Diel and seasonal courses of ambient carbon dioxide concentration and their effect on productivity of the epilithic lichen *Lecanora muralis* in a temperate, suburban habitat

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Abstract: Ambient CO₂ concentration (together with CO₂ exchange and microclimate) was recorded every 30 min for 15 months for *Lecanora muralis* growing in the Botanical Garden Würzburg (Germany, northern Bavaria), a habitat on the outskirts of the city. Annual mean CO₂ was around 17 ppm higher than the global average reported for the time of measurement (361 ppm; 1995/96), and daily values ranged from 317 to 490 ppm. Diel courses of CO₂ could be classified into three different types. Type A, when CO₂ levels rose overnight and then fell strongly to below global levels during the day, which predominated in the summer (about 75% of days); Type B, irregular diel courses occurred during all seasons with often very rapid changes apparently due to advective CO₂ transport; Type C, CO₂ concentration was typically almost stable at generally between c. 330 and 430 ppm which predominated in the winter (63% of days).

Under controlled conditions, CO₂ saturation of net photosynthesis (NP) of *L. muralis* at optimal hydration and light occurred at around 1000 ppm. NP was also affected by low CO₂ at limiting light and thallus water contents. Based upon these data, we estimated the improvement of NP of *L. muralis* due to transient increase of ambient CO₂ (as compared with the global average) for one selected combination of environmental factors (nocturnal dew or frost). This combination is an important source of water for the lichen, resulting in 40% of its annual production and, especially in these situations, photosynthesis was increased by high ambient CO₂ in the early morning under prevailing Type A conditions. After dew activation, light compensation point of NP occurred at an average concentration of 413 ppm and diel maxima of NP at 402 ppm. This allows a rough estimate that the transiently elevated CO₂ increased the photosynthetic gain of the lichen after dew of 7%, or an improvement to its annual carbon balance of about 3%. Conditions, especially interrelationships between lichen hydration, light and CO₂ are so complex that we are not yet able to extend our estimates to other environmental situations of photosynthetic activity of *L. muralis*.

Key words: CO₂, dew, global climate change, photosynthesis, water content

Introduction

The steady increase in global atmospheric CO₂ concentration has stimulated interest in its possible effects on the performance of plants. As a result, there is now a wealth of publications dealing with plant behaviour and growth under experimentally increased CO₂ concentrations under controlled conditions and in the field (e.g., Ehleringer et al. 2005; Nösberger et al. 2006). The base lines for such studies and, at the same time, for understanding the present situation, are natural time courses of ambient CO₂ and their spatial gradients from global to ecosystem and plant levels. Most research efforts concern phanerogamic vegetation and little is known about CO₂ as an environmental factor affecting cryptogamic communities, such as lichens and mosses, especially under natural conditions. Some special situations have been investigated where mosses and lichens experience shorter or longer periods of naturally extremely elevated ambient conditions and in the field (e.g., Ehleringer et al. 2005; Nösberger et al. 2006). The base lines for such studies and, at the same time, for understanding the present situation, are natural time courses of ambient CO₂ and their spatial gradients from global to ecosystem and plant levels. Most research efforts concern phanerogamic vegetation and little is known about CO₂ as an environmental factor affecting cryptogamic communities, such as lichens and mosses, especially under natural conditions. Some special situations have been investigated where mosses and lichens experience shorter or longer periods of naturally extremely elevated ambient
CO₂. Balагеr et al. (1999) studied acclimation of a lichen to elevated CO₂ in the vicinity of a natural CO₂ spring. Tarnawski et al. (1992) report an enormous increase in CO₂ concentration within and near to the phylloplane of cushion and turf forms of the antarctic moss Grimmia antarctica and this was confirmed by Green et al. (2000) for another antarctic moss, Bryum subrotundifolium. These authors discuss the possible source of carbon dioxide which, most probably, is not moss respiration. Mosses and lichens can benefit when soil respiratory carbon dioxide is trapped within the ground boundary layer. Examples include a subarctic birch forest site (Swedish Lapland; Sonesson 1992), a temperate Nothofagus rainforest (New Zealand; Tarnawski et al. 1994), a temperate podocarp forest (New Zealand; de Lucia et al. 2003) and a subalpine spruce-fir forest (Alberta; Coxson & Wilson 2004). Kappen et al. (1995) and Sommerkorn (2000) showed that arctic-alpine lichens benefit from natural CO₂ enrichment under a spring snow-cover (Swedish Lapland). However, we did not find a single example in the literature of longer-term monitoring of CO₂ concentration at an open lichen site.

