P_{\text{gross}} \) and PAR data using Eq. (7) are displayed for the synthetic measurement period 2004/2003 in Fig. 7. The canopy photosynthetic potential \( P_{\text{max}} \) generally followed the seasonal progression of the air temperature, reflecting the combined effect of the phenological development of the vegetation and the positive forcing of photosynthesis by temperature. When the first snow-free patches at the polygon rims appeared at the end of May, \( P_{\text{max}} \) was low but detectable with −0.07 g h\(^{-1}\) m\(^{-2}\). After 8 June, when the daily average air temperature rose above 0°C and the snow melt started, \( P_{\text{max}} \) increased steadily until mid-July. The period of maximum \( P_{\text{max}} \) lasted approximately three to four weeks from mid-July to the first week of August with \( P_{\text{max}} \) values around −0.8 g h\(^{-1}\) m\(^{-2}\) (range −0.5 g h\(^{-1}\) m\(^{-2}\)–1.1 g h\(^{-1}\) m\(^{-2}\)). This period indicates the mature phase of vegetation. As senescence started during August, \( P_{\text{max}} \) decreased at a high rate, later in September the decrease continued at lower rates. \( P_{\text{max}} \) decreased to zero at the end of September, when the daily average air
temperature fell below zero. The substantial scatter of $P_{\text{max}}$ around the general trend especially during midsummer was at least partly related to the precipitation pattern and subsequent moisture changes in the top soils and the moss layer. Particularly high $P_{\text{max}}$ values were observed during the periods of heavy rain at the end of July.

The initial canopy quantum efficiency $a$ increased similarly as $P_{\text{max}}$, starting at the appearance of the first snow-free patches and leading to the mature phase of the vegetation, but peaked delayed compared to $P_{\text{max}}$ during the second to third week of August. During autumn, $a$ decreased but still showed strong variations.

### 3.4 Seasonal and annual CO$_2$ balance

The daily integrated components of the CO$_2$ budget, $R_{\text{eco}}$, $P_{\text{gross}}$ and NEE are shown for the synthetic measurement period 29 May–21 October 2004/2003 in Fig. 8. At the beginning of June, $R_{\text{eco}}$ and $P_{\text{gross}}$ were low and of similar magnitude, and hence resulted in very low daily NEE. From 7 June to 9 July, the daily NEE alternated between net CO$_2$ uptake and net CO$_2$ release. This was related to the synoptic weather conditions and the dynamics of the underlying biological and soil-physical processes, i.e. the vegetation development and the thawing of the permafrost soils. From 10 July to 31 August, daily NEE was continuously negative and averaged to $-2.1$ g d$^{-1}$ m$^{-2}$. The daily CO$_2$ uptake reached a maximum of $-4.6$ g d$^{-1}$ m$^{-2}$ at the end of July. After the beginning of September, daily NEE was positive on most days except during the cold period 10–15 September. From mid-September to mid-October, the daily CO$_2$ release was on average $0.7$ g d$^{-1}$ m$^{-2}$; it decreased to the end of the campaign to about $0.3$ g d$^{-1}$ m$^{-2}$.

The sums of $P_{\text{gross}}$, $R_{\text{eco}}$ and NEE over the whole measurement campaign amounted to $-432$ g m$^{-2}$, $+344$ g m$^{-2}$ and $-90$ g m$^{-2}$, respectively. From 11 June to 31 August, the polygonal tundra was a net CO$_2$ sink. During this period, $P_{\text{gross}}$, $R_{\text{eco}}$ and NEE summed to $-375$ g m$^{-2}$, $+251$ g m$^{-2}$ and $-124$ g m$^{-2}$, respectively. From the beginning of
September, the polygonal tundra was a substantial CO$_2$ source.

An estimate of the annual CO$_2$ budget was calculated by combining the gap-filled NEE time series of 2003 and 2004 with estimated values of $R_{\text{eco}}$ for the winter and spring period 22 October 2003–28 May 2004 (220 days). The empirical models of $R_{\text{eco}}$ as a function of air, soil, and radiative ground surface temperature (Fig. 6) could not be used for the modelling of the wintertime $R_{\text{eco}}$. These models did not deliver sensible results when compared to measured CO$_2$ fluxes during the first days of measurements in 2004. We assumed that air and radiative ground surface temperature did not reflect the temperature regime in the soil, where respiration took place, and that the models based on soil temperature data were biased by the presence of a still unfrozen soil layer in early winter. Hence, this study followed a more simple but conservative approach for estimating $R_{\text{eco}}$ during winter and spring (Fig. 9). The modelling period was divided in two periods which were separated by the date when the soils of the polygonal tundra were completely frozen (17 November 2003, derived from soil temperature records). During the period 17 November–28 May, $R_{\text{eco}}$ was set constant to the value 0.003 g h$^{-1}$ m$^{-2}$, which is the value observed during the night 28/29 May 2004, when the soils were still completely frozen. This value falls at the lower end of the range of mean winter CO$_2$ fluxes reported by other authors from similar tundra sites (Fahnestock et al., 1998; Oechel et al., 1997a; Panikov and Dedysh, 2000; Zimov et al., 1993; Table 1). The resulting estimate of NEE during this period (194 days) was $+14$ g m$^{-2}$. During the period 22 October–17 November, $R_{\text{eco}}$ was estimated by linear interpolation between the value of $R_{\text{eco}}$ measured at 21 October 2003 (0.013 g h$^{-1}$ m$^{-2}$) and the value of $R_{\text{eco}}$ of the second period (0.003 g h$^{-1}$ m$^{-2}$). The resulting estimate of NEE during this period (26 days) was $+5$ g h$^{-1}$ m$^{-2}$. Hence, the cumulative NEE for the one-year period 20 July 2003–19 July 2004 was $−71$ g m$^{-2}$, and the polygonal tundra of the study area was a CO$_2$ sink during this period. Approximately 363 g CO$_2$ m$^{-2}$ was respired by the plants and the soil organisms, which was about 84% of the annual CO$_2$ assimilation by the vegetation of 432 g CO$_2$ m$^{-2}$.

