Interactions among vegetation and ozone, water and nitrogen fluxes in a coastal Mediterranean maquis ecosystem

G. Gerosa\textsuperscript{1}, A. Finco\textsuperscript{1}, S. Mereu\textsuperscript{3}, R. Marzuoli\textsuperscript{2}, and A. Ballarin-Denti\textsuperscript{1}

\textsuperscript{1}Dip.to di Matematica e Fisica, Università Cattolica del S.C., via Musei 41, 25121 Brescia, Italy
\textsuperscript{2}CRINES, via Galilei 2, Curno, Italy
\textsuperscript{3}Dip.to di Biologia Vegetale, Università La Sapienza, via Fermi 1, Roma, Italy

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Correspondence to: G. Gerosa (giacomo.gerosa@unicatt.it)

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Abstract

Ozone, water and energy fluxes were measured over a Mediterranean maquis ecosystem from 5 May until 31 July by means of the eddy covariance technique. Additional measurements of NO\textsubscript{x} fluxes were performed by the aerodynamic gradient technique. Stomatal ozone fluxes were obtained from water fluxes by a Dry Deposition Inferential Method based on a big leaf concept.

The maquis ecosystem acted as a net sink for ozone. The different water availability between late spring and summer was the major cause of the changes observed in stomatal fluxes, which decreased, together with evapotranspiration, when the season became drier.

NO\textsubscript{x} concentrations were significantly dependent on the local meteorology. NO\textsubscript{x} fluxes resulted less intense than the ozone fluxes. However an average upward flux of both NO and NO\textsubscript{2} was measured.

The non-stomatal pathways of ozone deposition were investigated. A correlation of non-stomatal deposition with air humidity and, in a minor way, with NO\textsubscript{2} fluxes was found.

Ozone risk assessment was performed by comparing the exposure and the dose metrics: AOT\textsubscript{40} (Accumulated dose over a threshold of 40 ppb) and AF\textsubscript{st} 1.6 (Accumulated stomatal flux of ozone over a threshold of 1.6 nmol m\textsuperscript{-2} s\textsuperscript{-1}). AOT\textsubscript{40}, both at the measurement height and at canopy height was greater than the Critical Level (5000 ppb\cdot h) adopted by UN-ECE. Also the AF\textsubscript{st} 1.6 value (12.6 mmol m\textsuperscript{-2} PLA, Projected Leaf Area) was higher than the provisional critical dose of 4 mmol m\textsuperscript{-2} PLA. The cumulated dose grew more regularly than the exposure but it showed two different growth rates in the spring and in the summer periods.
1 Introduction

The toxicity of ozone for plants has been widely documented over the past twenty years (Benton et al., 2000; Skärby et al., 1998). Once in the substomatal cavities of the leaves, ozone dissolves in the water film that covers the apoplast and sparks the generation of reactive oxygen species (ROS) which are considered to be responsible for biological damages (Baier et al., 2005; Iriti and Faoro, 2008, Heath, 2008) Among these injuries it is worth reminding the peroxidative destruction of membranes, the oxidation of the thiolic groups of proteins with a consequent alteration of enzymatic activity, the degradation of chlorophylls and the interference with the electron transport systems on which photosynthesis and respiration processes rely upon (Heath, 1996; Reichenauer et al., 1997; Bussotti et al., 2007a). These effects eventually lead to a visible damage on the leaf lamina (Bermejo et al., 2003; Novak et al., 2008; Marzuoli et al., 2008) Even if in most cases the effects remain latent they cause physiological alterations and a general loss in Net Primary Productivity (NPP) (Felzer et al., 2004; King et al., 2005).

Mediterranean ecosystems, because of their climatic conditions and their proximity to anthropic sources of ozone precursors, are among the most exposed ecosystems to this pollutant (Paoletti et al., 2006). The EMEP model estimated (Simpson et al., 2007) for the Mediterranean areas an exposure between 40,000 and 60,000 ppb·h (on a six months basis, April–September) for the year 2000, a value exceeding from 8 to 12 times the critical level of 5000 ppb·h set by UN-ECE for the protection of forests and seminatural vegetation.

Nevertheless field observations never reported a particularly strong plant injury (Bussotti and Gerosa, 2002; Bussotti et al., 2006, Paoletti et al., 2006), thus questioning the soundness of the exposure concept applied to ozone risk assessment. However, the lack of visible injuries in the Mediterranean vegetation could be due to extremely efficient physiological and biochemical defense mechanisms of these plants to oxidative stress as some experiments in controlled environments revealed (Nali et al., 2004; Elvira et al., 2004).
Among the physiological responses, stomatal regulation plays an important role. Following the seasonal decrease of water availability typical of the Mediterranean summer, stomata close to reduce transpiration and water losses, thus limiting the dose of pollutant effectively absorbed and hence its effects. Because of the crucial role of stomata in regulating the dose absorption process, the cumulative ozone flux (AF\textsubscript{st}Y) has been chosen by UN-ECE as a more reliable ozone risk index then AOT40 (Musselmann et al., 2006; Karlsson et al., 2007, Matyssek et al., 2007). This index is based on the accumulation of stomatal fluxes over a threshold (Y) which accounts for the biochemical detoxification mechanisms. This threshold value has been provisionally set to 1.6 nmol m\textsuperscript{-2} s\textsuperscript{-1} (Karlsson et al., 2004) for forests and 6 nmol m\textsuperscript{-2} s\textsuperscript{-1} for crops (Pleijel et al., 2004), however the meaning and usefulness of Y is still debated and discussed in the UN-ECE effects-based community. Despite the acknowledged better biological soundness of AF\textsubscript{st}Y over AOT40 (Matyssek et al., 2007, Karlsson et al., 2007) AOT40 is still widely used since field measurements of AF\textsubscript{st}Y are difficult. In fact direct measurements of ozone flux can be performed only by micrometeorological techniques such as Eddy Covariance (EC) (Keronen et al., 2003), which requires the use of not completely standardized complex equipment. As a consequence, flux based risk assessments is mostly performed with the aid of models, such as the deposition module DO\textsubscript{3}SE (Emberson et al., 2007; Ashmore et al., 2007) included in the EMEP model.