It is well known that atmospheric CO₂ concentration near the ground normally displays distinct seasonal and diel patterns with short-term variations superimposed upon the daily cycle (Geiger et al. 1995). The aim of the present work is to describe the atmospheric CO₂ environment for the epilithic lichen, Lecanora muralis (Schreb.) Rabenh., in a long-term study in a suburban, temperate habitat. We then analyse, under controlled conditions, the response of photosynthesis to changes in external CO₂ and discuss the possible influence of natural CO₂ concentrations on photosynthetic productivity of the lichen taking into account its activity and water status. It must be remembered that these measurements were made more than 10 years ago and, absolute concentrations of global CO₂ concentration have since increased. The response patterns that we found will have remained essentially identical.

Measuring site and Methods

Measurement of atmospheric CO₂ concentration (partial pressure, ppm) took place during a project in which CO₂ exchange, chlorophyll fluorescence, and microclimate conditions for several lichen species were monitored over a period of 2 years and 5 months under quasi-natural conditions in the Botanic Garden Würzburg (Germany, Bavaria, 49°45' N, 9°56' E).

Lichen photosynthesis and respiration were continuously recorded using two automatic cuvette systems (Walz Company, Effeltrich, Germany); the technical details of these 'klapp cuvettes' are described by Lange et al. (1997) and Lange (2002). For the gas exchange measurements, ambient air was continuously taken up from the lichen habitats (flow rate 1 l min⁻¹, regulated by mass flow controllers). The placing of the inlet beside the cuvettes c. 5 cm above the substratum covered by the lichen thalli, ensured that the air used for the measurements was identical to the ambient air experienced by the lichens. During winter snow cover, the air was taken from above the snow. The air stream was divided into a measurement gas stream which passed through the cuvette during CO₂ exchange measurements, and a reference gas stream. The absolute CO₂ concentration of the latter was measured every 30 minutes by an IRGA (Binos, Rosemount, Hanau, Germany). Before analysis, water content of the air streams (as well as all calibration air streams) was set by a Peltier-operated water vapor condenser. Teflon or steel tubing were used throughout the system. The analyser was calibrated at regular intervals using gas mixtures of known CO₂ concentrations (Linde A.G., Germany). Two calibration concentrations were used, one at the lower and one at the upper end of ambient CO₂ range both having been checked by means of Wösthoff pumps (see Byschlag et al. 1986). The same two cylinders were used during the entire measuring period, thus avoiding possible errors through long-term drift of the analyser. Reproducibility of the calibrations was extremely good, and we consider the absolute precision of our measurements better than ±1·5 ppm CO₂. The absolute error range of the calibration gases is given by the Linde Company as 2% (i.e. ±3·5 to ±5 ppm). Loss rate of data was very low. Downloading of data required that monitoring be stopped every four days normally for one reading, and IRGA calibrations caused additional interruptions. Technical problems usually concerned the functioning of the automatic cuvette especially during snow and frost in winter and did not affect recording of the other parameters. Thus, data set loss rate for ambient CO₂ was less than the 4% estimated for the measurements of lichen CO₂ exchange (Lange 2003a).