4 Discussion

4.1 Gross photosynthesis

4.1.1 Comparison with other tundra sites

The integrated value of $P_{\text{gross}}$ of the wet polygonal tundra in the Lena River Delta for the period June–August was $−383$ g m$^{-2}$. This value was low compared to $P_{\text{gross}}$ estimates for other arctic tundra sites. Estimates of $P_{\text{gross}}$ for the same seasonal period for wet sedge tundra and moist tussock tundra on the North Slope of Alaska were $−519$ g m$^{-2}$ and $−858$ g m$^{-2}$, respectively (Vourlites et al., 2000). The lower value observed in this study is partly related to later snow melt but also to generally lower $P_{\text{gross}}$ values during midsummer. The peak $P_{\text{gross}}$ value at the wet sedge tundra site described by Vourlites et al. (2000) was $−13.2$ g d$^{-1}$ m$^{-2}$ whereas the peak value at the Lena River Delta was only $−8.6$ g d$^{-1}$ m$^{-2}$. At mixed moist to wet tundra on the
Most tundra soils are characterized by a low nutrient availability due to the cold and often waterlogged soil conditions, which slow down microbial and soil fauna activity and consequently decomposition and mineralization of organic matter (Chapin et al., 1980a; Gersper et al., 1980; Shaver et al., 1998; Johnson et al., 2000; Hobbie et al., 2002). Furthermore, the soils of the polygonal tundra at the investigation site have to be considered as extremely poor, even when compared to other tundra soils. Water-logging is prominent due to the flat macro-relief and the development of low-centred polygons. The growing season is short, and the soils are extremely cold also during summer due to the very cold permafrost of Northern Siberia. The parent material of the soils consists mainly of fluvial, nutrient-poor sands of the Holocene river terrace. Moreover, the Holocene river terrace on which the polygonal tundra has formed is not flooded regularly anymore during the spring flood, so that no fresh nutrients are transported to the soils.

The soils at wet sedge tundra sites for which much higher gross photosynthesis was reported in the literature compared to this study (see above) are considered less nutrient-limited than the soils of the investigation site of this study as they were situated at the bottom of mountain valleys (Vourlites et al., 2000; Laurila et al., 2001) or on fine-grained marine sediments at coastal tundra (Brown et al., 1980; Harazono et al., 2003). On moist tundra sites, e.g. tussock tundra or shrub tundra, $P_{\text{gross}}$ appears to be generally larger than at wet sedge tundra sites due to more favourable conditions for the growth of vascular plants (Vourlites and Oechel, 1999; Vourlites et al., 2000; McFadden et al., 2003; Zamolodchikov et al., 2003).

4.1.2 Control by irradiation

Arctic tundra ecosystems have often been described as light-limited ecological systems (Tieszen et al., 1980). However, the results of this study showed that during the day $P_{\text{gross}}$ was often limited by saturation of the photosynthetic apparatus of the canopy. Although the canopy never reached full light saturation, the quantum efficiency declined substantially during midday high PAR intensities. A similar response of $P_{\text{gross}}$ to the diurnal PAR trend was described for wet sedge and tussock tundra in Alaska by Vourlites and Oechel (1997, 1999). The leaves of vascular tundra plant species approach light saturation at PAR values of $1300–1600 \mu\text{mol} \text{s}^{-1} \text{m}^{-2}$ (Tieszen et al., 1980), a value that was not reached at the investigation site during the growing season. However, arctic mosses tend to reach light saturation already at PAR values of around $450 \mu\text{mol} \text{s}^{-1} \text{m}^{-2}$, which is exceeded on average from 07:00 to 19:00 during June and July at the Lena River Delta. The observed limitation of $P_{\text{gross}}$ by the amount of photosynthetically active tissue in the tundra canopy under the current climate support the notion that enhanced plant

![Diagram showing cumulative net ecosystem exchange ΣNEE from July 2003 to July 2004. The periods of the micrometeorological measurements are highlighted by the grey background. The stars indicate: A=start of measurements 2003, B=start of modelling period with linearly decreasing $R_{\text{eco}}$, C=start of modelling period with constant $R_{\text{eco}}$ of 0.003 g h$^{-1}$ m$^{-2}$, D=start of measurements 2004, E=end of measurements 2004. ΣNEE is given as a cumulative CO$_2$ flux.](image-url)
Table 1. Comparison of cumulative net ecosystem CO\(_2\) exchange (\(\Sigma\)NEE) reported by other investigators and by this study. Note that the comparison periods differ depending on the availability of data. \(\Sigma\)NEE was calculated by integrating the CO\(_2\) fluxes over the comparison period specified.

