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published in
Biogeosciences
2006

document version
Publisher's PDF, also known as Version of record

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citation for published version (APA)

van der Tol, C., Dolman, A. J., Waterloo, M. J., & Raspor, K. (2006). Topography induced spatial variations in diurnal cycles of assimilation and latent heat of Mediterranean forest. *Biogeosciences*, 3, 1631-1677.
<http://www.biogeosciencesdiscussions.net>

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**Topography induced
variations in
assimilation and
latent heat**

C. van der Tol et al.

Topography induced spatial variations in diurnal cycles of assimilation and latent heat of Mediterranean forest

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Received: 4 September 2006 – Accepted: 29 September 2006 – Published: 6 October 2006

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Abstract

The aim of this study is to explain topography induced spatial variations in the diurnal cycles of assimilation and latent heat of Mediterranean forest. Spatial variations of the fluxes are caused by variations in weather conditions and variations in the vegetation characteristics. Weather conditions reflect short-term effects of climate, whereas vegetation characteristics, through adaptation and acclimation, long-term effects of climate. In this study measurements of plant physiology and weather conditions are used to explain observed differences in the fluxes. A model is used to study which part of the differences in the fluxes is caused by weather conditions and which part by vegetation characteristics. Data were collected at four experimental sub-Mediterranean deciduous forest plots in a heterogeneous terrain with contrasting slopes and aspect, soil water availability, humidity and temperature. We used a two leaf layer model to scale fluxes from leaf to canopy, and calculated the canopy energy balance. Parameter values were derived from measurements of light interception, leaf chamber photosynthesis, leaf nitrogen content and ^{13}C isotope discrimination in leaf material. Leaf nitrogen content is a measure of photosynthetic capacity, and ^{13}C isotope discrimination of water use efficiency. For validation, sap-flux based measurements of transpiration were used. The model accurately predicted diurnal cycles of transpiration and stomatal conductance, both their magnitudes and differences in afternoon stomatal closure between slopes of different aspect. The diurnal cycles were more strongly affected by spatial variations in vegetation parameters than by meteorological variables. This indicates that topography induced variations in vegetation parameters are of at least equal importance to the fluxes as topography induced variations in radiation, humidity and temperature. Weather conditions mainly affect the shape of the diurnal cycles, and vegetation parameters the magnitude of the fluxes.

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1 Introduction

Surface exchange models for water and carbon dioxide have developed in the last 30 years from simple conceptual models towards more accurate descriptions of the soil-vegetation-atmosphere system. This development is driven by an increasing emphasis on the carbon balance as a focal point of atmospheric modelling beside the energy and water balance (Sellers et al., 1997). A great improvement has been the discovery of a close relation between stomatal conductance and photosynthesis rate (Wong et al., 1979), and the consequent integration of the descriptions of transpiration and photosynthesis (Lloyd et al., 1995; Harley and Baldocchi, 1995; Tuzet et al., 2003).

Two important processes in these integrated models, are carboxylation of carbon dioxide in leaves, and transport of carbon dioxide from the air into leaves. Transport of carbon inevitably involves loss of water by transpiration, which travels along the same path as carbon but in opposite direction. The uptake of carbon dioxide and the loss of water by transpiration is regulated by stomatal conductance. The two most important characteristics of vegetation to describe these processes are unquestionably the photosynthetic capacity and the way in which vegetation regulates stomatal conductance (Farquhar and Sharkey, 1982).

Both weather conditions and vegetation characteristics affect the fluxes of water and carbon. The effect of vegetation is in fact an indirect effect of climate: the evolution of vegetation depends on long-term climate (Hetherington and Woodward, 2003). Thus, we can distinguish short term (or direct) from long-term (or indirect) effects of climate. Short term effects are those of radiation, temperature and vapour pressure deficit, and long term effects those of photosynthetic capacity and regulation of stomatal conductance. Both weather conditions and vegetation characteristics vary spatially, and are most likely correlated in some way.

This study focuses on the quantification of the effects of those spatial variations in weather conditions and vegetation characteristics on the diurnal cycle of latent heat flux. The two aims of the study are (1) to translate vegetation characteristics into

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diurnal cycles of photosynthesis and transpiration, and (2) to separate the effects of weather conditions and vegetation characteristics on the diurnal cycles of assimilation and latent heat flux using a sensitivity analysis.

We use a sun-shade model for photosynthesis and the energy balance to calculate the fluxes of carbon dioxide and water from leaf parameters. This model is similar to existing models for scaling of fluxes from leaf to canopy (Leuning et al., 1995). The novelty of our approach is the data we use for parametrization and validation of the model. Data were collected during a field campaign at four experimental plots in natural broadleaf sub-Mediterranean forests in Slovenia, which contrast in local hydrological and climate conditions, aspect and vegetation composition. Contrary to studies which calibrate vegetation parameters from measured fluxes, we used independent measurements at leaf level for parametrization, and sap flux density measurements for validation. For parametrization we use leaf nitrogen content, ^{13}C isotope discrimination of leaf material, and leaf chamber photosynthesis measurements. In this way, we test whether canopy scale fluxes can be predicted from leaf parameters. Spatial patterns in vegetation parameters and fluxes give an idea of how adaptation works. Next, we perform a sensitivity analysis of the model to separate the effects of vegetation parameters and weather conditions on surface conductance and the fluxes of carbon dioxide and water.

2 Method and materials

2.1 Model description

The model we used to calculate the fluxes of water and carbon dioxide consists of three components: photosynthesis and transpiration at leaf level as a function of biochemical parameters, scaling of the fluxes to canopy level, and an energy balance of the canopy. At leaf level, diffusion equations, a biochemical model for photosynthesis (Farquhar et al., 1980) and the model for optimal stomatal control of Cowan (1977) are

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used. Scaling from leaf to canopy is carried out with a two-leaf model that distinguishes between a sunlit and a shaded fraction of leaves. An energy balance of the canopy is used to solve canopy temperature, which in turn affects the processes in the leaves. Figure 1 shows the structure of the model. In what follows, the three components of the model are presented.

Water and carbon dioxide move by diffusion in opposite directions between the stomata and the air. Water evaporates from the cell walls, and travels from the stomata to the air, whereas carbon dioxide travels from the air, via the stomata into the mesophyll, where it is reduced to sugars by the chemical reactions in the Calvin cycle. If the resistance for transport of carbon dioxide from the stomata to the mesophyll is neglected, then the diffusion equations can be written as:

$$E = 1.6g \frac{\rho_a}{M_a} \frac{e_i - e_a}{p} = 1.6gD \quad (1)$$

$$A = g(C_a - C_i) \quad (2)$$

where E is evaporation and A assimilation ($\text{mol m}^{-2} \text{s}^{-1}$), g the effective aerodynamic and stomatal conductance (m s^{-1}), e_i and e_a the vapour pressure in the intercellular spaces and in the ambient air (Pa), respectively, p atmospheric pressure (Pa), ρ_a specific mass of air (kg m^{-3}), M_a the molar mass of air (kg mol^{-1}), $D = \frac{\rho_a}{M_a} \frac{e_i - e_a}{p}$ the molar vapour concentration gradient between the intercellular space and the air (mol m^{-3}), and C_a and C_i the molar carbon dioxide concentration in the ambient air and in the stomata (mol m^{-3}), respectively. The process of photosynthesis is described with the biochemical model of Farquhar et al. (1980):

$$A = \frac{v(C_i - \Gamma^*)}{C_i + \gamma} - R_d \quad (3)$$

$v = V_{cm}$ and $\gamma = K_c \left(1 + \frac{O}{K_o}\right)$ for Rubisco limited photosynthesis and $v = \frac{qI J_m}{4(I + 2.1 J_m)}$ and $\gamma = 2\Gamma^*$ for photon limited photosynthesis, V_{cm} is the maximum carboxylation capacity

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of Rubisco ($\text{mol m}^{-2} \text{s}^{-1}$), Γ^* the compensation point for carbon dioxide in absence of dark respiration (mol m^{-3}), O the oxygen concentration (mol m^{-3}), K_o and K_c the Michaelis-Menten constants for carbon dioxide and oxygen (mol m^{-3}), R_d dark respiration ($\text{mol m}^{-2} \text{s}^{-1}$), q the quantum yield efficiency, J_m the maximum potential electron transport rate ($\text{mol m}^{-2} \text{s}^{-1}$) and I the irradiance by photosynthetically active radiation (PAR) ($\text{mol m}^{-2} \text{s}^{-1}$).