The total data base for all lichens in the study covered the period from March 1995 until August 1997. However, we concentrate here on measurements on the epilithic lichen Lecanora muralis (Schreb.) Rabenh. [syn. Protoparmeliopsis muralis (Schreb.) M. Choisy] made continuously from June 23, 1995 to September 29, 1996, comprising 466 diel courses (see Lange 2002, 2003a, b). We define 'one day' or 'one diel course' as...
always being from 0 to 24 h, and the ‘reference year’ from July 1995 until June 1996, inclusive, and lying within the ‘total measuring period’ of 15 months.

The measuring cuvette with the exposed samples of *L. muralis* was built into a low, 40 cm-high brick wall, covered by sandstone slabs. This installation is described and depicted in detail by Lange (2002). The lichen samples were growing on thin sandstone slices which were included in the cuvette. Measurements were always made on several lichen pieces enclosed together in the cuvette and treated as a single sample. Control experiments have shown that the substratum did not affect CO2 exchange of the lichen thalli. The wall was at an open site in the southern part of the Garden, and was surrounded by other stonewalls, rocks and pavement, all covered for decades with epilithic lichens including *L. muralis*: a typical situation for a suburban lichen habitat. The Botanic Garden is located on the outskirts of the city of Würzburg (130 000 inhabitants), c. 3 km south of the city centre on an east-facing limestone slope of the Main river valley. The Würzburg area represents a relatively dry form of the temperate, central European climate with high sunshine duration (1600 h year−1), annual mean temperature of 9°C, and an annual rain fall of c. 600 mm. For detailed information about the weather conditions during the measuring period (Fig. 1) see Lange (2003a), and for the location of the Botanic Garden within its urban surrounding see www.bgw.uni-wuerzburg.de.

The instrumentation for CO2 exchange measurements in the laboratory (Figs. 3, 4, 5) are described by Lange (2002). A ‘minicuvette system’ was used (Walz Company, Effeltrich, Germany) operating under fully controlled conditions of temperature, light [photosynthetic photon flux density (PPFD) measured over the waveband 400–700 nm], humidity, and external CO2. Factor dependence of *L. muralis* was analysed with samples almost identical to that used for the field measurements, consisting of several sectors of circular thalli growing on thin sandstone slices. Handling of the samples and methodology of the measurements are described and depicted in detail by Lange (2002). Control experiments have shown that the substratum did not affect CO2 exchange of the lichen thalli. The wall was at an open site in the southern part of the Garden, and was surrounded by other stonewalls, rocks and pavement, all covered for decades with epilithic lichens including *L. muralis*: a typical situation for a suburban lichen habitat. The Botanic Garden is located on the outskirts of the city of Würzburg (130 000 inhabitants), c. 3 km south of the city centre on an east-facing limestone slope of the Main river valley. The Würzburg area represents a relatively dry form of the temperate, central European climate with high sunshine duration (1600 h year−1), annual mean temperature of 9°C, and an annual rain fall of c. 600 mm. For detailed information about the weather conditions during the measuring period (Fig. 1) see Lange (2003a), and for the location of the Botanic Garden within its urban surrounding see www.bgw.uni-wuerzburg.de.

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**Results**

**Diel and seasonal courses of ambient CO2 concentration**

At 378 ppm the annual mean ambient CO2 concentration at the lichen site (as calculated from the continuous measurements every 30 min) was around 17 ppm higher than the global average reported for the same period of time in 1995/96 (360·74 ppm, Mauna Loa, Hawaii, 3400 m altitude, and 361·5 ppm for the global average over marine surface sites; NOAA ESRL, Global Monitoring Division, www.cmdl.noaa.gov/index.php). The highest diel maximum was 490 ppm (June 1, 1996) and the lowest minimum was 317 ppm (September 25, 1995) so that the extremes deviated by +112 ppm (29·6%) and −61 ppm (16·1%), respectively, from the average. The mean weekly minima showed a clear seasonal pattern of lower concentrations in the summer months and higher in the winter (Fig. 1) that reflected the global atmospheric annual changes. In contrast, the maxima seemed to be more the result of special local weather situations: there was no clear annual trend and extreme values occurred during almost every month of the year. The maxima were the result of advective transport from larger sources of CO2 production in combination with specific climatic situations, but we have not been able to correlate these maxima with local events, for example, with changes in automobile traffic.