<table>
<thead>
<tr>
<th>Reference</th>
<th>Location</th>
<th>Tundra type</th>
<th>Comparison period</th>
<th>(\Sigma)NEE ((\text{g m}^{-2}))</th>
<th>(\Sigma)NEE this study ((\text{g m}^{-2}))</th>
</tr>
</thead>
<tbody>
<tr>
<td>Nordstroem et al. (2001)</td>
<td>Zackenberg, Greenland 74° N, 20° W</td>
<td>wet fen and grassland</td>
<td>Jun...Aug 1997</td>
<td>−120</td>
<td>−119</td>
</tr>
<tr>
<td>Groendahl et al. (2007)</td>
<td>Zackenberg, Greenland 74° N, 20° W</td>
<td>high Arctic heath</td>
<td>DOY 159...238</td>
<td>−5</td>
<td>−117</td>
</tr>
<tr>
<td>Oechel et al. (1993)</td>
<td>Prudhoe Bay, Alaska 70° N, 148° W</td>
<td>wet sedge</td>
<td>June...Sep (125 d) 1990</td>
<td>16</td>
<td>−116</td>
</tr>
<tr>
<td>Harazono et al. (2003)</td>
<td>Barrow, Alaska 71° N; 156° W</td>
<td>wet sedge</td>
<td>May...Sep 1999; 2000</td>
<td>−593</td>
<td>−104</td>
</tr>
<tr>
<td>Zamolodchikov and Karelin (2001)</td>
<td>whole Russian tundra</td>
<td>all types</td>
<td>June...Sep (117 d) model</td>
<td>−103</td>
<td>−121</td>
</tr>
<tr>
<td>Corradi et al. (2005)</td>
<td>Cherskii, Siberia 69° N, 162° E</td>
<td>moist to wet tussock</td>
<td>July...Aug 2002</td>
<td>−100</td>
<td>−112</td>
</tr>
<tr>
<td>Corradi et al. (2005)</td>
<td>Cherskii, Siberia 69° N, 162° E</td>
<td>moist to wet tussock</td>
<td>Sep...June 2002/2003</td>
<td>62</td>
<td>40</td>
</tr>
<tr>
<td>Fahnestock et al. (1998)</td>
<td>Prudhoe Bay, Alaska 70° N, 148° W</td>
<td>wet sedge</td>
<td>Feb...May 1996</td>
<td>1.84</td>
<td>9</td>
</tr>
</tbody>
</table>
growth under a warming climate may increase $P_{\text{gross}}$ and thus CO$_2$ uptake by arctic canopies provided that nutrient availability will increase in parallel (Shaver et al., 1992, 1998; Oechel et al., 2000; Hobbie et al., 2002).

4.1.3 Control by phenological development

The seasonal variation of $P_{\text{gross}}$ was mainly controlled by the phenological development of the canopy and the production of photosynthetically active leave area, which in turn was related to the general temperature progression over the growing season. Photosynthesis started directly when the first snow-free patches appeared on the polygon rims at the end of May. Most of this early photosynthesis was probably accomplished by mosses, whose photosynthetically active tissues can overwinter and start to assimilate CO$_2$ as soon as it is freed from the snow cover (Oechel, 1976). Also the vascular plants start to grow and photosynthesize within one day of snow melt (Tieszen et al., 1980). However, their photosynthetic rate is at first very low since their photosynthetically active tissues are not maintained over winter and have to be produced newly. From the start of photosynthesis, $P_{\text{gross}}$ increased until the mature stage of the vascular plants was reached at mid-July. Senescence of the vascular plants started after the first week of August and $P_{\text{max}}$ decreased as proteins and other materials were hydrolyzed and mobilized. In contrast, the mosses showed no senescence and continued to photosynthesize until the air temperature dropped below zero (Fig. 7). The shoulders of the seasonal progression curve of $P_{\text{max}}$ (28 May–10 June and 11–29 September) are considered to indicate periods of only moss photosynthesis. By interpolating between these dates, the contribution of moss photosynthesis to the total $P_{\text{gross}}$ during the photosynthetically active period was estimated to be approximately 40%. Miller et al. (1980) found by harvesting methods that mosses contributed about 30% to the gross primary production of a similar wet tundra canopy.