Unfortunately, DO\textsubscript{3}SE has been validated mostly with observations of total O\textsubscript{3} flux and stomatal conductance (gs) in ecosystem types which are representative of the Central and Northern Europe (e.g. Tuovinen et al., 2001). In Mediterranean conditions only one comparative study on wheat (Tuovinen et al., 2004) has been conducted in Italy and, to date, validation and comparison are still missing for an evergreen Mediterranean forest or the Mediterranean maquis. In fact it is important to test the model in conditions where high ozone concentrations can occur with high soil and atmospheric water deficits (Emberson et al., 2005).
This article is aimed to the analysis of the ozone flux dynamics and their interaction with NOx in a water limited environment during the dry season, and to offer a dataset of measurements suitable for model calibration and validation. It is also aimed at assessing the ozone risk for a Mediterranean maquis ecosystem, by the measurement of the effective dose absorbed by the vegetation through stomata and its comparison with AOT40.

2 Materials and methods

Measurements were performed from 5 June to 31 July 2007 in a coastal Mediterranean maquis at Castelporziano, Italy. Details on the measuring site can be found in Fares et al. (2009). Two different techniques were used to measure turbulent fluxes of energy and matter: the eddy covariance technique and the gradient approach. A Dry Deposition Inferential Method approach (Wesely and Hicks, 2000; Gerosa et al., 2003, 2005) was then applied to calculate the ozone stomatal fluxes and the toxicological dose absorbed by the ecosystem.

2.1 Instrumentation

Sensible and latent heat fluxes as well as ozone fluxes were measured using the eddy covariance technique (Swinbank 1951; Hicks and Matt, 1988). An ultrasonic anemometer (USA-1, Metek, Elmshorn, Germany), a CO2/H2O open path fast sensor (LI-7500, LI-COR, Lincoln, Neb., USA) and a fast ozone analyser (COFA, Ecometrics, Italy) were mounted on the top of a 3.8 m tall scaffold.

One net radiometer (NR lite, Kipp&Zonen, Holland), one PAR meter (190SA, LI-COR, Lincoln, Neb., USA) and a temperature and relative humidity probe (50Y, Campbell Scientific, Shepshed, United Kingdom) were placed at the same height than the anemometer.

An additional reference O3 analyzer (SIR S-5014, DASIBIE, Spain), sampling air at the top of the scaffold near the fast ozone sensor, was also used.
NO and NO\textsubscript{2} fluxes were measured by a NO\textsubscript{x} analyzer (SIR 2308, DASIBIE, Spain) using the gradient approach, measuring the nitrogen oxides concentrations alternatively at two different heights by means of an electro-valve switching system controlled by a computer with a LabView (National Instruments, Austin, Tx, USA) software. The two sampling points were chosen at 3.8 m and 1.3 m.

The measuring site was equipped with additional instrumentation to better describe the temperature and humidity profile, the soil water status, the energy fluxes and the microclimate of the area: two additional temperature and relative humidity probes (50Y, Campbell Scientific, Shepshed, United Kingdom) at 1 m and 0.1 m; three soil heat flux plates, (HFP01SC, Hukseflux, Delft, Holland); three TDR reflectometers (C616, Campbell Scientific, Shepshed, United Kingdom); three leaf temperature probes (Pt100, DeltaT, United Kingdom); two surrogate leaves (237, Campbell Scientific, United Kingdom) to measure leaf wetness, one rain gauge (52202, Young/Campbell Scientific, Cambridge, United Kingdom) and one barometer (PTB101B,Vaisala, Finland).

Fast sensors were sampled at 20 Hz by a computer with a customised software written in Delphi 5.0 (Borland). Slow sensors were sampled every 15 s and data were collected by a datalogger (CR10x, Campbell Scientific, Shepshed, United Kingdom) equipped with a signal multiplexer device (AM16/32, Campbell Sci., United Kingdom), and data were averaged every 30 min.

2.2 Eddy-covariance

Eddy covariance is a turbulence based technique which states that fluxes are equal to the covariance between the vertical component of the wind (\( w \)) and the measured scalar quantity. Originally developed by Swinbank (1951), it has been accurately described and widely used for gas exchange measurements (e.g. Stull, 1988; Kaimal and Finnigan, 1994; Foken, 2008).

Under some condition that have to be fulfilled (stationarity of the variables for which vertical fluxes are calculated, horizontal homogeneity, absence of chemical sources and sinks between the measuring height and the exchanging surface, and average
vertical wind component equal to zero; Grunhage et al., 2000) vertical fluxes are constant with height and ozone, sensible and latent heat fluxes can be calculated as follows

\[ F_{O_3} = \overline{w'C'} \]  

\[ H = \rho c_p \overline{w'T'} \]  

\[ \lambda E = \lambda \rho \overline{w'q'} \]

where \( C \) is the ozone concentration, \( T \) the air temperature, \( q \) the specific humidity of the air, \( \rho \) the air density, \( \lambda \) the constant of water vaporization, and \( c_p \) the specific heat of the air. The primes (’) indicate fluctuations of each variable around their mean and the overbars represent averages over a chosen time period, in our case a 30 min averaging period. This period is short enough to separate synoptic and diurnal variations from the turbulent data (van der Hoven, 1957), and long enough to include all turbulent fluctuations occurring in the atmospheric surface layer.