Once values for the biochemical parameters are known, the diffusion equations (Eqs. 1 and 2) and the biochemical model (Eq. 3) form a set of three equations containing four unknowns (A , E , g and C_i). A fourth equation, describing the stomatal behaviour, is required to yield a unique solution. Cowan (1977) and Cowan and Farquhar (1977) suggested that stomata operate such as to minimize the evaporative cost of plant carbon gain. This condition is met if the marginal water cost of assimilation Λ , is constant with time:

$$\frac{\delta E / \delta g}{\delta A / \delta g} = \Lambda = \text{constant} \quad (4)$$

This model does not explain how stomatal regulation works physiologically, but only describes the stomatal behaviour that yields the highest mean assimilation rate over a time period with variable environmental conditions, during which a certain positive amount of water evaporates. The parameter Λ is a measure for the intrinsic water use efficiency (Lloyd and Farquhar, 1994). Low values of Λ refer to more water efficient vegetation than high values. The advantage of this model for stomatal behaviour compared to empirical relations between stomatal conductance and humidity deficit, is that only one parameter, which has a conceptually clear meaning, is used. The model works best for the diurnal cycle, although it has also been applied to longer time periods, including dry conditions (Cowan, 1986). A problem with longer time scales and dry periods is that Λ does not remain constant (Makela et al., 1996; Arneth et al., 2002), because of hydraulic limitation of transport of water (Tyree and Sperry, 1988; Jones, 1998; Mencuccini, 2003) and because stomata respond to abscisic acid transmitted by

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roots (Zhang and Davies, 1989).

With Eqs. (1) to (4), the variables A , E , g and C_i are uniquely defined. Solving Eqs. (1) to (4) results in a C_i that is a function of the vapour gradient D between the intercellular space and the air (Appendix A). The solution is different for the case in which assimilation is enzyme limited and for the case in which assimilation is electron limited. Because it is not known a-priori whether assimilation is enzyme or electron limited, C_i is solved by iteration of Eqs. (A1) to (A3) and the biochemical model (Eq. 3).

Climate variables in this model are photosynthetically active radiation (PAR), vapour pressure deficit and carbon dioxide concentration. Biochemical parameters are maximum carboxylation capacity V_{cm} and maximum electron transport J_m , quantum yield efficiency q , marginal cost of assimilation Λ and Michaelis-Menten coefficients for the chemical reactions in the Calvin cycle.

The approach used to scale from leaf to canopy level (Appendix B) is similar to that of De Pury and Farquhar (1997). Lambert-Beer's equation is used to calculate the vertical distribution of light in the canopy, discriminating between indirect (diffuse) and direct light, which have different extinction coefficients. Because the experimental sites were located on steep slopes, a coordinate rotation was used to correct for the effect of topography on the extinction coefficients for direct light (Appendix B). The model calculates the exposed and shaded fraction of leaves (f_e and f_s), and the intensities of PAR on the exposed and shaded leaves (I_e and I_s), which are variable over the day. The fluxes and the surface conductance at canopy level are calculated by adding the contributions of the two fractions:

$$V = L (f_e v_e + f_s v_s) \quad (5)$$

where V and v refer to any of the variables A , E and g at canopy and leaf level, respectively, the index e to exposed and s to shaded, and L is leaf area index. The effective internal carbon dioxide concentration for the canopy is:

$$C_i = C_a - \frac{A}{G} \quad (6)$$

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where A is canopy assimilation and G the canopy-scale equivalent of g .

For the calculation of E with Eq. (1), an estimate of the internal vapour pressure e_i is needed, which cannot be measured directly. The intercellular vapour pressure is in equilibrium with leaf water potential and leaf temperature. Because air in stomata is always close to saturation, we assume that e_i is the saturated vapour pressure e_s at surface temperature T_s :

$$e_i = e_s(T_s) \quad (7)$$

Surface temperature is solved from the energy balance. Neglecting soil heat flux and changes in heat storage, the energy balance is:

$$R_n = H + \lambda E \quad (8)$$

where R_n is net radiation, H sensible and λE latent heat flux (all in W m^{-2}). Latent heat flux is calculated by converting E from units of $\text{mol m}^{-2} \text{s}^{-1}$ to $\text{kg m}^{-2} \text{s}^{-1}$ and multiplying by the latent heat of vaporisation of water, $\lambda (=2.501 - 0.0024 T(^{\circ}\text{C})) \text{ MJ kg}^{-1}$.

$$\lambda E = 1.6\lambda\rho_a G \frac{M_{\text{H}_2\text{O}}}{M_a} \frac{e_i - e_a}{\rho} \quad (9)$$

where $M_{\text{H}_2\text{O}}$ the molar mass of water (kg mol^{-1}). Sensible heat flux is calculated as:

$$H = \rho_a c_p G_a (T_s - T_a) \quad (10)$$

where G_a the aerodynamic conductance. Net radiation is calculated from incoming and outgoing shortwave and longwave radiation:

$$R_n = (1 - \alpha)R_{si} + R_{lj} - R_{lo} \quad (11)$$

where R_{si} , R_{lj} and R_{lo} incoming shortwave, incoming longwave and outgoing longwave radiation, respectively, and α the reflection coefficient for shortwave radiation (albedo). Outgoing longwave radiation is calculated with Stefan-Boltzman's equation:

$$R_{lo} = \varepsilon\sigma T_s^4 \quad (12)$$

where $\varepsilon=0.98$ the emissivity of the canopy and σ Stefan-Boltzman constant ($=5.67 \times 10^{-8} \text{ W m}^{-2} \text{ K}^{-4}$).

From the biochemical model (Eqs. 1 to 4), A , E , C_i and g can be solved, provided that e_i is known. Equation (5) is used to scale from leaf to canopy, and from the energy balance (Eqs. 7 to 12), e_i , T_l , R_{l0} , R_n , H and λE can be solved, provided that G and G_a are known. Both sub-models should be solved simultaneously. An analytical solution is not possible due to the non-linearity of the interactions between the sub-models. For this reason, surface temperature is adjusted iteratively in order to force energy balance closure.

The aerodynamic conductance G_a is calculated with a wind function:

$$G_a = G_{a0} + Bu \quad (13)$$

where G_{a0} a convective term, u wind speed measured at the meteorological station (m s^{-1}), and B an empirical wind function. The terms G_{a0} and B are calibrated by optimising calculated surface temperatures T_s with measured ones. Because T_s is measured only at the south plot, the parameters G_{a0} and B of the south plot are also used at the other three plots.

2.2 Site description

The study was part of a broader project to study the effects of natural reforestation on the water balance and geomorphology of the river catchment of the Dragonja River in Mediterranean Slovenia (N $45^{\circ}28'$ E $13^{\circ}46'$, Fig. 2).

The Dragonja catchment is located within a 30 km wide band along the Adriatic coast of the peninsula of Istria with has a Sub-Mediterranean climate. The Julian Alps and up to 1500 m high Karst plateaus form sharp orographic boundaries at the north and the east with a more continental climate, with lower temperatures and higher precipitation. The Sub-Mediterranean climate is classified as Caf (mild winter, hot summer, no dry season) in the Köppen system. Mean annual precipitation varies from 1300 mm at the source to 1000 mm at the outlet of the Dragonja and is distributed evenly over the year.

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The parent material in the Dragonja catchment is flysch: a sequence of calcareous shales and thin sandstone banks. In the upper part of the catchment, broad plateaus are intersected with narrow, steep river valleys of two contributing streams. In the lower part, the valley is broad and the plateaus narrow. The elevation ranges between 0 and 330 m above sea level. Soils in the whole catchment are Rendzina soils (Keesstra, 2006) and consist of clay loam (30 percent sand, 50 percent silt, 20 percent clay). Soil depth ranges from a few decimeters on the slopes to several meters of alluvial deposits in the valley.

Four experimental plots were selected in deciduous forests, which contrasted in aspect, local hydrological and climate conditions and vegetation composition (Fig. 2). The forests had developed with minimum human interference during the last 50 years. Both texture and chemical composition of the soils at the plots were similar. One plot was located on a north and one on a south facing slope (north and south plot), and one at the foot of a converging west facing slope (west plot) and one on a diverging south facing slope (east plot). Although the east plot has a south facing aspect, it was named east plot to distinguish it from the south plot, and because it is located the most to the east.

The differences in water availability, light, temperature and vapour pressure deficit among the plots are presented schematically in Fig. 3. Two plots are predominantly sunlit and experience a high vapour pressure deficit, temperature and radiation input (south and east), and two plots predominantly shaded and experience a low vapour pressure deficit, temperature and radiation input (north and west). Two plots experience a high (north and east) and two plots a low (south and west) water availability. In this way, each of the four combinations of high and low vapour pressure deficit and high and low soil moisture content was present.

The plots not only contrasted in micro-environment, but also in species composition (Fig. 4), stem density and forest structure (Table 1). The dominant species were *Carpinus betulus* at the two shaded plots (north and west), and *Quercus pubescens* at the two exposed plots (south and east). Trees at the shaded plots were taller than at

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the sunlit plots. The forest at the east plot was younger, and pioneer vegetation was present (*Juniperus communis*).

2.3 Measurements

At each plot, vegetation and soil parameters, meteorological variables and the water balance were measured between May and September 2004. We selected only those data for which vegetation was not limited by water availability. The reason for doing so is that we are interested in the diurnal cycle during a period in which biochemical parameters remain constant, and water stress causes biochemical parameters to change in time (Lambers et al., 2000). Those temporal variations of biochemical parameters in a changing environment are the subject of a study we will publish separately.