The non-synchronous seasonal trends of $P_{\text{max}}$ and $\alpha$ is probably related to the different seasonal development of vascular plants and mosses. The coefficients of the light response model of $P_{\text{gross}}$ have a firm physiological basis: $P_{\text{max}}$ is indicative of the development and capacity of the CO$_2$ fixation apparatus, such as the amount and activity of the carboxylation enzyme Rubisco, whereas $\alpha$ is determined by the leaf chlorophyll content and photosynthetically active leaf area (Vourlites and Oechel, 1999). The moss layer can be considered to have a large photosynthetically active leaf area (high $\alpha$) but a markedly lower content of Rubisco than vascular plants (low $P_{\text{max}}$) (Miller et al., 1980). Thus, senescence of the vascular plants affected $P_{\text{max}}$ stronger and earlier than $\alpha$.

4.1.4 Control by temperature

Temperature is generally a major control factor on $P_{\text{gross}}$. While the light reactions of photosynthesis are insensitive to temperature, the dark reactions are highly sensitive to temperature since the activity of Rubisco has a distinct temperature optimum. However, the temperature optima of the photosynthesis of arctic plants are broad. The photosynthesis optima for vascular plants and bryophytes were found to be 10–15°C and 10–19°C, respectively (Oechel, 1976; Tieszen et al., 1980). Even at 0°C, photosynthetic activity of Arctic plant species is substantial. The lower limit of photosynthesis was shown to be about −4°C (Tieszen et al., 1980).

Despite the adaptation of Arctic vegetation to the low temperatures in the Arctic, plant growth and consequently $P_{\text{gross}}$ is severely constrained by the low temperatures and the short growing season. The timing of the snow melt has a great influence on the length of the growing season and the annual $P_{\text{gross}}$. Variations in growing season length probably accounted for much of the large interannual variation in $P_{\text{gross}}$ observed by several studies in the Arctic (Vourlites and Oechel, 1997; Lloyd, 2001b; Harazono et al., 2003). Atmospheric warming may prolong the summer period free of frost and snow and permit a longer growing season, increased plant growth and higher CO$_2$ assimilation of both vascular plants and mosses (Sveinbjörnsson and Sonesson, 1997; Oechel et al., 1998). When projecting future changes of CO$_2$ assimilation due to climate change, it has to be kept in mind that phenological events such as bud break, culm elongation and flower development in spring and summer as well as senescence and bud dormancy initiation in autumn are controlled by a complex suite of environmental and plant-internal variables including not only temperature but also photoperiod or species internal periodicity (Heide, 1990, 1997, 2005; Oberbauer et al., 1998; Keller and Köper, 2003). In the Arctic, however, the timing of snowmelt and soil thaw has to be considered as the key factor in determining the start of the growing season (Pop et al., 2000; Chapin et al., 2005). The photoperiod would only limit significantly the plant development in spring if the snowmelt would be drastically earlier (for example April) because the timing of snowmelt in the Arctic under the actual climate occurs during all-day-light conditions near the summer solstice. On the other hand, the triggering of the end of the growing season is less well understood and probably the result of several interacting factors (Oberbauer et al., 1998; Van Wijk et al., 2003). In the long-term, an extension of the growing season will favour species which are able to adapt to and benefit from a longer growing season enhancing thus vegetation community changes in the Arctic (Chapin et al., 2005).

4.1.5 Control by water availability

Water availability is considered to be of minor importance for the regulation of whole canopy gross photosynthesis at the wet polygonal tundra. As neither a depression of $P_{\text{gross}}$ nor a decrease of evapotranspiration at midday by stomatal regulation was observed, the vascular plants were considered to not experience appreciable water stress. However,
the meso-scale variations around the general trend of $P_{\text{max}}$ (Fig. 7) could in part be attributed to the response of mosses to water availability. Mosses are very sensitive to changes in available water since they cannot control their tissue water content (Miller et al., 1980). In the periods between precipitation or dew events, the mosses at the moist polygon rims desiccated, particularly during warm and dry weather (Lloyd, 2001a). The desiccation of mosses led to a decrease of the moss photosynthetic potential which consequently also reduced the whole canopy photosynthetic potential $P_{\text{max}}$. During events of rain, the tissue water content of the mosses was replenished and moss photosynthetic potential resumed quickly (Sveinbjörnsson and Sonesson, 1997). These processes were clearly observable from the middle of July to the beginning of August, when periods of high precipitation alternated with dry and warm periods and the calculated $P_{\text{max}}$ fluctuated substantially on the scale of several days. An alternative or complementing explanation for the increased $P_{\text{max}}$ during rainy periods could be that during cloudy weather the diffuse radiation is higher, which is more effective in feeding photosynthesis than direct sunlight (Roderick et al., 2001; Stanhill and Cohen, 2001; Gu et al., 2003; Rocha et al., 2004).