At the end of each 30 min averaging period, the fluctuations around the means were calculated after linear detrending of the data series and a covariance matrix was calculated, i.e. the covariances between every considered parameter and each other. Then the covariance matrix was rotated following the three rotation suggested by McMillen (1988) in order to eliminate the advective components resulting from small non-homogeneities of the exchanging surface and an eventual slight vertical tilt of the instrumentation.

In order to ensure a perfect synchronization of the data series and to account for different instrumental delays, the data series of the variables acquired by fast sensors were lagged with successive steps of 0.05 s, with respect to the wind data series, until the calculated fluxes reached their maximum value. When a maximum flux value was not found within the 60th lag, the sample was recognized as not stationary and discarded.
2.2.1 Data selection

In addition to the previous method, the fulfillment of the stationarity requirement has been checked by using the selection criterion proposed by Dutaur et al. (1999). This criterion requires that fluctuations are calculated in two different ways: as the difference of the linear detrended series with the 30-min average, and as the difference with an instantaneous local running mean obtained by passing a mathematical R-C recursive filter (analogous of an electric circuit of a resistor and a capacitor in series) over the original time series.

If the normalized absolute difference between the covariances calculated with the fluctuations obtained with the two methods is below 1, the sample was considered as stationary and reliable. On the contrary, the sample was discarded.

Other data selection criteria were that the data capturing efficiency of each sample had to be greater than 85%, and that the canopy had to be completely dry. Only the data that passed these selections were used for successive analysis.

2.2.2 Dry Deposition Inferential Method (resistive analysis)

The deposition of a gas depends on many variables such as wind velocity, friction velocity, incoming radiation, temperature, surface type, etc. In the Dry Deposition Inferential Method approach (DDIM) the deposition surface is treated as a “big leaf” located at a height \( d + z_0 \) over the soil, where \( d \) is the displacement height accounting for the canopy height \( h \) (set to 2/3 of the \( h \)) and \( z_0 \) is the roughness length which accounts for the canopy roughness. Three main phases are considered in the deposition process: first of all the gas must overcome the aerodynamic resistance \( (R_a) \) existing in the turbulent layers of the atmosphere above the studied surface; then the gas must move across the quasi-laminar sub-layer which is characterized by a molecular diffusion against the so-called sub-laminar resistance \( (R_b) \); finally, in order to reach the surface, the gas must overcome the resistance of the surface itself \( (R_c) \). The deposition flux of a gas is hence considered as the analogous of a current flowing through a...
electric circuit composed by these three resistances in series. Their equivalent resistance is called total resistance ($R_{tot}$) and it is equal to the concentration of the gas at the measuring point divided by the flux:

$$R_{tot} = R_a + R_b + R_c = C_{zm}/F_{O_3}$$ (4)

The aerodynamic resistance $R_a$ was calculated using the well known similarity relation introduced by Monin and Obukhov (1954), while the sub-laminar resistance $R_b$ for ozone was calculated following the general purpose parameterization proposed by Hicks et al. (1987). The surface resistance $R_c$ is hence obtained as a residual since $R_{tot}$ is known, because it is derived from directly measured entities ($C_{zm}$ and $F_{O_3}$).

In order to estimate the fraction of gas penetrating through the stomata of plants, the surface resistance $R_c$ (also called the canopy resistance in ecology) is broken down into a stomatal resistance and a non-stomatal one mounted in parallel.

The stomatal resistance to ozone, $R_{ST}$, has been calculated from the stomatal resistance to water evaporation $R_w$ by inverting the Penmann-Monteith equation (Monteith, 1981) – i.e. by solving this equation for $R_w$ since all the other entities are directly known from measurements – and by considering the relative diffusivity ratio of ozone in air to that of water vapour.

The non-stomatal resistance was calculated as a residual from $R_c$ and $R_{ST}$, following the rules of the parallel resistances.

The stomatal ozone flux was hence obtained as

$$F_{ST} = \frac{R_c}{(R_a + R_b + R_c) R_{ST}} C_{zm}$$ (5)

Further details can be found in Gerosa et al. (2005).

The inferred resistances that showed values exceeding 10 000 s m$^{-1}$ were rejected, because of being unrealistic and causing numerical problems when used. The same happened for samples where $R_c$ resulted lower than 0 or $R_{ST}$ was not computable because the Penman-Monteith equation could not be inverted.
2.2.3 Ozone dose and exposure

In this work both exposure and dose approaches were compared. The exposure was calculated as AOT40 for daylight hours only:

\[
AOT40 = \sum_{\forall \text{GlobRad} \geq 50 \text{W/m}^2} \max(0; C_{d+z_0} - 40) \Delta t
\]

(6)

where \( \Delta t \) is the averaging period for the ozone concentration measurements (1 h). The concentration at \( d+z_0 \), recommended by the ICP modelling and mapping manual (2004), was calculated with the Dry Deposition Inferential Method (Gerosa et al., 2005):

\[
C_{d+z_0} = C_{zm}(1 - R_a / R_{tot})
\]

The ozone dose received by the ecosystem in the whole measuring period (May - July) was calculated as \( AF_{st0} \) by summing up all the 30-min ozone stomatal fluxes \( F_{st} \) of the period

\[
AF_{st0} = \sum \max(0; F_{st}) \Delta t
\]

(7)

where \( \Delta t \) is the averaging period chosen for eddy covariance measurements.

The dose was also calculated as \( AF_{st1.6} \):

\[
AF_{st1.6} = \sum \max(0; F_{st} - 1.6) \Delta t
\]

(8)

where \( F_{st} \) is the stomatal ozone flux obtained by the DDIM, 1.6 nmol m\(^{-2}\) s\(^{-1}\) is the UN-ECE detoxification threshold and \( \Delta t \) is the averaging period of the flux measurements.