As is the general rule (Geiger *et al.* 1995; Buwalda *et al.* 1992), during the summer the diel course of CO2 concentration was generally enriched with CO2 at night due to CO2 production by soil and plant respiration (Fig. 2, Type A). Increased air stability at reduced wind speed also caused high CO2 at night and this increase was especially marked with strong inversion weather situations. The increase in solar radiation in the morning produced convection and turbulence, and photosynthesis also began, causing a sharp decrease in CO2 concentration. The depression lasted for several hours with CO2 concentrations often remaining lower than 350 ppm. CO2 values increased again at dusk when photosynthesis ceased and air movement was slowed. Large changes in CO2 concentration could occur within a few hours during a single day, the maximal diel differences exceeded 150 ppm. The amplitude of such diel variability decreased in late summer and autumn and, in winter, CO2 concentration was typically almost stable at generally between c. 330 and 430 ppm (Fig. 2, Type C) with only small and irregular
ambient CO₂ concentration and weather on a weekly basis for the research period July 1995 to September 1996 for *Lecanora muralis*, Botanical Garden, Würzburg. Abscissa: weeks (every second week marked) and months. A, weekly mean, maximum and minimum of CO₂ concentration; B, weekly sum of incident photosynthetically active photon flux density (PPFD); C, weekly maxima and minima of ambient air temperature and of thallus temperature (infrared thermometer).
The three different types (A, B and C) of diel CO₂ concentration during the research period for *Lecanora muralis*, sampling frequency 30 min. A, Type A; B, Type B; C, Type C (see text).
oscillations of around 20 ppm. In addition to these regular patterns, irregular diel courses occurred during all seasons with often very rapid changes apparently due to advective CO₂ transport (Fig. 2, Type B). They resulted from sudden changes in wind speed for example, during thunderstorms, air pressure changes, heavy rain, or strong changes in solar radiation, and occurred more frequently in winter than in summer. The seasonal distribution of the three types of diel courses in CO₂ concentration is given in Table 1. Summer was characterized by a predominance of Type A, and almost 62% of all winter days belonged to Type C.

**Dependence of lichen photosynthesis on external CO₂ concentration**

As expected, *L. muralis* under controlled conditions exhibited a saturation type response curve of net photosynthesis to external CO₂ concentration (Fig. 3; Lange 2002). Under saturating light and at optimal thallus water content, CO₂ saturation (defined as lowest concentration which allowed 95% of maximal NP) was at around 1000 ppm; CO₂ compensation point was as low as 16 ppm. Minimum (317 ppm) and maximum (490 ppm) natural CO₂ concentrations recorded at the site during daylight conditions, produced 56% and 73%, respectively of maximal NP (under saturating CO₂). Net photosynthetic carbon uptake at the highest natural CO₂ was c. 30% greater than that at the lowest natural concentrations. Typically, the lichen was operating in the field at the beginning of the convex part of its CO₂ response curve, that is at the beginning of the transition between the linear response