4.2 Ecosystem respiration

4.2.1 Comparison with other tundra sites

The average $R_{\text{eco}}$ observed at the polygonal tundra of the Lena River Delta during summer and the autumnal period of refreezing was comparatively low. It amounted to 60% of average $R_{\text{eco}}$ observed at a moist to wet tussock tundra at the Kolyma River lowlands in North-East Siberia (Corradi et al., 2005), to 60% of $R_{\text{eco}}$ at moist to wet tundra on the Chukotskiy Peninsula (Zamolodchikov et al., 2003), to 47–54% of $R_{\text{eco}}$ at a high arctic fen (Soegaard and Nordstroem, 1999; Nordstroem et al., 2001), to 38–44% of $R_{\text{eco}}$ at tussock tundra at an Alaskan mountain valley (Vourlites and Oechel, 1999; Vourlites et al., 2000) and to 50% of the average $R_{\text{eco}}$ modelled for the whole Russian tundra area (Zamolodchikov and Karelin, 2001). On the other hand, $R_{\text{eco}}$ observed in this study was equal to $R_{\text{eco}}$ at wet sedge tundra at an Alaskan mountain valley (Vourlites et al., 2000), about two times higher than $R_{\text{eco}}$ reported for flooded wet sedge tundra at the coastal plain of Alaska (Harazono et al., 2003) and about 2.3 times higher than $R_{\text{eco}}$ at an high-arctic semi-desert at Svalbard (Lloyd, 2001a).

Major controls on $R_{\text{eco}}$ are temperature, soil moisture, water table position, soil redox conditions, nutrient availability, vegetation type and litter quality (Hobbie et al., 2002; Ping et al., 1997; Christensen et al., 1998; Zamolodchikov and Karelin, 2001). Most of these control factors are not favourable for a high $R_{\text{eco}}$ at the wet polygonal tundra of the Lena River Delta: Soil temperatures are low due to the very cold permafrost in the region. Widespread water-logged conditions cause anaerobic soil conditions, and nutrient availability is poor as described before. Furthermore, the vegetation is characterized by a low coverage of vascular plants and a high coverage of mosses, which are known to produce extremely recalcitrant litter and even bactericidal substances in their tissues (Zimov et al., 1993; Hobbie et al., 2002).

The comparison of values of $R_{\text{eco}}$ and $P_{\text{gross}}$ given by other studies for tundra ecosystems reveals that $R_{\text{eco}}$ is generally higher in ecosystems with higher $P_{\text{gross}}$, with the exception of permanently flooded tundra types which showed relatively low $R_{\text{eco}}$ and high $P_{\text{gross}}$ (Harazono et al., 2003). More CO$_2$ is assimilated by the vegetation in tussock tundra (Vourlites and Oechel, 1999; Vourlites et al., 2000; Corradi et al., 2005) than in wet sedge tundra (Vourlites and Oechel, 1997; this study), but also more CO$_2$ is released by the respiratory processes. At arctic semi-desert sites, the rates of both $R_{\text{eco}}$ and $P_{\text{gross}}$ are much lower compared to all real tundra sites (Lloyd 2001a, b). Regional comparison studies on Greenland and arctic Alaska found that $R_{\text{eco}}$ increased with the leaf area index LAI. This was explained by a combination of greater leaf maintenance respiration increasing with LAI and increased soil respiration due to better litter quality and larger root biomass at higher productivity sites (Soegaard et al., 2000; McFadden et al., 2003).

4.2.2 Control by temperature

Most of the variation of $R_{\text{eco}}$ during summer and autumn could be well modelled by an exponential function between $R_{\text{eco}}$ and the surface temperature ($R^2=0.79$ for the 2003 campaign). The good performance of the $R_{\text{eco}}$ model based on surface temperature can be explained by the importance of the above-ground plant respiration. Although most of the biomass in wet sedge-dominated tundra ecosystems (80–88%) is belowground in roots and rhizomes of the grasses and sedges (Billings et al., 1977; Chapin et al., 1980b), 30–46% of summer $R_{\text{eco}}$ in grass and sedge tundra was found to originate from above-ground sources (Peterson and Billings, 1975; Nordstroem et al., 2001; Zamolodchikov and Karelin, 2001). This high respiration in the relatively small aboveground biomass reflects the intense biological activity inside the arctic plants during the short growing season. Within only three months, the vascular tundra plants have to develop their complete photosynthetically active tissue, to flower and to senesce. All these processes imply intense allocation of nutrients and carbohydrates, which is powered by autotrophic respiration. Furthermore, soil and root respiration was shown to be most prominent in the uppermost centimetres of tundra soils (Billings et al., 1977; Sommerkorn, 1998). However, heterotrophic respiration can not be assumed to have a significant contribution to the above-ground respiration, because there is not much litter above the ground surface, which would mean on the dense moss layer or between the living moss stems. An additional explanation for the poor fits of $R_{\text{eco}}$ to the soil temperatures compared to the
surface temperature is that the soil respiration probably declined gradually during the refreeze period in response to the continuously low temperatures while the temperatures itself stayed stable due to the “zero curtain” effect.