2.2.4 Data selection and data gap-filling

The results presented in this paper rely on the measured data that fulfilled all the selection criteria. One unique exception has been made for the assessment of the exposure and the dose.
In fact, since both AOT40 and the ozone dose are cumulative metrics, a gap in ozone concentrations and ozone fluxes will result in an unavoidable underestimation of their values.

To reduce such underestimation a gap-filling was performed. Gaps of no more than 3 consecutive 30-min averages were linearly interpolated, while large gaps were gap-filled with time series reconstructed by multiple linear regression based on available predictors as local meteorological parameters and ozone concentrations from a nearby measuring station.

Gaps in stomatal fluxes could result also as a consequence from data rejection at the output of the DDIM process. In these cases stomatal fluxes were estimated by taking the average stomatal fraction (the ratio between \( F_{st} \) and \( F_{tot} \), based solely on measured data) at the corresponding half an hour, and by multiplying it by the available total ozone flux.

2.3 Gradient approach

The aerodynamic gradient has been widely used to estimate surface fluxes (Grunhage et al., 2000; Foken, 2008) since, unlike eddy covariance, it does not require fast analyzers. Here the turbulent diffusion coefficient for heat \( K_{H} \), that takes into account also the vertical stability of the atmosphere, was easily available from eddy covariance data and applied to both nitric oxide and nitrogen dioxide fluxes, following the similarity of the transport of each scalar entity found by Monin and Obhukhov (1954) (Stull, 1988; Monteith and Unsworth, 1990; Pal Arya, 1988). Nitrogen oxides fluxes were hence calculated for each semi-hour as follows:

\[
F_{NO_x} = -K_{H} \cdot \Delta[NO_x]/\Delta z
\]

where \( NO_x \) indicates alternatively NO and \( NO_2 \), \( \Delta[NO_x] \) is the mean difference of NO and \( NO_2 \) concentrations between the two measuring heights \( z_2 \) and \( z_1 \), and \( \Delta z \) is \( z_2 - z_1 \), equal to 2.5 m in our case.
3 Results

A total of 4176 semi-hourly samples were gathered during the whole measuring period. The samples referred to completely dry canopy conditions were 1894 (45.3%), because nearly all the samples between 08:00 p.m. and 08:30 a.m. were excluded due to the presence of dew on the leaves.

This sub-dataset was further reduced by 16% because instrumental drawbacks (e.g. shut down, instrument breakings and substitutions), and by another 33.2% following the exclusion of the data which did not fulfill the stationarity conditions.

Hence the output of the DDIM analysis is based on the 50.8% of the data gathered from 08:30 a.m. to 08:00 p.m., except when explicitly indicated.

In order to highlight different meteorological and physiological traits, most of the results are split in two different periods: the first one from 5 May to 12 June (late spring) and the second one from 13 June to 31 July (summer).

3.1 Ozone concentrations and fluxes

The average ozone concentration at the measuring height was 32.6 ppb in the first period and 38.6 ppb in the second one. The overall average was 35.9 ppb and the maximum peak concentration was 107 ppb. Ozone concentrations were strongly influenced by the local meteorology: higher concentrations were observed when the wind was blowing from the sea and lower concentrations downwind to the city of Rome. Ozone concentrations usually began to increase early in the morning, around 08:00 a.m. reaching their daily maximum in the first afternoon, and decreased to the lower nighttime values afterwards (Fig. 1). With the only exception of some days with high nighttime values, ozone concentrations showed a typical bell-shaped behavior.

On the contrary, the total ozone fluxes appear more irregular because of their intrinsic link to the atmospheric turbulence. Substantial differences in the daily maximum absolute values (Fig. 1), which are usually reached in the first hours of the afternoon, are evident. A significant reduction of the average total ozone fluxes occurred in the
second period: in the first period total fluxes were between 15 and 20 nmol m\(^{-2}\) s\(^{-1}\) for a large part of the day (08:00 a.m. to 07:00 p.m.), while in the second period they were around 10 to 12 nmol m\(^{-2}\) s\(^{-1}\) (Fig. 2). Also nighttime values of the total fluxes were slightly greater in the first period (about 7 nmol m\(^{-2}\) s\(^{-1}\)) than the second one (about 5 nmol m\(^{-2}\) s\(^{-1}\)).

The stomatal component of the ozone flux decreased as a consequence of stomatal response to a decreased water availability in the soil and an increased VPD in air (Mereu et al., 2009).

Figure 3 highlights the reduction in evapotranspiration that follows the decrease in soil water content (SWC hereafter). In the late spring the ecosystem could rely on a relatively high SWC, especially from the lower soil strata. In summer, the ecosystem experienced dryer conditions, since water depletion involved the deeper soil layers and the water table itself became shallower. Consequently, an increasing fraction of the available energy was thermically dissipated, while the fraction used for evapotranspiration decreased by more than 60%, from 105 W m\(^{-2}\) to 45 W m\(^{-2}\) in the central hours of the second period (Fig. 4).

Also stomatal fluxes decreased their absolute diurnal mean value by 44% between the first and the second period. Moreover their diurnal behavior changed too, as described in Fig. 2. In the first period ozone stomatal uptake showed a slight increase from the morning hours to the first hours of the afternoon (from 5 to 8 nmol m\(^{-2}\) s\(^{-1}\)), and afterwards it decreased back to 5 nmol m\(^{-2}\) s\(^{-1}\) in the evening. In the second period stomatal fluxes were about 4 nmol m\(^{-2}\) s\(^{-1}\) around 09:00 a.m. and slightly decreased to a constant value reached at midday (around 2.5 nmol m\(^{-2}\) s\(^{-1}\)), then remained unvaried until sunset.