<table>
<thead>
<tr>
<th>Season</th>
<th>Diel course type A (% of days)</th>
<th>Diel course type B (% of days)</th>
<th>Diel course type C (% of days)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Summer 1995</td>
<td>79·1</td>
<td>16·5</td>
<td>4·4</td>
</tr>
<tr>
<td>(June 23–Sept. 22)</td>
<td></td>
<td></td>
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<tr>
<td>Autumn 1995</td>
<td>25·8</td>
<td>29·3</td>
<td>44·9</td>
</tr>
<tr>
<td>(Sept. 23–Dec. 20)</td>
<td></td>
<td></td>
<td></td>
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<tr>
<td>Winter 1995/96</td>
<td>0·0</td>
<td>36·7</td>
<td>63·3</td>
</tr>
<tr>
<td>(Dec. 21–March 20)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Spring 1996</td>
<td>53·3</td>
<td>23·9</td>
<td>22·8</td>
</tr>
<tr>
<td>(March 21–June 20)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Summer 1996</td>
<td>72·3</td>
<td>23·4</td>
<td>4·3</td>
</tr>
<tr>
<td>(June 21–Sept. 21)</td>
<td></td>
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**FIG. 3.** Net photosynthesis of *Lecanora muralis* as a function of external CO₂ concentration under controlled conditions (optimal thallus water content, saturating light). The numbers 317 and 490 ppm indicate minimum and maximum external CO₂ concentration measured under field conditions. The response data (one sample consisting of several thalli, see text) were analysed by fitting of a Smith function, see Smith (1938) and Green *et al.* (1997):

$$\text{NP} = a \frac{[\text{CO}_2]}{\sqrt{1 + \left(\frac{a [\text{CO}_2]}{\text{PML}}\right)^2}} - \text{DR}$$

In this equation the dependence of net photosynthesis, NP (μmol m⁻² s⁻¹), at a given dark respiration (DR, 0·421 μmol m⁻² s⁻¹), on carbon dioxide concentration [CO₂] (ppm) is expressed by the parameters $a$, the initial slope of the curve (0·0262), and PML, the maximal rate of [NP+DR] (11·9062).
at low CO₂ concentrations and saturated values.

Optimal conditions in terms of water content (WC) and PPFD occurred, at most, only very briefly under natural conditions for the epilithic lichen in a temperate climate. CO₂ exchange was usually limited by low hydration or by suprasaturation (Lange 2003a). Changes in ambient CO₂ concentration also affected lichen net photosynthesis at suboptimal thallus water contents. Figure 4 shows drying-down curves of *L. muralis* at three different external CO₂ concentrations (274, 351, and 430 ppm). At optimal WC (c. 0.5 mm precipitation equivalent) the influence of CO₂ was similar to that in Fig. 3. The relative reduction in NP due to decreasing CO₂ (after NP_max) was initially almost constant at different thallus water contents and remained detectable at very low thallus WC, in some cases close to the moisture compensation point of the lichen. In contrast, changes in external CO₂ concentration over the range found in nature had no detectable impact on NP when CO₂ uptake was depressed at a WC higher than c. 0.85 mm (the right portion of the response curves in Fig. 4). Net photosynthesis was reduced almost to zero because thallus diffusion resistance had increased dramatically, restricting CO₂ uptake almost completely. We know that much higher concentrations of external CO₂ are necessary to overcome this barrier (Lange 2002).

Changes in external CO₂ concentration also affected NP at light intensities below light saturation but only at WC lower than that causing suprasaturation. This is shown in Fig. 5 where light response curves of *L. muralis* are depicted for four different ranges of thallus water content and CO₂ concentrations of 330 and 430 ppm. Differences in CO₂ concentration affected NP down to about 100 μmol m⁻² s⁻¹ PPFD (the three lowest WC ranges). At still lower light intensities, scatter of measuring points hid differences in this data set. In other experiments with *L. muralis* and with other species we found that larger differences in external CO₂ concentration (data not shown) affected the light compensation point of CO₂ exchange, confirming earlier findings by Sommerkorn (2000) and by Coxson and Wilson (2004). At thallus suprasaturation (Fig. 5, lowest pair of curves), again, there was no
detectable difference between 330 and 430 external CO₂.