4.2.3 Seasonal progression of $R_{\text{eco}}$

At the polygonal tundra of the Lena River Delta, 70% of the estimated annual $R_{\text{eco}}$ took place during the summer months June–August. This value is well within the range of 60–80% estimated by Coyne and Kelley (1975). However, also the autumn was found to be important for the dynamics of $R_{\text{eco}}$ at the investigation site. About 20% of the estimated annual $R_{\text{eco}}$ was achieved in September. During the freezing of the soils from October to mid-November, $R_{\text{eco}}$ amounted to 6% of the annual ecosystem respiration. The so-called “period of autumnal carbon emission” (Zamolodchikov and Karelin, 2001) is a common feature of high-latitude ecosystems and is one of the causes of the high atmospheric $CO_2$ concentration during winter which was observed over northern Siberia (Fung et al., 1987; Zimov et al., 1993; Zamolodchikov and Karelin, 2001).

On the other hand, no pronounced period of $CO_2$ net emission during spring snow melt was observed as reported for a range of tundra sites (Vourlites and Oechel, 1997; Soegaard et al., 2000; Vourlites et al., 2000; Nordstroem et al., 2001; Zamolodchikov and Karelin, 2001; Corradi et al., 2005). During the snow melt period 2004, daily $R_{\text{eco}}$ exceeded $P_{\text{gross}}$ only slightly and for only a few days. Photosynthesis started directly when the first snow-free patches appeared and outweighed the low respiration. Similar early season trends of $CO_2$ exchange were reported for tussock tundra and flooded sedge tundra in Alaska by Vourlites and Oechel (1999), and Harazono et al. (2003). It is suggested here that the reason for this pronounced difference in the seasonal $CO_2$ progression may be related to the interannual variability of the timing of snow melt, which is thought to have a strong effect on the balance of early season $P_{\text{gross}}$ and $R_{\text{eco}}$ (Groendahl et al., 2007).

The approach for the estimation of $R_{\text{eco}}$ during winter and spring performed in this study resulted in a comparatively low value of winter and spring $R_{\text{eco}}$ of 4% of the annual $R_{\text{eco}}$. It is well known that microbial (bacterial and fungal) respiration takes place also during much of the winter and spring performed in this study resulted in a comparatively low value of winter and spring $R_{\text{eco}}$ of 4% of the annual $R_{\text{eco}}$. It is well known that microbial (bacterial and fungal) respiration takes place also during much of the winter and spring at wet tundra sites (Vourlites and Oechel, 1997; Soegaard et al., 2000; Vourlites et al., 2000; Nordstroem et al., 2001; Zamolodchikov and Karelin, 2001; Corradi et al., 2005). During the snow melt period 2004, daily $R_{\text{eco}}$ exceeded $P_{\text{gross}}$ only slightly and for only a few days. Photosynthesis started directly when the first snow-free patches appeared and outweighed the low respiration. Similar early season trends of $CO_2$ exchange were reported for tussock tundra and flooded sedge tundra in Alaska by Vourlites and Oechel (1999), and Harazono et al. (2003). It is suggested here that the reason for this pronounced difference in the seasonal $CO_2$ progression may be related to the interannual variability of the timing of snow melt, which is thought to have a strong effect on the balance of early season $P_{\text{gross}}$ and $R_{\text{eco}}$ (Groendahl et al., 2007).

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The ecosystem of the wet polygonal tundra is characterized by a comparatively low intensity of carbon cycling. Both main $CO_2$ exchange processes between the ecosystem and the atmosphere, the gross photosynthesis $P_{\text{gross}}$ and the ecosystem respiration $R_{\text{eco}}$, were low due to the environmental conditions at the site, which include climatic as well as pedogenetic factors. The net ecosystem exchange NEE depends on the balance of $CO_2$ uptake by $P_{\text{gross}}$ and $CO_2$ emission by $R_{\text{eco}}$. Since the two opposed fluxes $P_{\text{gross}}$ and $R_{\text{eco}}$ are much larger than their balance NEE, small relative changes in $P_{\text{gross}}$ or $R_{\text{eco}}$ can cause large relative changes of NEE. The measurements presented in this study showed that the wet polygonal tundra of the Lena River Delta was a substantial net $CO_2$ sink during the summer ($-119 g m^{-2}$ during June–August). Also on the annual basis, the polygonal tundra was estimated to be a $CO_2$ sink ($-71 g m^{-2}$) because the $CO_2$ efflux during autumn, winter and spring was assessed to be moderate ($+48 g m^{-2}$ during September–May).