These stomatal fluxes were compared with the ozone stomatal uptake derived from sap flow measurements performed simultaneously on three species at the same site (Mereu et al., 2009). In Fig. 5 the sap flow uptake is obtained by summing up each species uptake, upscaled by the species specific LAI and percentage cover as reported by (Fares et al., 2009). The ozone flux attributed to these species shows a similar trend...
to that estimated from EC. In the first period, fluxes rapidly increased in the morning hours from values of less than 1 nmol m$^{-2}$ s$^{-1}$ to values of about 7 nmol m$^{-2}$ s$^{-1}$ around 11:30 a.m. and gradually decrease afterwards. In the second period, the morning increment was significantly lower and fluxes reached a value of 2.5 nmol m$^{-2}$ s$^{-1}$ already at 10:30 and remained steady for most of the day.

As a consequence of the different water availability, the stomatal fraction of the ozone flux absorbed by vegetation varied from a range of 40 to 50% of the total flux in the first period, to a range of 20–30% in the second period. In the latter case, it is interesting to note that the stomatal fraction was higher in the morning (45% around 09:00 a.m.), it rapidly decreased to a lower percentage for most of the day (Fig. 6 and slightly recovered to a 30% in the late afternoon.

3.2 Ozone exposure and dose

During the whole experimental period the ecosystem experienced an ozone exposure, expressed in terms of AOT40 for daylight hours of 20 650 ppb·h (Fig. 7). Such a value is computed using ozone concentrations at $z_m$=3.8 m a.s.l., but if it is computed with the concentrations at top canopy height, i.e. at $d + z_0$, total exposure lowers to 8600 ppb·h.

In the same period the stomatal dose, computed as a cumulative stomatal flux without any cutting threshold ($AF_{st0}$), was 22.8 mmol m$^{-2}$ PLA.

The exposure at $z_m$ during the measuring period shows an irregular growth which reflects the alternation of photochemical episodes with meteorological perturbations. The exposure at $d + z_0$, after the initial increment between 20 May and 25 May, grew more regularly due to the lower ozone concentrations at leaf level.

Dose development ($AF_{st0}$), instead, was less variable, but a more careful analysis reveals two distinct periods with two different ozone assimilation rates, periods corresponding to late spring and summer (Fig. 7). In both periods the dose grew almost linearly, at a rate of 0.17 mmol m$^{-2}$ day$^{-1}$ in the first period and of 0.11 mmol m$^{-2}$ day$^{-1}$ in the second. The two different rates clearly reflect the lower stomatal response in the
second period. Also $AF_{st} \ 1.6$ showed a similar trait, but with a lower growth rate during summer. The $AF_{st} \ 1.6$ dose at the end of the measured period was 12.6 mmol m$^{-2}$ PLA, about the half of the $AF_{st} \ 0$ (55.3%).

3.3 Nitrogen oxides concentrations and fluxes

Mean nitric oxide (NO) concentrations remained fairly constant in the central hours of the day around a value of 3–4 ppb (Fig. 8a). Concentrations slowly increased during the night reaching a peak value of 20 ppb around 05:00 a.m., after which they decreased back to the typical diurnal values.

Nitrogen dioxide (NO$_2$) concentrations remained around 1–2 ppb during the diurnal hours and rose to 6 ppb after sunset. Similarly to NO, also NO$_2$ concentrations showed a morning peak of 10 ppb which occurred between 08:00 and 09:00 a.m.

Vertical fluxes of nitrogen oxides were considerably weaker than the already described ozone fluxes and were in general more variable. Nevertheless, the mean day trend (Fig. 8b) reveals some regularities. During the measuring period NO showed a median positive flux (exiting the ecosystem) during the day (between 08:00 and 18:00 a.m.) and an almost null flux in the remaining night hours. The peak value of the median flux was about 0.2 ppb m s$^{-1}$ ($\pm$0.28 std. dev.)

On the contrary, the NO$_2$ flux shows an almost constant median deposition rate around 0.06 ppb m s$^{-1}$ at night, with a weak peak of about 0.1 ppb m s$^{-1}$ ($\pm$0.19) at 08:00 a.m. followed by an intense diurnal emission peak of 0.6 ppb m s$^{-1}$ ($\pm$0.39 std. dev.) around 01:00 p.m.

The net NO efflux could be the result of soil emissions below the vegetation, while the efflux of NO$_2$ may be interpreted as a photochemical effect. As refereed by Gao et al. (1993) and Monson and Holland (2001), NO concentration show a steep decrease with height above the soil under a canopy, reaching a minimum near the middle of the canopy, while the O$_3$ induced oxidation of NO to O$_3$ within the canopy causes NO$_2$ concentrations to be higher within the canopy than above resulting in a net efflux of
NO$_2$. The origin of such imbalance is not completely clear and could be attributed to advection, deposition and transformation by nighttime chemistry of nitrogen species transported from Rome by the city plume.

4 Discussion

The order of magnitude of the ozone dose received by the maquis ecosystem is comparable to the dose received by the Holm Oak forest positioned just 0.8 km inland (Gerosa et al., 2008a) and it is also similar to the amount absorbed by barley and wheat crops (Gerosa et al., 2003, 2005). Apparently, the dose does not vary greatly despite the evident structural differences of these ecosystems and the different leaf and canopy stomatal conductance. In all these cases, only a small portion of the ozone received by the ecosystem is effectively absorbed by the stomata. The greatest part of ozone, instead, is depleted in chemical-physical processes that altogether are termed non-stomatal deposition.