Basic meteorological variables (wind speed, diffuse and direct incoming shortwave radiation and reflected shortwave radiation) were measured at a meteorological station 3 km east of the experimental plots. Parameters for the biochemical model at leaf level were derived from leaf chamber photosynthesis measurements carried out on two species at the south and the east plot, and leaf nitrogen content and ^{13}C isotope discrimination at all four plots.

Light response curves of photosynthesis and transpiration of leaves of *Quercus pubescens* and *Fraxinus ornus* were measured using a broadleaf leaf chamber with portable light unit connected to an LCA3 gas analyser (ADC BioScientific Lt., UK) at the east and the south plot between 14 and 21 July 2004. At the east plot, the trees were so small that measurements could be carried out at breast height. At the south plot, measurements were carried out on a scaffolding tower of 9 m height. The two species sampled are the most abundant species at the south and the east plot.

2.4 Leaf sample analysis

Leaf samples for analysis of carbon and nitrogen content and ^{13}C isotope discrimination were collected by a professional tree climber at the start (a few weeks after bud

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break) and the end (four weeks before the onset of senescence) of the growing season at all four plots. The total number of leaf samples was 83 (15 at the north, 31 at the south, 16 at the west and 21 at the east plot). One third of the samples was collected between 5 May and 8 June 2004, and two third between 8 and 10 September 2004.

Each sample consisted of 4 to 15 leaves of different size from neighbouring branches of a tree. The number of samples of a species was chosen such that it approximated the relative contribution to the total sapwood area of that species at the plot. Species which contributed less than 5 percent to the total sapwood area were not sampled. At all plots, the sampled species represented over 85 percent of the sapwood area. Both predominantly sunlit and predominantly shaded leaves were collected. Samples were classified by plot, species, sunlit or shaded, and young or old leaves.

The leaves were air dried, oven dried at 50°C, minced in a mincing machine and grounded in centrifugal ball mill. Carbon and nitrogen content (percentage by weight) and discrimination of ^{13}C were determined using an elemental CHNO-analyzer Flash EA 1112 (Finnegan MAT, Bremen, Germany).

The ^{13}C discrimination against ambient air was calculated as (Farquhar and Richards, 1984):

$$\Delta^{13}\text{C}_l = \frac{\delta^{13}\text{C}_a - \delta^{13}\text{C}_l}{1 + \Delta^{13}\text{C}_l} \quad (14)$$

where $\delta^{13}\text{C}$ the isotope ratio per mil compared to the PDB standard, subscript *a* and *l* indicate air and leaf, respectively, and $\delta^{13}\text{C}_a = -8$ ppm. The long-term internal carbon dioxide concentration is calculated as (Farquhar et al., 1989):

$$\frac{C_l}{C_a} = \frac{\Delta^{13}\text{C}_l - c_1}{c_2 - c_1} \quad (15)$$

where $c_1 = 4.4$ per mil the discrimination by diffusion in air and $c_2 = 27$ per mil the discrimination by Rubisco.

The leaf samples were used to calculate effective values for leaf nitrogen content and ^{13}C isotope discrimination for each plot. First, a Kolmogorov-Smirnov test showed that the measurements of leaf nitrogen and ^{13}C within each plot had normal distributions, and a Levene test showed the variances within the plots were not different from each other. Next, the estimated mean values \hat{m} and variances \hat{s}^2 for each plot were calculated as:

$$\hat{m} = \sum_{n_s} F_i m \quad (16)$$

$$\hat{s}^2 = \sum_{n_s} F_i^2 \hat{s}_i^2 \quad (17)$$

where F_i the contribution of species i to total sapwood area, and n_s the number of sampled trees of species i . A 95% confidence interval for the mean was calculated as:

$$m = \hat{m} \pm s_{0.95} \sqrt{\frac{\hat{s}^2}{n_p}} \quad (18)$$

where $s_{0.95}$ the Student-t statistic for $p=0.95$ and n_p the number of samples at each plot.

The parameters for the model of Farquhar and internal carbon dioxide concentration for each plot were derived in the following way. First, maximum carboxylation capacity and electron transport capacity for each plot were calibrated from the leaf chamber photosynthesis measurements. Next, a linear relationship between maximum carboxylation capacity and leaf nitrogen content was calibrated for the leaves for which measurements of both leaf nitrogen content and maximum carboxylation capacity were available. Finally, this linear relationship was used to derive values for the maximum carboxylation capacity at all four plots from measurements of leaf nitrogen content.

The average values of internal carbon dioxide concentration \bar{C}_i from the isotope analysis were used to calibrate the model parameter Λ for each plot, by minimising the

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absolute difference between measured and calculated internal carbon dioxide concentration. Modelled internal carbon dioxide concentration changes during the day. An average was calculated by weighing C_i with the instantaneous rate of photosynthesis (Farquhar et al., 1982):

$$\bar{C}_i = \int \frac{A(t)C_i(t)}{\bar{A}} dt \quad (19)$$

Temperature, relative humidity, vertical profiles of soil moisture content and sap flux density were measured continuously at each plot, and data stored at 30-min intervals. Precipitation was measured at 3 stations within 500 m of the forest plots. Transpiration was calculated from sap flux density measurements with the method of Granier (1987).

At each plot, 12 sensors were installed in 6 trees (2 sensors per tree). The trees were selected such that they best represented the distribution of species and stem diameters (Figure 4). From each of the three most abundant species at each plot, at least one tree was sampled. Two species, *Fraxinus ornus* and *Quercus*, were sampled at all four plots because they were present at all plots. Both ring porous (*Quercus* and *Fraxinus ornus*) and diffuse porous species (*Carpinus betulus*, *Juniperus communis* and *Acer campestre*) were sampled.

Effective mean sap flux density and a 95% confidence interval were calculated from the individual sensors weighed by the contribution of each species to the total sapwood area, in the same way as for leaf nitrogen (Eqs. 16 to 18). Latent heat flux, λE (W m^{-2}), was calculated by multiplying sap flux density by the area of sapwood per unit forest floor f_A and by the latent heat of vaporization of water λ (2.5 MJ kg^{-1}):

$$\lambda E = \lambda f_A m \quad (20)$$

The area of sapwood per unit forest floor, f_A , was calculated in the following way. Because the sensors had a length of 20 mm and were inserted in heat-conducting material, it was assumed that the measured sap flux density is the effective sap flux density of the outer 20 mm of the stem. However, for most trees the sapwood area extended

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deeper than 20 mm into the stem. The actual sapwood area was inferred from microscopic analysis of tree cores taken perpendicular to the tree rings of all sampled trees at the end of the growing season of 2004. A film of approximately 0.1 mm thickness was planed off the tree cores with a razor blade and examined visually under a microscope for the presence of active xylem vessels. The majority of xylem vessels were present in the outer 20 mm, and few extended up to 40 mm depth. Because 10 to 30% of active xylem vessels were present at depths greater than 20 mm, we used a value of $25\text{ mm} \pm 10\%$ for the sapwood depths for all trees at all plots. The total sapwood area per unit forest floor was calculated by measuring the diameters of all trees in each plot.

The sun-shade model for light interception was validated using PAR measurements along horizontal transects at the forest floor of the north and the south plot, and along vertical transects at the south plot, at different weather conditions, times of the day and days of the year. Leaf area index L was derived from PAR measurements on completely overcast days.

3 Results

In this section we present the derivation of the parameters for the biochemical model at leaf level from leaf chamber photosynthesis measurements and leaf samples, the validation of the sun-shade model, the model predictions of diurnal cycles of the fluxes. The discussion section is dedicated to a sensitivity analysis of the model.

3.1 Biochemical model

Figure 5 shows leaf chamber photosynthesis measurements used for parametrization of the biochemical model, and model predictions (solid lines). The upper panels show net assimilation as a function of photosynthetically active radiation (PAR), the lower panels C_i/C_a versus PAR. The parameters V_{cm} , R_d and q of the photosynthesis model were calibrated (Table 3) using these data, whereas a-priori values were used for other

parameters (Table 2).

The measurements were carried out at similar vapour pressure deficit (10 hPa) and temperature (28°C). If the hypothesis of Cowan (1977) holds, then C_i should be independent of PAR. However, Fig. 5 shows a sharp increase of C_i/C_a when PAR decreases to low values. This is either caused by a minimum stomatal conductance that prevents the stomata to fully close, or by the fact that the time interval of 2 minutes between the measurements was too short for the stomata to reach an equilibrium. The solid line in Fig. 5 was derived by assuming a relation between assimilation and stomatal conductance as proposed by Leuning (1995).

The values for V_{cm} for the two sampled species (Table 3) were used to calibrate the linear relationship between leaf nitrogen content and V_{cm} . Figure 6 shows leaf nitrogen versus V_{cm} for the two sampled species at the south and the east plot, with 0.95 confidence intervals for both nitrogen content and V_{cm} . We adopt a linear relation of Field and Mooney (1986) to relate V_{cm} to nitrogen content:

$$V = x(N - N_0) \quad (21)$$

where N_0 the residual leaf nitrogen content (=25 mmol m⁻² or 0.5%), and x an empirical coefficient. We calibrated x for the two measurement points (solid line in Figure 6), acknowledging the ambiguousness to use this relationship based on two data points for all plots. However, our data agree with literature data (dots in Fig. 6), calculated from values of maximum photosynthesis by Reich et al. (1999) for different vegetation types in North and South America. Our data are in the same order of magnitude and the relation between leaf nitrogen content and photosynthetic capacity is similar.