Possible significance of external CO₂ changes for lichen productivity

Metabolic activity of lichens is restricted by the occurrence of periods of desiccation. The thin thallus of *L. muralis* had a low water holding capacity and showed high rates of water loss after hydration by dew, fog, rain or high air humidity alone. It was active for only 35·6% of the total time of the year, made up of 16·7% net photosynthetic CO₂ uptake and 18·9% respiratory CO₂ release (Lange 2003a). Only the periods of photosynthesis are influenced by changes in ambient CO₂ concentration. We characterized the environmental CO₂ conditions on each day when the lichen was photosynthetically active by three values, namely the CO₂ concentration: (1) at the time of light compensation during sunrise when the thallus had been hydrated during night or at the beginning of net photosynthesis if hydration took place later during the daylight, (2) at the time of diurnal maximum of CO₂ uptake, and (3) at the end of the photosynthetically active period, either, and most commonly, marked by the moisture compensation point after desiccation, or by the light compensation point at dusk. The high variability of these values throughout the year means that the averages of these numbers (Table 2) are not statistically different, but they do show characteristic tendencies. The annual mean of daily ambient CO₂ concentration at (1), 395 ppm, was higher than it was during the daily maximum of net photosynthesis (2), 380 ppm, and at the end of the photosynthetically active period (3), 371 ppm (Table 2, upper row). If we assume that the average of these three parameters was the mean CO₂ concentration at which *L. muralis* performed photosynthetic CO₂ exchange at its site in the Botanic Garden Würzburg, we arrive at 382 ppm, which is 21 ppm higher than the global standard concentration for the same time period.

Days with dew (and fog) or frost are very important for the carbon budget of *L. muralis*; they contribute 40% to the total annual carbon balance of the lichen (Lange 2003b). Usually, these periods of photosynthetic activity when the lichen could substantially benefit from increased concentrations of ambient CO₂ lasted only for a short period of time after sun rise. Therefore, these situations were especially suited for a more
detailed analysis. Nocturnal dew and fog formation usually occurred on nights with high long wave radiation loss and a weather situation with stable air and low turbulence near the ground, that is, during conditions which promote nocturnal CO₂ enrichment (Type A situation, Fig. 2). Figure 6 shows an example of a typical lichen performance in summer under such conditions. The thallus became moistened by dew during the night which resulted in low rates of respiration. When the light compensation point was surpassed at 4:45 a.m. (below c. 30 μmol m⁻² s⁻¹ PPFD) ambient CO₂ was 467 ppm. It was 460 ppm when maximal NP was reached and had decreased to 428 ppm at the moisture compensation point at 7:30 a.m. when the lichen became dry again. Thus, the average CO₂ concentration during its photosynthetic activity at

<table>
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<tr>
<th></th>
<th>CO₂ at beginning of positive NP</th>
<th>CO₂ at maximum of positive NP</th>
<th>CO₂ at end of positive NP</th>
<th>CO₂ average during positive activity</th>
</tr>
</thead>
<tbody>
<tr>
<td>All days of the year</td>
<td>395</td>
<td>380</td>
<td>371</td>
<td>382</td>
</tr>
<tr>
<td>Days with dew activation</td>
<td>410</td>
<td>401</td>
<td>384</td>
<td>398</td>
</tr>
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Fig. 6. Time courses for *Lecanora muralis* on 20 July 1995, Botanical Garden, Würzburg. A, ambient CO₂ concentration and area-related CO₂ exchange; the CO₂ concentrations are indicated for the light compensation point in the morning, maximally achieved net photosynthetic rate, and moisture compensation point after drying; B, incident light (PPFD) and air temperature.
that day was c. 452 ppm which is substantially higher than the global annual concentration. During the two summer periods, 1995 and 1996, the lichen experienced 74 days with dew hydration of which 68 days (92%) showed a Type A CO₂ pattern, only 5 days were Type B, and 1 day Type C. In Fig. 7 ambient CO₂ concentration is shown at light compensation point, maximal NP, and end of the photosynthetic activity for all dew (and frost) events of the total measuring period, with average values of 413, 402, and 385 ppm, respectively. The respective average values for the reference year are given in Table 2 (lower row).