4.1 Stomatal uptake

It is important to notice that night time hours are frequently characterized by thermal inversions, which determine the accumulation of dew over the canopy and soil (Cieslik et al., 2009; Mereu et al., 2009). The formation of dew between 02:00 and 06:00 a.m., as well as the immediate drying of the leaf lamina and the evaporation of dew immediately after 06:00 a.m., are confirmed by the leaf wetness sensors (Fig. 9). When this event takes place, the morning flux due to evaporation from the stand surfaces could be erroneously attributed solely to a stomatal activity. It is for this reason that the presented results were filtered in order to exclude fluxes measured when the canopy was wet or in the following hour. However it cannot be excluded that the evaporation of dew hides a prominent stomatal opening (Fig. 10). In fact, light intensity is sufficient to cue stomatal aperture already from the first hours of the morning (PPFD at 07:00 a.m. was
around 300 $\mu$E m$^{-2}$ s$^{-1}$) and it could be that after the evaporation of dew and as the VPD increases, guard cells adjust in order to reduce gs to the lower values of the day.

However, the sap flux measurements performed on three species, (Mereu et al., 2009) contradict this hypothesis. With the only exception of the driest period (late July), sap flux showed a bell shaped trend and a peak in sap flux never occurred in the morning (Fig. 11). However, the measured species only represent 38% of the total cover (Fares et al., 2009), hence if a morning transpiration peak is really taking place, it could be attributed to the other species. But this eventuality has to be excluded by the fact that also the unmeasured species ($Q. ilex$, $P. latifolia$, $C. incanus$) are known to behave in a similar way for the transpiration rate (Bombelli and Gratani, 2003). The only exception could be Rosmarinus officinalis (17% of the cover), but the SWC ($\Psi_s < -1$ MPa) was low enough to ensure a reduction of its stomatal conductance of more than 80% (Clary et al., 2004). It is hence most likely that the observed morning peak is an artifact attributable simply to dew evaporation.

This underline the importance of verifying dry canopy conditions in EC based flux measurements, even in xeric environments, and the potential advantage of coupling EC measurements with sap flow gauges.

4.2 Non-stomatal deposition

The nature of the non-stomatal deposition ($F_{nstrom}$), more than 50% percent in this study, is still not understood and different hypothesis have been made to explain it. Van Pul and Jacobs (1994), after observing that the $F_{nstrom}$ increased with the efficiency of ozone transport inside the canopy, suggested the cause to be the destruction of ozone over the plant and soil surfaces, since they observed that the ozone transport efficiency is proportional to $u^*$ and inversely related to the Leaf Area Index (LAI) and to the vegetation height. Instead, in a laboratory experiment Rondón (1993) showed that cuticular uptake by leaf waxes increased from a negligible rate at low levels of light intensity, to rates comparable with stomatal uptake at light levels equivalent to strong sunlight conditions, thus suggesting an important role of solar radiation. Fowler
et al. (2001) suggested that the relationship between $F_{\text{nstrom}}$ and radiation reported by Rondón (1993) and by Coe et al. (1995) could be explained as thermal decomposition of the ozone molecules intercepting the surfaces heated by radiation; they also estimated the energy of activation of this reaction to be 36 KJ mol$^{-1}$. Kurpius and Goldstein (2003), instead, observed that the exponential form of the relationship between $F_{\text{nstrom}}$ and temperature was similar to the relationship between temperature and the emission rate of Volatile Organic Compounds (VOC). Based on this similarity, they advanced the hypothesis that a great fraction of the daytime ozone deposition could be a consequence of gas-phase reactions with biogenic hydrocarbons and estimated that 45 to 55% of the total ozone flux could depend on such reactions. However Mikkelsen et al. (2000) estimated from concurrently measured $\alpha$- and $\beta$-pinene fluxes a maximum destruction potential of monoterpene emissions corresponding only to 10% of the total ozone flux.

A role in non-stomatal ozone depletion has been also attributed to gas-phase tri-tration reactions with biogenic emissions of NO. The measurements of Dorsey et al. (2004) and of Pilegaard et al. (1999), and the precedent models of Duyzer et al. (1995) and Walton et al. (1997), highlighted that NO emissions from forest soils not only affect the magnitude and the direction of the NO$_2$ fluxes, but also the intensity of ozone fluxes. Nevertheless, the influence on the latter is usually weak, but not negligible, and can become relevant in case of a high soil NO efflux (Pilegaard et al., 1999), when a substantial ($\sim$30%) amount of the total ozone flux could be accounted for by chemical reactions, especially at night (Pilegaard, 2001; Walton et al., 1997). Finally, Altimir et al. (2004) found an influence of the atmospheric humidity on non-stomatal deposition at night, with an hyperbolic increase of surface conductance to ozone with saturating RH %.

In this study, the analysis of the data presented in Fig. 12 allow to support only the hypothesis of a direct influence of atmospheric humidity on non-stomatal deposition. A gas-phase reaction with NO of biogenic and antropic origin could be observed but only at high efflux rates as reported by Pilegaard et al. (1999).
The occurrence of this reaction is revealed by the hyperbolic dependence ($R^2=0.95$) of $F_{\text{nstom}}$ on the NO$_2$ fluxes observed over the canopy (Fig. 12f).