Table 4 shows the biochemical parameters as derived from the leaf sample analysis and leaf photosynthesis measurements, and leaf area index for the four plots. For R_d and q , we used equal values for all plots. Differences in nitrogen content between sunlit and shaded leaves were not significant, perhaps due to the open structure of the canopy. It is remarkable that the two plots with the lowest vapour pressure deficit (north and west) have higher carboxylation capacity V_{cm} than the two plots with the

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highest vapour pressure deficit (south and east), and that the two plots with the lowest soil moisture content (south and west) have a higher intrinsic water use efficiency (lower Λ) than the two plots with a higher soil moisture content. This topic will be discussed in detail in a separate study which focusses on the relation between climate and biochemical parameters.

3.2 Sun-shade model

Figure 7 shows the modelled versus the measured fraction of direct light that reaches the forest floor I/I_0 at the north and the south plot. The model prediction is reasonable for the south plot ($r^2=0.77$), and poor for the north plot ($r^2=0.31$), which can be attributed to the high spatial variability of sunfleck distribution at the north plot due to the heterogeneous structure of the forest. The values in the circle correspond to low light conditions in the late afternoon. Figure 8 shows vertical profiles of measured (x) and modelled (line) light distribution in the canopy at the south plot, measured at different times of the day and for different weather conditions. The depth is in units of leaf area index, assuming a homogeneous leaf distribution over depth. From left to right and from top to bottom, the fraction of diffuse ambient irradiance increases from 13 to 100%. The curves are similar in shape, and the model performs well in all conditions except for low solar angles and low light intensity (lower left and lower middle panel). Although the shape of the curves are similar, large differences exist in the irradiance on sunlit and shaded leaves depending on light conditions and time of the day. The insets show calculated values for the fractions of sunlit (open boxes) and shaded (shaded boxes) leaves, and the mean intensities of PAR on sunlit and shaded leaves. With increasing fraction of indirect radiation, the difference in irradiance between sunlit and shaded leaves decreases.

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3.3 Diurnal cycle of assimilation and latent heat

Figure 9 shows the mean diurnal cycles of A , λE and G_s for all 20 clear days between 29 May and 8 July 2004. During this period, the time of sunrise and sunset shifted by approximately 14 min. Because this is smaller than the time resolution of the data (30 min), we did not correct for this. The lines are model predictions, the bars 0.95 confidence intervals for latent heat flux estimated from sap flux measurements (from now on referred to as measured latent heat), and surface conductance calculated from measured latent heat using the inverse Penman-Monteith equation.

The diurnal cycles of assimilation show some remarkable features. The increase of assimilation in the morning and the decrease in the evening are slower at the north and the west than at the south and the east plot. This can be attributed to the lower fraction of sunlit leaves at the north and the west plot than at the south and the east plot, and consequently a greater contribution of leaves that assimilate at a light limited rate, even late in the morning. Although assimilation reaches its peak later at the north and the west plot than at the south and the east plot, the peak values are higher due to a higher value of maximum carboxylation capacity V_{cm} .

The modelled diurnal cycles of latent heat closely match the measured latent heat. This is remarkable, because the measured latent heat is entirely independent of the data we used for deriving model parameters. The biochemical parameters V_{cm} (the maximum carboxylation rate) and Λ (the marginal cost of assimilation) affect latent heat flux in the following way. The north plot has the highest latent heat flux, because it has both a relatively high V_{cm} and a relatively high Λ , i.e. a high photosynthetic capacity and a high marginal cost of assimilation. The west plot also has a relatively high V_{cm} , but a low Λ , and therefore a lower latent heat flux than the north plot. The south and the east plot both have a relatively low V_{cm} , but the east plot has a higher Λ and therefore a higher latent heat flux than the south plot.

The model accurately reproduces the diurnal cycles of surface conductance. The north plot does not show afternoon stomatal closure, whereas the other three plots

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show a typical pattern of stomatal closure in the late morning and afternoon. This can be explained by the combined effects of vapour pressure deficit and the parameter Λ . Low values of Λ (low marginal cost of assimilation) indicate early stomatal closure in response to vapour pressure deficit, whereas high values of Λ indicate that stomata remain open relatively long. At the north plot, stomata remain open because Λ is high and vapour pressure deficit low, whereas on all other plots, Λ is low, or vapour pressure deficits are high, or both. At south plot, Λ is low and vapour pressure deficit high, at the west plot, Λ is low, and at the east plot, vapour pressure deficit high.

In Fig. 10, the data are presented in a different way. This figure shows modelled versus measured latent heat flux for all half-hourly data between 29 May and 8 July 2004. The solid lines are 1:1 lines. For all plots, the squared correlation coefficients are above 0.90. Latent heat flux at the north plot is slightly underestimated, and maximum latent heat flux at the south and east plot overestimated. The significance of this difference between measured and modelled latent heat flux is discussed in the next section using an error propagation analysis of the model.

The model also predicts surface temperature. Modelled surface temperature is not independent of the measurements, because surface temperature was used to calibrate aerodynamic conductance at the south plot. Figure 11 shows for 27 June 2004, the measured and modelled difference between surface temperature and air temperature at the south plot (left panel), and the modelled temperature difference versus the measured temperature difference for all half hourly values between 19 May and 8 July 2004 (right panel). The correlation between modelled and measured surface temperature is lower than that for latent heat. This is a minor problem, because $T_s - T_a$ is relatively small compared to the diurnal cycle of T_a , and the model prediction of latent heat is not very sensitive to errors in $T_s - T_a$.

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4 Discussion

The agreement between modelled and measured fluxes depends on the accuracy of four components: (1) measurements used for validation, (2) input variables, (3) parameter values and (4) the model description itself.

5 The uncertainty in the measurements for validation includes the variation of sap flux density measurements among sensors and the sapwood-surface area ratio. An additional error is that we ignored the time lag that exists between transpiration and sap flux due to storage of water in stems (Schulze et al., 1985). The data indicate that indeed for some time after sunset, sap flow continues. However, there is no reason
10 why storage in stems and the time lag would be equal for all four plots (tree heights vary from 3 m at the east plot to 18 m at the north plot). To account for the time lag requires at least four additional parameters to be estimated (one for each plot). To calibrate these parameters against modelled latent heat would compromise our aim to use independent data for validation of the model. An alternative would be to use radiation
15 data to estimate the time lag, but doing so also creates a dependence between the model and the validation data, because the same radiation data are also used as input in the model. Any other parametrization would be highly subjective. For this reason, we did not account for the time lag.

20 The sensitivity of the model to the most relevant input variables and parameters was calculated with an error propagation analysis, assuming all errors are uncorrelated. The variance of the model prediction σ_y^2 of an output variable y can be calculated from the variances of the variables and parameters x as:

$$\sigma_y^2 = \sum_i \sigma_x^2 \left(\frac{\delta y}{\delta x} \right)^2 \quad (22)$$

25 The greatest uncertainty in the input variables is the vapour pressure deficit, which was measured at 2 m height rather than in or above the canopy. For four weeks at the end of the growing season in 2004, the instrument for temperature and relative humidity

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was moved to just above the canopy at 9 m height at the south plot. A comparison with measurements before and after relocation of the instrument with data of the other plots showed that both temperature and relative humidity are lower above than below the canopy. The net effect was that vapour pressure deficit was <2 kPa higher above than below the canopy. At the north and the west plot, the effect might have been larger because of the taller trees and the denser canopy. In the sensitivity study, we assumed $\sigma_{e_s-e} = 3$ kPa. The greatest uncertainty in the biochemical parameters is the maximum carboxylation capacity V_{cm} . The accuracy of V_{cm} depends on the accuracy of nitrogen measurements and the relation with leaf chamber photosynthesis measurements. Based on the confidence intervals of nitrogen and leaf chamber photosynthesis measurements and the coarse relationship between nitrogen content and V_{cm} , we assumed $\sigma_{V_{cm}} = 10 \mu\text{mol m}^{-2} \text{s}^{-1}$, which is about 20% of the actual values.

Table 5 shows the sensitivity of the mean value of the fluxes of water and carbon dioxide to variations in e_s-e , V_{cm} , Λ and L , absolute and as a percentage of the actual values. In the table we present standard deviations instead of variances. The difference between modelled and measured latent heat flux (Fig. 9) is smaller than the standard deviation of the error of the model prediction, i.e. the difference between modelled and measured latent heat falls within the accuracy of the parameter values and input variables. Thus, there is no reason to improve the accuracy of the physical model description itself. The uncertainty of the vapour pressure alone is insufficient to explain the difference between measured and modelled latent heat flux.