We can obtain a rough estimate of the increase in net photosynthetic carbon gain of the lichen under the actual ambient CO₂ conditions as compared with a CO₂ environment of 361 ppm, the globally averaged concentration for the years 1995/96 (see above), by looking at selected situations. Restricting the calculation to dew (and frost) hydration and using the regression function depicted in Fig. 3, NP increases from 100% at 361 ppm to 107% at 402 ppm, the average ambient CO₂ during maximal NP after dew moistening. We assume that the lichen would respond similarly to changes in CO₂ under the field conditions at maximal NP after dew as it did under optimal water content and saturating light, a view supported by experiments shown in Figs. 4 and 5. If this assumption is accepted, and if we exclude adaptive responses of the lichen photosynthetic and respiratory apparatus (see e.g., Lange & Green 2005), then average maximal NP after dew under the given environmental conditions will be around 7% higher than at a level of 361 ppm. In the extreme case, at an ambient CO₂ concentration of 482 ppm, the increase would reach 18%. In Fig. 8, for all dew events with L. muralis, diel photosynthetic carbon income showed a strong linear relationship to maximal rate of NP. This means, that increase of ambient CO₂ from 361 ppm to an average of 402 ppm for NPₘₐₓ would also result in an increase in gross primary production of 7% which is substantial if one considers that dew and frost hydration provide a large proportion of the total annual carbon budget of the lichen. Interactions between ambient CO₂ and carbon income for the other weather response types (see Lange 2003a, b) such as days with heavy rain or with nocturnal rain and subsequent diurnal drying of the lichen are so complex, that a realistic guess of the impact of ambient CO₂ is not yet possible.

**Discussion**

The mean CO₂ concentrations calculated from measurements directly adjacent to the experimental lichens monitored for over a year (1995/1996) were slightly higher than the accepted global mean of 361 ppm. Over the whole year the mean elevation was 17 ppm and, during the period in the early morning when the lichen was physiologically active, about 21 ppm. The mean values camouflage some high variability in daily and seasonal patterns during the year. The daily range was substantially higher in the summer, reaching over 180 ppm, compared to a much more stable winter value often of around 20 ppm. The cause of this change is well known and it reflects the more rapid life processes (photosynthesis and respiration) in the warmer summer. Although the lichen might have had 120 ppm higher CO₂ at dawn, it could spend the day many ppm (up to 60) below the accepted global value. These values are for a lichen within the ground boundary layer and it is almost certain that they would be different for fruticose or epiphytic species. Other studies have also found elevated CO₂ concentrations, in particular when a boundary layer occurs together with a soil respiratory source. Good examples of this would be two studies in forests in New Zealand (Tarnawski et al. 1994; de Lucia et al. 2003). In these two cases the respiratory CO₂ was probably generated by tree roots in the soil and it was estimated that the bryophyte soil mat recycled 10% of the tree respiration (de Lucia et al. 2003). The extreme values are found in two antarctic studies where CO₂ concentrations can reach many times the accepted normal values during the day (Tarnawski
F. Ambient CO₂ concentration for *Lecanora muralis* on days with metabolic activity initiated by nocturnal moistening with dew (fog) or frost during the measuring period (abscissa, weeks and months). A, CO₂ concentration at light compensation point in the early morning; B, CO₂ concentration at the daily maximum net photosynthetic activity; C, CO₂ concentration at the end of the daily activity (moisture compensation point) when the lichen had become dry again (bottom). In each panel the average CO₂ concentration is given for the measurement period.