In Table 1, the cumulative NEE observed in this study is compared with NEE values reported from other tundra sites for different time periods. The cumulative $CO_2$ uptake observed during summer at the polygonal tundra in the Lena River Delta was similar to values reported for a high-arctic fen in Greenland (Nordstroem et al., 2001), a moist to wet tussock tundra at the Kolyma River lowlands in North-East Siberia (Corradi et al., 2005) and a mixed moist and wet tundra at the Chukotsky Peninsula (Zamolodchikov et al., 2003). The mean summer $CO_2$ uptake for the whole Russian tundra area modelled by Zamolodchikov and Karelin (2001) was slightly lower than the value observed in this study. On the other hand, summer cumulative $CO_2$ uptake was several times higher at wet sedge and tussock tundra sites at an Alaskan mountain valley (Vourlites and Oechel, 1999; Vourlites et al., 2000) and at wet sedge tundra at the Alaskan
coastal plain (Harazono et al., 2003). However, cumulative summer CO$_2$ uptake at a moist to wet polygonal tundra at the coastal plain of Alaska was only about half the value measured in this study (Vourlites and Oechel, 1997). The cumulative summer CO$_2$ uptake at high-Arctic dry ecosystems was observed to be – though very variable over different years – substantially lower than at the studied polygonal tundra (Lloyd, 2001b; Groendahl et al., 2007). The estimated annual CO$_2$ sink strength at the polygonal tundra of the Lena River Delta was equal to the sink strength of a high-arctic fen at Greenland (Soegaard et al., 2000) and about half of the sink strength of wet tussock tundra at the Kolyma River lowlands (Corradi et al., 2005). On the other hand, it was about four times the average sink strength estimated for the whole Russian tundra area (Zamolodchikov and Karelin, 2001), which is due to a markedly higher estimate for $R_{ecos}$, especially for the winter period by the latter authors. Given the observed high spatial variability of NEE between tundra ecosystems, generalised conclusions on the response of the biome tundra on climatic change appear questionable (Oechel et al., 1993, 2000).

The function of tundra ecosystems as CO$_2$ sources or sinks was found to fluctuate considerably on the inter-annual and decadal scales in response to fluctuating meteorological conditions and changing climate (Groendahl et al., 2007; Oechel et al., 1993, 2000). The only eddy covariance study on NEE performed for more than two growing seasons in the Arctic demonstrated large inter-annual differences of summer CO$_2$ net ecosystem exchange at a high-Arctic heath tundra ranging over more than one order of magnitude, i.e. from $-5$ g m$^{-2}$ to $-85$ g m$^{-2}$ (Groendahl et al., 2007). This shows that CO$_2$ flux results from one or two years as given by most of the published work including this study can not be generalised easily to be valid on the multi-annual time scale which is most relevant for regional and global carbon budgets as well as climate change projections. Oechel et al. (1993) stated that Alaskan tussock and wet-sedge tundra ecosystems, which were strong CO$_2$ sinks in the cool and wet 1970s (Coyne and Kelley, 1975; Chapin et al., 1980a), had changed to a pronounced net CO$_2$ source during the mid-1980s and the early 1990s due to the acceleration of soil decomposition under a warming and drying climate. Even during the warm, photosynthetically active season (June–September), the examined tundra ecosystems were net CO$_2$ sources. However, the same authors reported that between 1992 and 1996, in response to further warming and drying, net summer releases of CO$_2$ to the atmosphere of both ecosystems declined, and they eventually became CO$_2$ sinks again (Vourlites and Oechel, 1997; Oechel et al., 2000). The authors suggested that the return to CO$_2$ sink activity of the tundra ecosystems was related to changes in nutrient cycling, physiological acclimation, and population and community reorganization which enhanced the gross primary productivity of the tundra vegetation (Shaver et al., 1998; Chapin et al., 1995, 2005; Sturm et al., 2001b). These findings underline the importance of long-term flux studies for the assessment of how Arctic ecosystems will respond to a changing climate and how this response will feed back to the climate system.

The carbon balance of the wet polygonal tundra is not complete with the study of CO$_2$, because the polygonal tundra is also a source of methane (CH$_4$). Methane is an important radiatively active trace gas, second only to CO$_2$ in its cumulative effect on the additional anthropogenic greenhouse effect. The global warming potential (GWP) of CH$_4$ for a 100-year time horizon is about 25 times the GWP of CO$_2$ (IPCC, 2007). During the micrometeorological campaigns 2003 and 2004 described in this study, also the ecosystem exchange of CH$_4$ was measured by the eddy covariance method (Wille et al., 2007$^2$). The cumulative CH$_4$ emission over the combined measurement periods 2003 and 2004 was calculated by Wille et al. to be 2.4 g m$^{-2}$ which corresponds to a carbon (C-CH$_4$) release of 1.8 g m$^{-2}$. The cumulative CO$_2$ ecosystem exchange NEE over the combined measurement periods of $-90$ g m$^{-2}$ calculated in this study corresponds to a carbon (C-CO$_2$) uptake of 24.5 g m$^{-2}$. Consequently, the C-CH$_4$ release was about 7% of the net C-CO$_2$ uptake over the combined measurement period 2003 and 2004 (end of May to middle of October). The annual cumulative C-CH$_4$ release of 2.25 g m$^{-2}$ estimated by Wille et al. was 12% of the annual cumulative C-CO$_2$ uptake of 19.4 g m$^{-2}$ estimated in this study. The ratio of the CH$_4$ emission to the CO$_2$ uptake was substantially lower at the studied Arctic polygonal tundra compared to other Siberian wetlands with higher CH$_4$ emissions (Friborg et al., 2003; Corradi et al., 2005). The greenhouse gas balance expressed as mass flux of CO$_2$ equivalents was calculated considering the 100-year GWP of CH$_4$ of 25 (IPCC, 2007). It was around $-60$ g m$^{-2}$ (uptake) for the combined measurement periods (end of May to middle of October) and not significantly different from zero ($+4$ g m$^{-2}$) for the annual estimates of CO$_2$ and CH$_4$ exchange. Further discussions on the greenhouse gas balance can be found in Wille et al. (2007$^2$).