The direct effect of climate and the effect of biochemical parameters on the fluxes can be separated by the following modelling experiment. The north and the south slope have different climate conditions, and also different biochemical parameters. In one experiment, we reverse the meteorological variables at the north and the south plot while leaving the biochemical parameters the same, and in another experiment, we reverse the leaves of the north and the south plot while leaving the weather conditions the same. Figure 12 shows the result of these model experiments for the surface conductance. The bold curve is the reference scenario, as in Fig. 9, the fine line is the

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model where vapour pressure deficit, temperature, incoming radiation and aspect have been reversed, and the dashed line the model where parameters V_{cm} , J_m and Λ have been reversed. Reversing the parameters has a greater effect on surface conductance than reversing the meteorological variables. Reversing the biochemical parameters results in a large change of the magnitude of surface conductance, but a similar shape of the diurnal cycle. Reversing the meteorological variables also changes the shape of the diurnal cycle, especially the hour of the peak.

Wilson et al. (2003) studied the time lag between the peaks of radiation and the fluxes of carbon and latent heat for different climates. The time lag between the peak of radiation and surface conductance we find is for the south plot similar to what they found for Mediterranean forests, and for the north plot similar to what they found for boreal forests. Our results indicate that this difference is mainly caused by the diurnal cycles of radiation, temperature and vapour pressure deficit.

The large effect of the spatial variations of biochemical parameters on the fluxes suggests that models which use uniform biochemical parameters would not predict the observed spatial variability of the fluxes. If average biochemical parameters are used for the north and the south plot, then surface conductance is overestimated at the south plot and underestimated at the north plot.

Although we demonstrated that spatial variations in biochemical parameters are important, we did not address the questions why biochemical parameters vary among the plots the way they do, and whether we can predict spatial patterns. Biochemical parameters are functions of environmental conditions: water potentials in soil and air, the availability of water and nutrients and temporal variations therein, and stand age, succession, pests and diseases and anthropogenic influence. A model to explain biochemical parameters from long term climate will be the subject of a separate study.

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This study showed that both the magnitude and the shape of the diurnal cycle of transpiration and stomatal conductance can be calculated from measurements of leaf nitrogen, ^{13}C isotope discrimination and leaf photosynthesis measurements. The diurnal cycles were more strongly affected by spatial variations in vegetation parameters than by meteorological variables. This indicates that topography induced variations in vegetation parameters are of at least equal importance for the fluxes as topography induced variations in radiation, humidity and temperature.

Appendix A Cowan-Farquhar model after Arneth et al. (2002)

In their Appendix A, Arneth et al. (2002) present a solution of the model of combined model of Farquhar et al. (1980) for photosynthesis, the diffusion equations and the model of Cowan (1977). Combining Eqs. (1), (2), (3) and (4), gives two solutions for C_i : one for enzyme limited photosynthesis, and one for electron limited photosynthesis. In both cases, the equation for C_i is quadratic:

$$k_2 C_i^2 + k_1 C_i + k_0 = 0 \quad (\text{A1})$$

where

$$\begin{aligned} k_2 &= \Lambda + \frac{1.6D}{k' + \Gamma^*} \\ k_1 &= 1.6D - 2C_a\Lambda + \frac{1.6D(\Gamma^* - k')}{k' + \Gamma^*} \\ k_0 &= (\Lambda C_a - 1.6D)C_a + \frac{1.6D\Gamma^*k'}{k' + \Gamma^*} \\ k' &= K_c(1 + O/K_o) \end{aligned} \quad (\text{A2})$$

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for the enzyme limited case and

$$\begin{aligned}
 k_2 &= \Lambda - \frac{1.6D}{3\Gamma^*} \\
 k_1 &= 1.6D - 2C_a\Lambda + \frac{1.6D\Gamma^*}{3\Gamma^*} \\
 k_0 &= (\Lambda C_a - 1.6D)C_a + \frac{1.6D2\Gamma^{*2}}{3\Gamma^*}
 \end{aligned}
 \tag{A3}$$

5 for the electron limited case.

Appendix B Light distribution model

The extinction of both indirect and direct light in a canopy is calculated analogous to light absorption in homogeneous media with the law of Lambert-Beer:

$$\frac{I(l)}{I_0} = \exp^{-\kappa l}
 \tag{B1}$$

10 where I light intensity, l the depth in the canopy in units of L from the top of the canopy and κ an extinction coefficient, which depends on the zenith angle of the light beam θ and the orientation of the leaves. The orientation of the leaves is expressed by the ellipsoidal leaf angle distribution parameter x , where $x < 1$ for mainly vertical leaves, $x = 1$ for a spherical leaf distribution or $x > 1$ for mainly horizontal leaves. The extinction coefficient κ is (Campbell, 1986):

$$\begin{aligned}
 \kappa(x, \theta) &= \frac{\sqrt{x^2 + \tan^2(\theta)^2}}{x + 1 / (2\epsilon_1 x) \ln[(1 + \epsilon_1) / (1 - \epsilon_1)]} & \text{if } x \geq 1 \\
 \kappa(x, \theta) &= \frac{\sqrt{x^2 + \tan^2(\theta)^2}}{x + \arcsin(\epsilon_2) / \epsilon_2} & \text{if } x \leq 1
 \end{aligned}
 \tag{B2}$$

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where $\epsilon_1 = \sqrt{1 - 1/x^2}$ and $\epsilon_2 = \sqrt{1 - x^2}$

The extinction coefficients for indirect and direct light are different, because they origin from different directions. The calculation of the extinction of direct light is straightforward, but the extinction of indirect light must be calculated by integration of Eqs. (B1) and (B2) over the sky area. In practice, this results in an extinction coefficient of 0.7 if $x=1$, independent of geographical location, time of the year or time of the day.

In terrains of steep topography, the penetration of direct light into the canopy is different from that in flat areas. To include the effect of topography, a modified zenith angle is used in Eq. (B2). The coordinate system is rotated such that the surface becomes horizontal, and the zenith angle is calculated for the rotated coordinate system. A sloped surface is described by its steepest angle ϕ and the orientation of the slope ω , which is the horizontal direction of the steepest downward slope, measured clockwise from north. The angle of the surface in the plane of the direct light beam and the vertical ϕ' , is defined as (Fig. 13):

$$\phi' = \arctan(\tan(\phi) \cos(\omega_s - \omega)) \quad (\text{B3})$$

where ω_s the hour angle of the sun. The rotated zenith angle, θ' , is defined as the angle between the vector perpendicular to the slope and the solar beam, i.e.:

$$\theta' = \theta + \phi' \quad (\text{B4})$$

In the calculation of the vertical profile of light in the canopy with Eqs. (B1) and (B2), θ' is used instead of θ . By doing so, it is implicitly assumed that leaf angle distribution x is unaffected by the coordinate rotation. This is an acceptable assumption if leaf angle distribution is spherical, but may not be acceptable if leaf angle distribution is strongly erectophile or planophile. In this study, it is assumed that $x=1$.

It is assumed that leaves are either sunlit or shaded. The effect of a partial eclipse due to the fact that direct radiation does not origin from a point source, or due to light bending over edges of leaves, is ignored. Sunlit leaves receive direct and diffuse light, shaded leaves receive only diffuse light. Scattering and transmission of direct light is

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ignored, and it is assumed that reflected direct light does not meet other leaves on its way back to the atmosphere. The fractions of sunlit leaves f_e , and shaded leaves f_s , are functions of depth in the canopy l in units of leaf area index:

$$f_e(l) = \frac{I_d(l)}{I_{d0}} \quad (\text{B5})$$

$$f_s = 1 - f_e \quad (\text{B6})$$

where I_{d0} the intensity of ambient direct light. The intensities of irradiance on the two fractions are:

$$I_e(l) = I_{d0} + I_i(l) \quad (\text{B7})$$

$$I_s(l) = I_i(l) \quad (\text{B8})$$

where I_{i0} the intensity of ambient indirect light. Total light intensity at depth l in the canopy is:

$$I(l) = f_e I_e(l) + f_s I_s(l) = I_i(l) + I_d(l) \quad (\text{B9})$$

The exposed and shaded fractions and the irradiance on the fractions for the whole canopy are calculated by integrating Eqs. (B5) to (B8) over the leaf area index. In this study, the integration was done numerically using intervals of units leaf area index of 0.1.

Acknowledgements. The authors thank J. de Lange, Ron Lootens, K. de Bruine and H. Visch of the Vrije Universiteit (VU) for developing the sapflow sensors and other equipment, M. Groen (VU) for his technical support, M. Hooyen, S. Verdegaal, M. Konert, H. Vonhof and N. Slimmen (VU) for laboratory analyses, F. Batic of the University of Ljubljana (Lj), R. Aerts and P. van Bodegom (VU) for allowing me to use their equipment, L. Globevnik of the institute for Water in Ljubljana for her administrative support, V. Zupanc and M. Padenik (Lj) for their work in the field, and Peter for climbing the trees.