The dependence on RH, instead, is revealed by the median values of non-stomatal conductance, that, increased linearly together with the absolute humidity of air ($R^2=0.63$) when the canopy was completely dry (Fig. 12d). When the canopy was wet (mostly at night in our site) the relationship became hyperbolic ($R^2=0.60$) (Fig. 13) as already found by Altimir et al. (2004). In this last case, ozone is not only removed by reactions with atmospheric water, but also dissolved in the water film deposited over the colder surfaces, i.e. leaves, stems and soil. The chemical process of ozone decomposition and reaction with air water vapour and droplets has been described by Seinfeld and Pandis (1997) and de Paula and Atkins (2006). Nevertheless, the proximity of the coastline could bring to suppose a contribution of marine aerosols which can modify the chemical properties of the water films and consequently enhance ozone solubility and its removal. Recent findings on the chemistry of halogen compounds (Br, I, Cl) in marine boundary layers MBL (Monks, 2005), suggest a possible role of these gas compounds in the removal of ozone in our coastal site. In fact, bromine and chlorine in the MBL are emitted through the production of sea salt aerosol, with a smaller fraction released from biogenic organic halogens. Iodine compounds on the other hand are largely released as organic compounds and molecular iodine from micro and macro algae that accumulate iodine from the sea water (von Glasow, 2008). Chemical and photochemical reactions rapidly convert these compounds in halogenated monoxides (Monks, 2005). It is these latter compounds that are directly responsible for the catalytic removal through one principal and simple mechanism (von Glasow et al., 2002):

$$XO + YO \rightarrow X + Y + O_2 \quad \text{with } X \text{ and } Y = \text{Br, I, Cl}$$  \hspace{1cm} (10)

$$X + O_3 \rightarrow XO + O_2$$ \hspace{1cm} (11)

Model calculations have indicated that the presence of only 0.5–4 pmol mol$^{-1}$ BrO can significantly impact the O$_3$ budget in the MBL (von Glasow et al., 2002; von Glasow and Crutzen, 2004).
Recently, BrO and IO measurements in Green Cape, not only confirmed the occurrence of such mechanism but also brought to the conclusion that the ozone photochemistry is largely dominated by halogen chemistry (Read et al., 2008). Interestingly, the daily profile reported for BrO and IO concentrations is similar to the ozone deposition found in this study. Concentrations of both gasses reach a maximum around 10:00 a.m., remain constant until 14:00 p.m. for IO and until 07:00 p.m. for BrO, and decrease to 0 afterwards. In the coastal site subject to a considerable external source of NOx at night, as in this case, the formation of halogenated nitrates – as BrONO2 – can take place. At sunrise, these compounds photolyse freeing NO2 and halogenated oxides that may trigger and amplify the catalytic destruction of ozone. Such a process is what is suggested by the so-called “sunrise ozone destruction” reported by different authors (Nagao et al., 1999, Galbally et al., 2000, Watanabe et al., 2005). Such a morning peak of ozone depletion was already reported for a close site (Gerosa et al., 2005, 2008) and it occurred again during this campaign. The morning peak, especially in the first period, can be inferred from Figs. 2 and 6, where a rise in total ozone deposition is not supported by a concomitant rise in ozone stomatal uptake. This hypothesis would also explain the NO2 efflux in excess observed in the maquis during the day. In fact, the air masses of the city plume rich in nitrate species, would first be transported offshore from the night breeze and return to land during the day under the form of organic and halogenated nitrous compounds. Hence, if the hypothesis of the role of the chemistry of halogenated compounds should be confirmed also for this site, the correlation found with humidity could simply be an indicator of the transport of halogenated species from the sea when the wind was blowing from offshore.

4.3 Ozone risk assessment

Ozone exposure was calculated as AOT40 using ozone concentrations both at the measuring point and at canopy level, calculating the latter ones by means of the DDIM. In both cases the ozone exposure exceeded the critical level (CL) of 5000 ppb·h established for plants protection, revealing a potentially ozone hazard condition for the
maquis ecosystem. Considering that the vegetative period of maquis is much longer than our measuring period, it can be reasonably supposed that the exposure to which these ecosystems are usually subject to is very high.

It is worth noticing that the CL was exceeded very soon by the AOT40 evaluated at the measuring height (23 May), and only one month and half later (5 July) at the canopy height \( d + z_0 \). The two metrics gave very different results, highlighting the importance of following the indications of the Mapping Manual (ICP Modelling and Mapping, 2004), the disrespect of which may bring to considerable overestimation of the risks – a three fold higher in this case – and to erroneous conclusions. The risk of overestimating the negative effects of ozone on the vegetation is particularly high in the Mediterranean area, where the concentrations of this pollutant are usually high (Paoletti, 2006).

The failure to estimate ozone concentrations at top canopy height is not necessarily due to negligence, but to the lack of the necessary information to infer the ozone gradient above the canopy when using data from monitoring network stations, which usually sample at a height of 3m. The determination of such gradient, in fact, requires knowledge of the aerodynamic state of the atmospheric surface layer (turbulence/stability) and the conductance to ozone of the ecosystem, which is known to vary rapidly as a response to the environmental conditions and to the physiological state of the plant. Ultimately, ozone exposure at top canopy height can be determined only through direct flux measurements. Measurements that, as in this case, can be more profitably used to determine the stomatal ozone uptake, the only toxicological significant parameter.

The dose of ozone absorbed by the vegetation during the measuring period, appears well above the provisional critical flux level of 4 mmol m\(^{-2}\), expressed as \( A F_{st} 1.6 \), for the reduction of 5% of biomass growing in beech and birch (ICP Modelling and Mapping, 2004; Karlsson et al., 2004)

This dose, however, is below the critical dose for the appearance of visible injury symptoms on leaves (30 mmol m\(^{-2}\)) of beech and poplar found in recent OTC experiments (Gerosa et al., 2008b). These experiments reported also that even at lower doses, even asymptomatic species such as \( Q. \ robur \) showed a marked reduction in
the photosynthetic efficiency (Bussotti et al., 2007). So, even if leaf injuries were not reported in this study, it cannot be excluded that photosynthetic assimilation was negatively influenced by ozone and ozone activated antioxidant systems capable of protecting the vegetation from photo-oxidative stress (Nali et al., 2004; Paoletti, 2006).

In any case, in a multi specific ecosystem the ozone risk assessment might to be more precise and take into account the specific physiology of each specie. Some species, in fact, can largely account for the dose of ozone absorbed by the ecosystem and hence may be relatively more effected by this pollutant and trigger future changes in ecosystem composition. This is what happened, for example, for the three species A. unedo, Q. ilex, E. arborea considered in drawing the Fig. 4.