**Fig. 7.** Ambient CO₂ concentration for *Lecanora muralis* on days with metabolic activity initiated by nocturnal moistening with dew (fog) or frost during the measuring period (abscissa, weeks and months). A, CO₂ concentration at light compensation point in the early morning; B, CO₂ concentration at the daily maximum net photosynthetic activity; C, CO₂ concentration at the end of the daily activity (moisture compensation point) when the lichen had become dry again (bottom). In each panel the average CO₂ concentration is given for the measurement period.
et al. 1992; Green et al. 2000). In these cases the source of the CO₂ remains a mystery. It seems, therefore, that the CO₂ concentrations around lichens are often different from the global values. The question is, does this situation affect the productivity of the lichens by altering their photosynthetic rate? The lichen studied here, *L. muralis*, has been extensively researched and its photosynthetic response to CO₂ and other environmental factors such as temperature and light, are well known (Lange 2002, 2003a, b). By taking a selected combination of environmental factors, namely the hydration of the lichen by dew or frost in the early morning, it was possible to show that an additional gain of around 7% in photosynthetic rate occurred due to elevated CO₂. The calculation was possible for several reasons: first, we could use the regression line depicted in Fig. 3 as an approximation for NP response to CO₂ concentration in the field; second, the WC of the thallus is optimal or suboptimal when hydrated by dew (there is no CO₂ enhancement at high WC); and third, the proportional effect of the higher CO₂ occurs at all optimal and suboptimal WC and light levels.

It is less easy to estimate gains or losses under other conditions. When the lichen is moistened by extended rain, NP is usually temporarily heavily depressed by high diffusion resistances so that changes in external CO₂ concentration have practically no effect on rates of NP. This situation occurs on 74 days, is the most productive in photosynthetic gain but makes almost no contribution to the annual carbon budget (4%) because of concomitant high respiration. Two other environmental combinations, wetted overnight and drying during the day, and light showers and continuously wet, make a substantial 35-5% contribution to the annual carbon gain, but, and it is a large but, in these cases most photosynthetic activity occurs during the day when CO₂ concentrations are most likely to be decreased below normal global values. At these times the lichens are likely to be losing carbon fixation potential because of the decreased CO₂ concentration. However, the data available do not allow a detailed analysis of this complex problem.

To fully quantify the advantage of elevated ambient CO₂ for carbon gain in *L. muralis* in its suburban habitat, a model would be necessary which takes into account CO₂ dependence of its photosynthesis at different light, temperature and hydration conditions including, for example, CO₂ dependence of the light compensation point. However, sufficient data for these complex interrelationships are not available. At best we can say that the hydration by dew contributes 40-0% of *L. muralis* carbon gain (Lange 2003b) so that we are able to calculate an overall increase of (7·0 × 0·4) about 3% of total lichen carbon gain due to transiently increased external CO₂.

We cannot easily extend this to other lichens because not only is the interaction between NP and WC almost species-specific but so, also, is the response of NP to CO₂ concentration. Many lichens appear to be saturated at lower CO₂ levels than *L. muralis*. Examples include *Flavoparmelia caperata* (400 ppm; Balaguér et al. 1999), *Sticta latifrons*, *Pseudocyphellaria billardierii*, *Peltigera dolichorhiza* and *Sphaerophorus ramulosum* (450 ppm; Green & Snelgar 1981). These species might be expected to show little or...
no response to changes in ambient CO₂ concentrations. However, these responses were all measured under optimal conditions of light and WC and it is known that these optimal combinations are actually very rare (Lange et al. 1993). Certainly, at suboptimal WC we might well expect an improved NP with higher CO₂ concentrations.

It is also not so easy to predict what the effects of the present steady increase in global atmospheric CO₂ concentrations will be. For L. muralis we can with some confidence say that increases up to around 600 ppm will have an almost linear affect on NP as the lichen is only entering the convex part of its CO₂ response curve. At higher levels, the gain will decline as saturation is approached. For other lichens, particularly those with very low CO₂ concentrations for saturating NP, we can do little more than guess that, because most lichens operate at non-optimal conditions, there will be a slight increase in carbon gain.

Constructing a model to predict the effects of changes in CO₂ concentrations will also be very difficult. The major difference between lichens and higher plants is the poikilohydric nature of lichens which not only means that they spend substantial times desiccated and dormant but also means that there are species-specific differences in water relations and that there is a complex and species-specific interaction between thallus water content and net photosynthesis. As is true for many other lichen carbon fixation problems, the accurate modelling and prediction of thallus water content remains the Holy Grail of lichen ecophysiology.

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