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Table 1. Characteristics of the four experimental forest plots in the Dragonja catchment.

	north	south	west	east
elevation (m)	180	190	120	150
slope	30°	30°	30°	30°
aspect	330°	210°	270°	210°
plot size (m ²)	625	313	250	100
soil depth (m)	1.0	1.0	1.0	0.8
soil type	clay loam	clay loam	clay loam	clay loam
no of stems ha ⁻¹ (10 ³)	2.3	7.2	3.4	14.4
average diameter (cm)	12.8	7.3	8.6	4.0
average height (m)	16	8	14	4
mid-season LAI	4.0	5.2	4.5	2.7
age (y)	>100	>100	>100	60
management (1900)	wood gathering	cattle grazing	wood gathering	crop field
management (current)	wood gathering	wood gathering	wood gathering	wood gathering

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Table 2. A-priori parameter values for the biochemical model (Farquhar et al., 1980).

K_c (mmol m ⁻³)	K_o (mbar)	O (mbar)	Γ^* (mmol m ⁻³)	J_m/V_{cm}
18.7	330	210	1.22	2.5

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Table 3. Calibrated values of maximum carboxylation capacity V_{cm} , dark respiration rate R_d and quantum yield efficiency q with 0.95-confidence intervals, calibrated for measurements on 13 leaves of *Quercus pubescens* and 6 leaves of *Fraxinus ornus* at the south and the east plot between 14 and 21 July 2004.

	V_{cm} ($\mu\text{mol m}^{-2} \text{s}^{-1}$)	R_d ($\mu\text{mol m}^{-2} \text{s}^{-1}$)	q
<i>Quercus pubescens</i>	54±9	1.0±0.2	0.46±0.15
<i>Fraxinus ornus</i>	44±7	1.1±0.0	0.39±0.09

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Table 4. Mean values of leaf nitrogen concentration [N] and ¹³C isotope discrimination with 95% confidence intervals, maximum carboxylation capacity V_{cm} , dark respiration R_d , marginal cost of assimilation Λ , quantum yield efficiency q , and leaf area index L at the north, south, west and east plot. V_{cm} was derived from leaf nitrogen concentration and leaf chamber measurements, Λ from ¹³C isotope discrimination, q and R_d from leaf chamber measurements and L from PAR measurements.

	north	south	west	east
[N] (g 100 g ⁻¹)	1.61±.08	1.33±0.03	1.74±0.16	1.34±0.04
$\Delta^{13}\text{C}$ (ppm)	21.20±0.24	19.95±0.31	19.76±0.21	20.73±0.22
V_{cm} (μmol m ⁻² s ⁻¹)	68	51	70	52
Λ	1233	622	507	1030
R_d (μmol m ⁻² s ⁻¹)	1.0	1.0	1.0	1.0
q	0.45	0.45	0.45	0.45
L	3.9	4.4	4.2	2.5

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Table 5. Sensitivity of assimilation A and latent heat λE to parameter or variable x with standard deviation σ_x , calculated with Eq 22. Standard deviation are also expressed as percentage of the mean values of assimilation and latent heat.

x	σ_x	σ_A $\mu\text{mol m}^{-2} \text{s}^{-1}$	%	$\sigma_{\lambda E}$ W m^{-2}	%
D	3 hPa	0.30	5	4.0	5
V_{cm}	$10 \mu\text{mol m}^{-2} \text{s}^{-1}$	0.72	12	7.2	10
Λ	100	0.27	4	5.8	8
L	0.5	0.64	11	6.5	9
total		1.05	13	12	11

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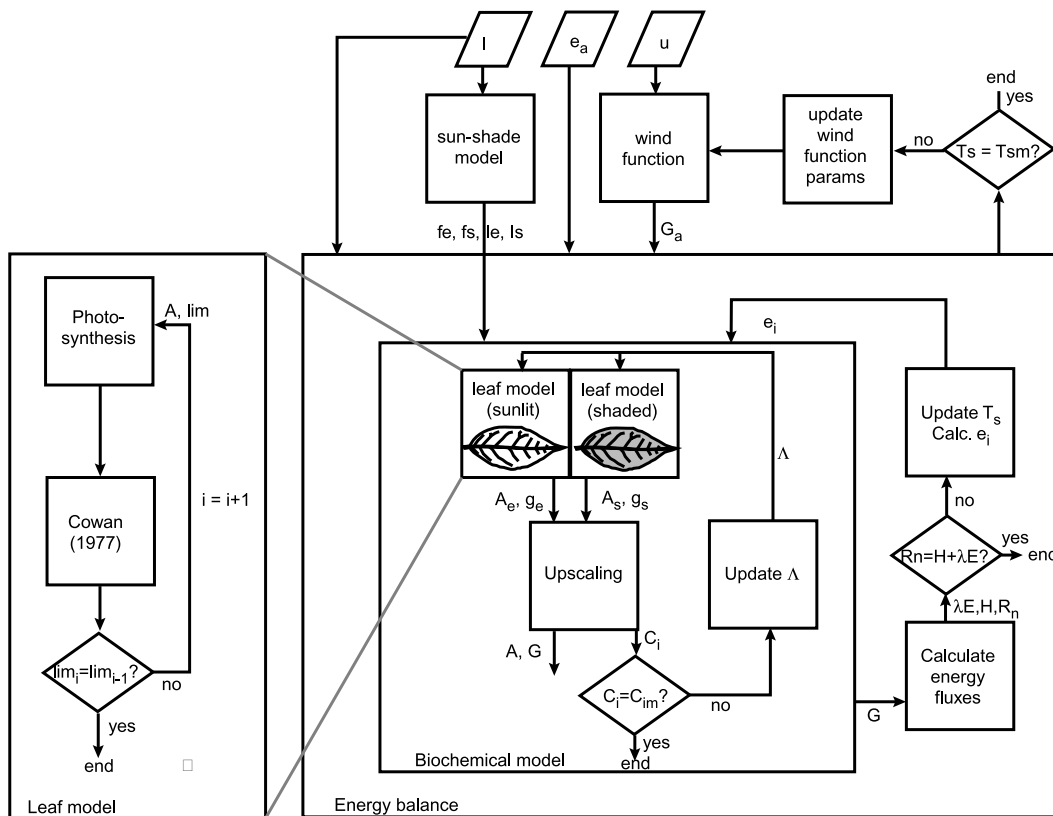


Fig. 1. Flow chart of the combined photosynthesis-transpiration model. The enlargement at the left represents the biochemical model at leaf level.

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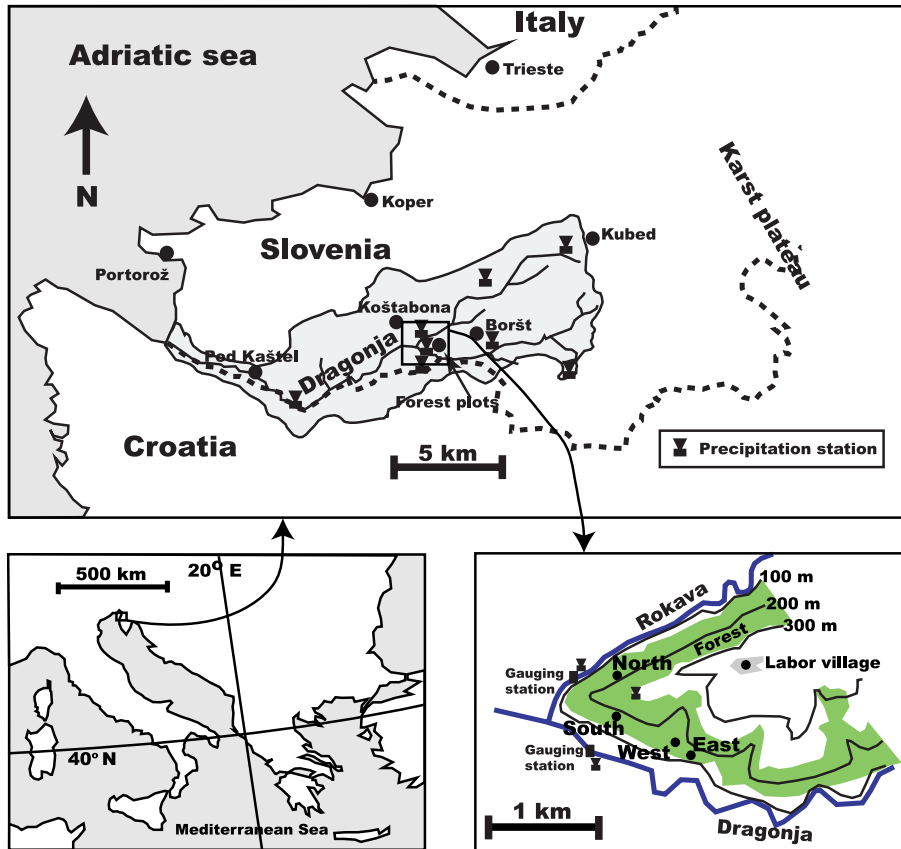


Fig. 2. Map showing the location of the four forest plots and meteorological stations.