Finally, the measured dose is very close to the 24 mmol m$^{-2}$ estimated for generic forests, in the same area and for the year 2000, by the renovated deposition module DO$_3$SE-EMEP (Simpson et al., 2007). But it should be noted that the model estimation covered 6 months of the entire growing season and not only the central three months, as in our case.

In any case model parameterizations for this ecosystem are still necessary.

5 Conclusions

The maquis ecosystem acted as a net sink for ozone and the ozone deposition was quite high. Nevertheless, only a minor part of the ozone flux (32.8%) was absorbed by vegetation through the stomata. The stomatal uptake was influenced by water availability and decreased throughout the measuring period as the season became dryer.

The remaining part of the ozone deposition, the so called non-stomatal one, was positively influenced by air humidity and by nitrogen oxides. Nevertheless the influence of these latter was weak, and was evident only when nitrogen fluxes were particularly high. No influence with other measured factors, such as temperature, solar radiation and turbulence intensity were found. Moreover, due to the coastal location of the measuring site, a possible role on ozone depletion of halogenated species avvected from
the sea was also suggested as a working hypothesis, even if it should be confirmed by new measurements.

The maquis ecosystem resulted at high risk for ozone considering both AOT40 and $AF_{st}$1.6 approaches, just on a three months base instead of six months as suggested by UN-ECE. Hence, negative effects on the vegetation community cannot be excluded for this particular ecosystem, and more in general for the Mediterranean maquis ecosystems. However the different behaviors shown by the exposure and the dose metrics highlighted the need for field measurements in order to realize a better risk assessment.

Finally a significant dataset is now available for testing, refining and validating deposition models on this type of ecosystems and Mediterranean climatic conditions

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Fig. 1. Ozone concentration and ozone fluxes. The dark line is the total ozone flux to the ecosystem and the gray line is the stomatal flux (left axis), i.e. the amount of ozone taken up by vegetation through stomata. Circles represent ozone concentrations (right axis) and dashed dark and gray lines are gap-filled values.
Fig. 2. Mean daily course of total and stomatal ozone fluxes. (a) Spring period (6 May–12 June). (b) Summer period (13 June–31 July). Deposition fluxes are here indicated with positive values. $F_{\text{tot}}$ and $F_{\text{stom}}$ are the total and the stomatal ozone fluxes when the canopy were completely dry, i.e. excluding the periods where dew was found on the leaves; $F_{\text{totDEW}}$ and $F_{\text{stomDEW}}$ are the same fluxes but obtained including also the periods where canopy were wet. Vertical bars are the standard deviations.
Fig. 3. Daily averages of evapotranspiration, rain and soil water content (v/v) during the whole measuring period. Soil water was measured at three different levels: 30 cm, 60 cm, 100 cm. Vertical dotted line is the separation between the two periods.
Fig. 4. Energy fluxes: available energy (net-radiation), sensible heat and latent heat. Mean daily course in (a) the spring period and (b) the summer period. Vertical bars are the standard deviations.
**Fig. 5.** Comparison of the ozone uptake (stomatal fluxes) obtained from the sap flow measurements and the eddy covariance (EC) measurements in the late spring and summer period.
Fig. 6. Average stomatal fraction in the two periods (a) and (b). $F_{\text{stom}}/F_{\text{tot}}$ is the ratio of the stomatal flux to the total flux when the canopy were completely dry, i.e. excluding the periods where dew was found on the leaves; $F_{\text{stom}}/F_{\text{totDEW}}$ is the same ratio but obtained including also the periods where canopy were wet. Vertical bars are the standard deviations.
Fig. 7. Evolution of Ozone exposure and stomatal dose during the measuring period. The exposure is expressed as AOT40 calculated at measurement height and at canopy height (d+z0). The dose is expressed as cumulated stomatal flux $AF_{st\,0}$, without any flux threshold, and as $AF_{st\,1.6}$ after the application of the instantaneous flux threshold of 1.6 nmol m$^{-2}$ s$^{-1}$. 
Fig. 8. (a) Mean daily course of NO and NO\textsubscript{2} concentrations; (b) Median daily course of vertical fluxes. Positive values are efflux and negative values are depositions. Vertical bars are the standard deviations.
Fig. 9. Canopy wetness as measured by surrogate leaves. Frequency of canopy wetness in the different hours of (a) first and (b) second period.
Fig. 10. Stomatal conductance of the ecosystem canopy. Median daily course in (a) the spring period and (b) the summer period. $G_{\text{stom}}$ is the stomatal conductance to ozone when the canopy were completely dry, i.e. excluding the periods where dew was found on the leaves; $G_{\text{stomDEW}}$ is the same conductance but obtained including also the periods where canopy were wet. Vertical bars are the standard deviations.
Fig. 11. Average days of sap flow for three species (A. unedo, Q. ilex, E. arborea). (a) mid May–12 June (b) 13 June–end July (data from Mereu et al., 2009). Vertical bars are the standard deviations.
Fig. 12. Non-stomatal deposition ($G_{\text{nonStom}}$) versus air turbulence ($u^*$), air temperature ($T$), solar radiation (PPFD) and absolute air humidity ($\chi$), NO and NO$_2$ fluxes ($F_{\text{NO}}$ and $F_{\text{NO}_2}$). Points are median values of non-stomatal conductance for each bin of the variable in x-axis and vertical bars represent standard deviations. Note than the y-axis in the $F_{\text{NO}_2}$ graph has a different scale. Positive flux values indicate efflux and negative values depositions.
Fig. 13. Non-stomatal deposition and relative air humidity in (a) dry canopy and (b) wet canopy conditions. Vertical bars are the standard deviations.