### 5 Conclusions

This study delivered a range of new results on the processes of the CO$_2$ cycling at wet arctic tundra of Northern Siberia which are considered to be of concern not only for the Lena River Delta but also for the extensive area of the North Siberian lowland tundra as a whole. On the basis of the presented results and discussions, the following main conclusions are drawn:

The CO$_2$ budget of the investigated tundra was determined by (1.) the polar and distinctly continental climate, (2.) the very cold and ice-rich permafrost which underlies the tundra of Northern Siberia, (3.) the wetland character of polygonal

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tundra, (4.) the position at the interface between the Arctic Ocean and the Siberian mainland, (5.) the long duration of the snow coverage, (6.) the generally low nutrient status of the soils at the site, and (7.) a vegetation cover which is dominated by sedges and mosses.

The exchange fluxes of CO₂ showed clear seasonal trends on which the diurnal oscillation and pronounced meso-scale variations were superimposed. The meso-scale variations were related to the synoptic weather conditions and strongly affected the exchange fluxes through changes of cloudiness, precipitation and pronounced changes of air temperature. Thus, the large-scale atmospheric circulation patterns, for example the phase of the Arctic Oscillation, have a strong influence on the function of the North-Siberian tundra as a sink or source of CO₂.

The CO₂ budget of the wet polygonal tundra was characterized by a generally low intensity of the main CO₂ exchange processes between the ecosystem and the atmosphere, namely the gross photosynthesis $P_{\text{gross}}$ and the ecosystem respiration $R_{\text{eco}}$. Both processes were attenuated by the unfavourable environmental conditions at the site, which included climatic as well as pedogenetic factors.

The cumulative $P_{\text{gross}}$ amounted to $-432 \text{ g m}^{-2}$ over the photosynthetically active period (June–September). The rather low $P_{\text{gross}}$ was related to the low coverage of vascular plants, mainly sedges, and a high coverage of mosses at the polygonal tundra. The contribution of moss photosynthesis to the annual $P_{\text{gross}}$ was estimated to be about 40%. The gross primary productivity of the vegetation of the wet polygonal tundra was constrained by the low nutrient availability in the soils.

The diurnal response of $P_{\text{gross}}$ was mainly controlled by the irradiation. During midday the photosynthetic apparatus of the canopy was frequently near saturation and represented the limiting factor on $P_{\text{gross}}$. The seasonal progression of $P_{\text{gross}}$ was controlled by the combination of the phenological development of the vegetation and the general temperature progression over the summer. The phenological development of the plants was largely controlled by intrinsic factors. However, temperature was also a major control on $P_{\text{gross}}$ at the investigation site since photosynthesis took place for most of the time below its temperature optimum. On the other hand, water availability had only minor importance as control on $P_{\text{gross}}$ due to the wet soil conditions at polygonal tundra. Only when the mosses at the drier microsites of the polygon rim experienced water stress, $P_{\text{gross}}$ was reduced significantly.

The cumulative $R_{\text{eco}}$ amounted to $+327 \text{ g m}^{-2}$ over the photosynthetically active period (June–September), which corresponded to 76% of the cumulative $P_{\text{gross}}$. However, $R_{\text{eco}}$ continued at substantial rates during autumn when photosynthesis had ceased and the soils were still largely unfrozen and to a lesser degree throughout the winter and spring. The significant $R_{\text{eco}}$ during autumn, winter and spring was a major but highly uncertain factor in the annual CO₂ balance which should be addressed in future winter campaigns. The temporal variability of $R_{\text{eco}}$ during summer was best explained by the surface temperature and not by the soil temperature. This finding demonstrated the high and often overlooked importance of the autotrophic plant respiration within the CO₂ balance. Indeed, the composition and productivity of the vegetation had to be considered as major controls on $R_{\text{eco}}$.

Under the observed meteorological conditions, the wet polygonal tundra of the Lena River Delta acted as a CO₂ sink with a cumulative net ecosystem CO₂ exchange NEE of $-119 \text{ g m}^{-2}$ over the summer (June–August) and an estimated annual NEE of $-71 \text{ g m}^{-2}$. A more complete carbon balance must include the release of CH₄ from the ecosystem to the atmosphere. Since the carbon released as CH₄ was observed to be about 7% of the carbon uptake as CO₂ over the combined measurement period 2003 and 2004 and estimated to be about 12% of the carbon uptake as CO₂ over the whole year (Wille et al., 2007), this will still result in a clear carbon sink function of the investigated wet arctic tundra, both for the vegetation period and the whole year. However, considering the high global warming potential of CH₄, the annual greenhouse gas balance of the investigated Siberian tundra can be estimated to be near-neutral.

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