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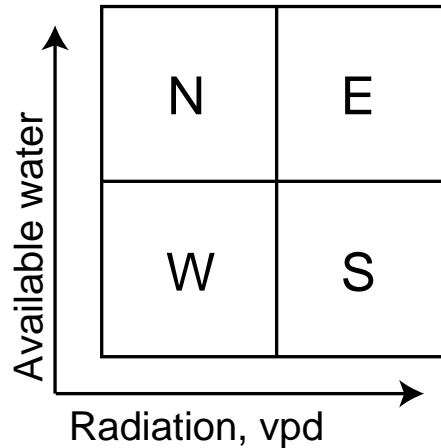


Fig. 3. Schematic representation of vapour pressure deficit and soil water availability the four forest plots: north (N), south (S), west (W) and east (E).

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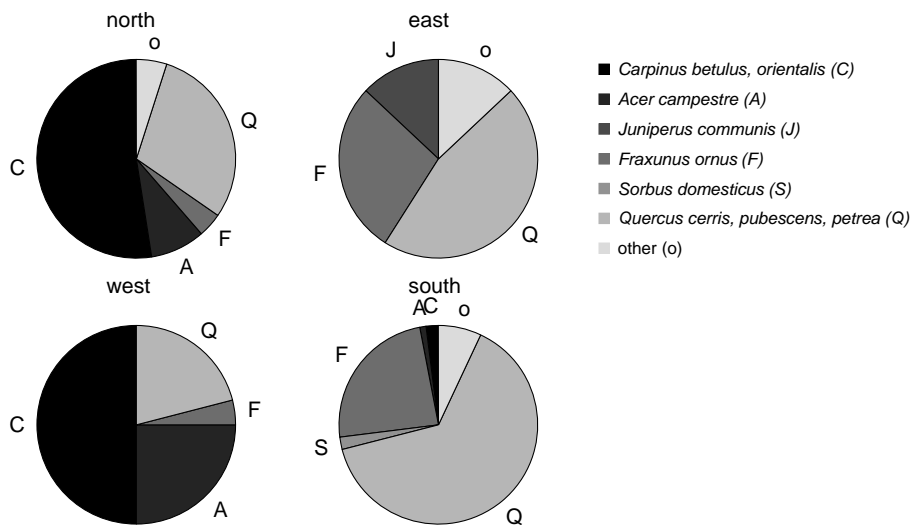


Fig. 4. Distribution of sap wood area over different species at the four forest plots.

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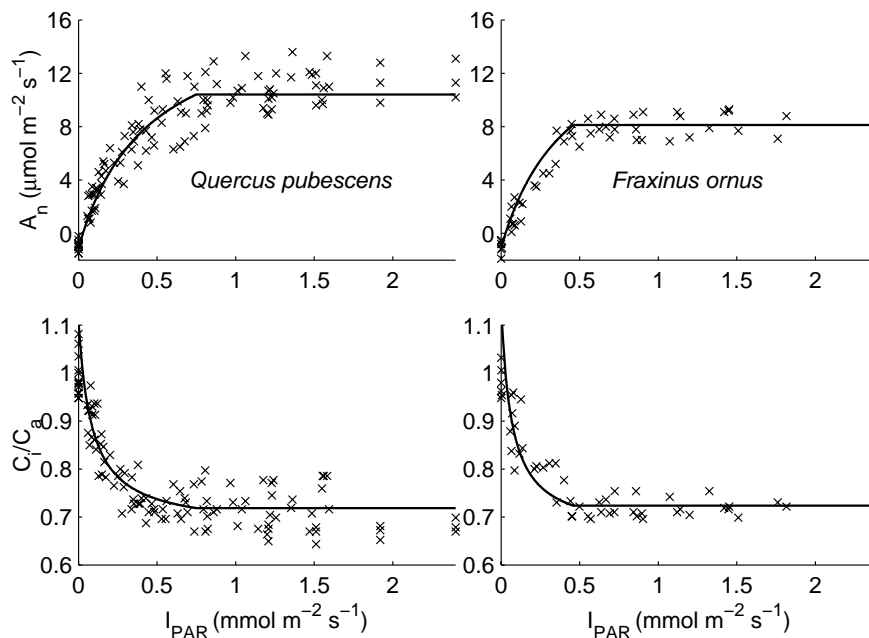


Fig. 5. Rates of net photosynthesis ($\mu\text{mol m}^{-2} \text{s}^{-1}$) (upper panels), and C_i/C_s versus intensity of photosynthetically active radiation PAR ($\text{mmol m}^{-2} \text{s}^{-1}$) (lower panels) for *Quercus pubescens* and *Fraxinus ornus*. Measurements were carried out at the south and the east plot between 14 and 21 July 2004. The solid line is a prediction of the calibrated biochemical model. For the lines in the lower panels, the model of Leuning (1995) was used.

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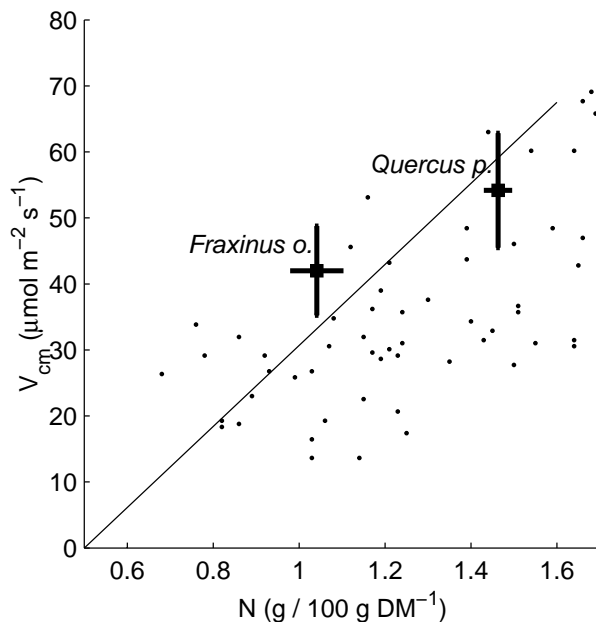


Fig. 6. Mean maximum carboxylation capacity V_{cm} , with 0.95-confidence intervals for *Quercus pubescens* and *Fraxinus ornus*, versus mean leaf nitrogen with 0.95-confidence intervals, measured at the south and the east plot, and a linear regression line through the two data points, forced through $V_{cm}(0.5)=0$. Dots refer measurements by Reich et al. (1999), and were calculated with Eq. (3) from light saturated photosynthesis, assuming $C_i=0.7C_a$ and using the constants of Table 2.

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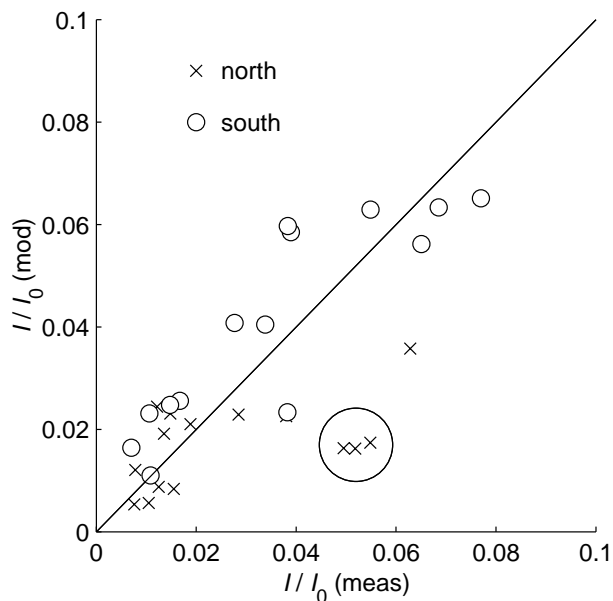


Fig. 7. Modelled versus measured fraction of ambient PAR that reaches the forest floor, for different weather conditions and different times of the day (6 to 19 h) at the north and the south plot, for different days between May and September 2004. The values in the circle refer to low-light conditions in the late afternoon.

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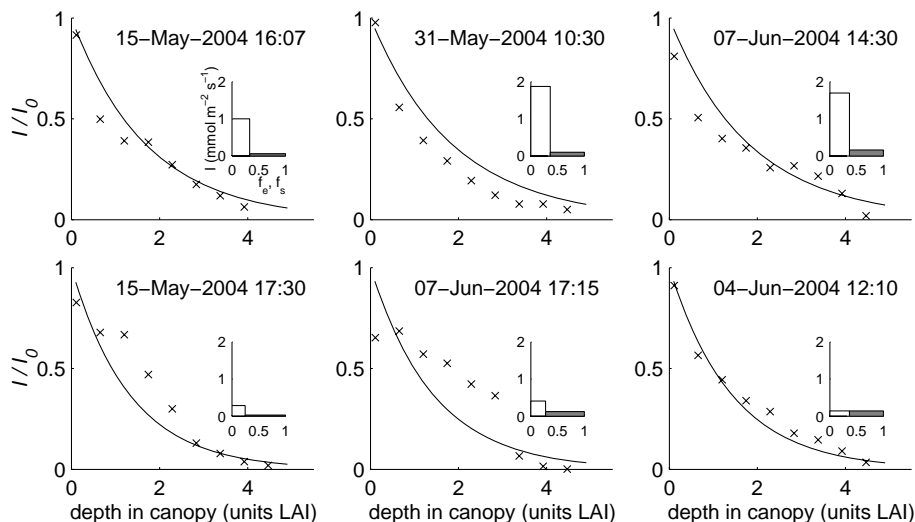


Fig. 8. Measured (x) and modelled (lines) vertical profiles of light intensity relative to ambient light (I/I_0) for different weather conditions at the south plot on different days in May and June 2004. The insets indicate the difference between sunlit and shaded leaves: the open boxes refer to the sunlit fraction, the shaded boxes to the shaded fraction. The width of the boxes denotes the size of the fraction, and the height the intensity of PAR. From left to right and from top to bottom, the fraction of diffuse ambient irradiance increases from 13 to 100%.

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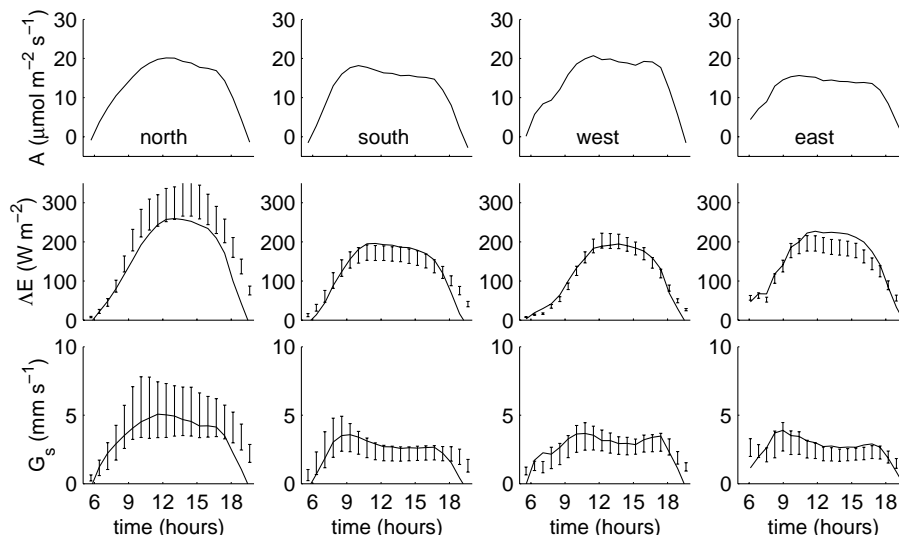


Fig. 9. Modelled rates of net photosynthesis ($\mu\text{mol m}^{-2}\text{s}^{-1}$), latent heat flux λE (W m^{-2}), and surface conductance G_s (mm s^{-1}) at the north, south, west and east plot for 20 clear days between 29 May and 8 July 2004. Lines are model predictions, and bars latent heat flux and surface conductance derived from independent sap flux measurements, and 95% confidence intervals, derived from measurements of 12 sap flux sensors per plot. Bars for surface conductance were derived by inverting the Penman-Monteith equation, using the sap-flux based estimates of latent heat flux.

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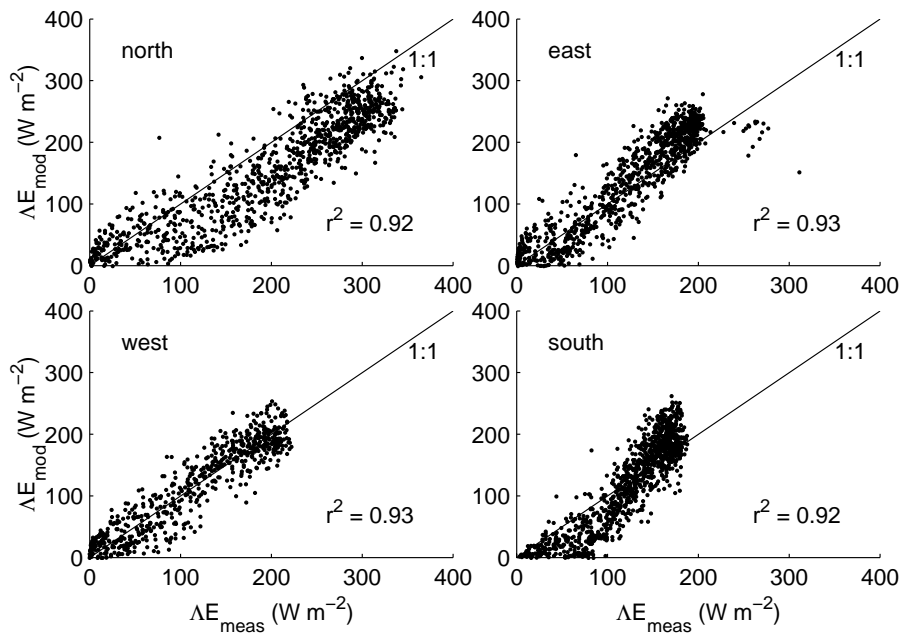


Fig. 10. Modelled versus sap-flux based latent heat flux at the north, south, west and east plot for half hourly data between 29 May and 8 July 2004.

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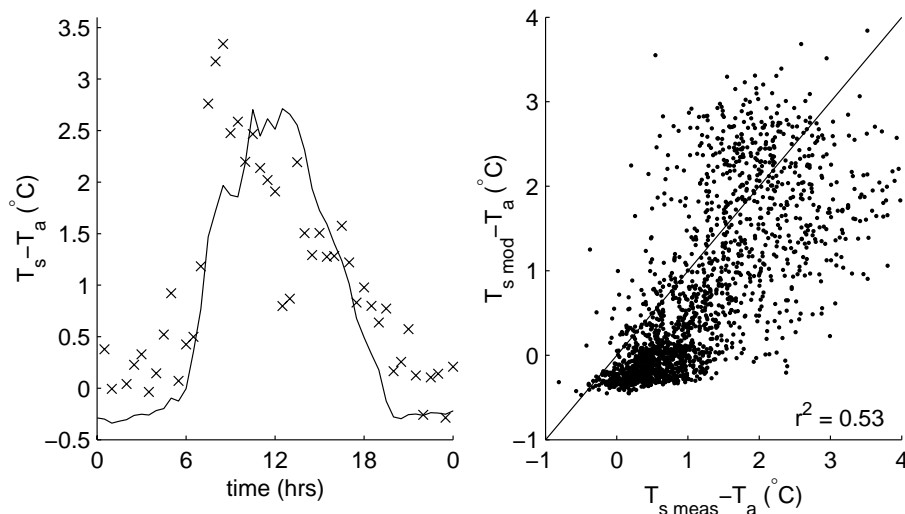


Fig. 11. Diurnal cycle of the measured (x) and modelled (line) difference between surface and air temperature at the south plot on 27 June 2004 (left), and the modelled versus the measured difference between surface and air temperature for half hourly values between 29 May and 8 July 2004 (right).

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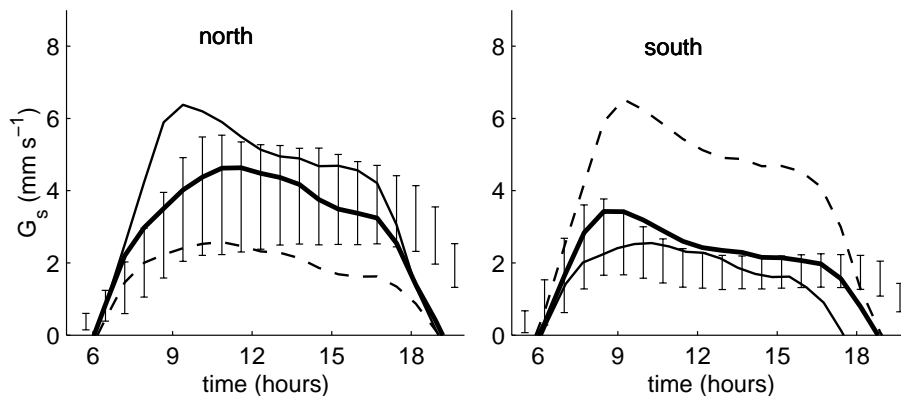


Fig. 12. Modelled and measured surface conductance for the north and the south plot, as in Fig. 9 (bold line), modelled surface conductance after reversing aspect, radiation, temperature and vapour pressure deficit of the north and south plot (fine line), and after reversing V_{cm} , J_m and Λ of the north and south plot (dashed line).

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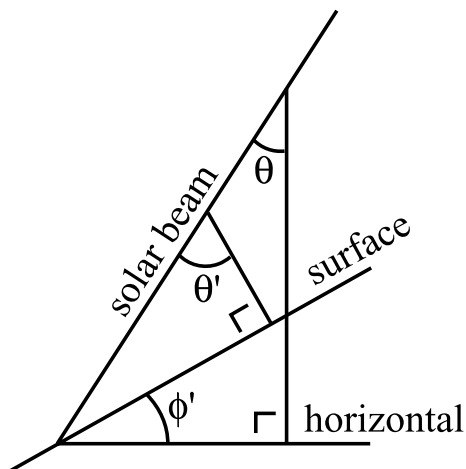


Fig. 13. Definition sketch of the zenith angle θ , the angle of the surface in the plane of the direct light beam ϕ' , and the modified zenith angle θ' for sloped terrain.

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