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Seasonal cycles of biogenic volatile organic compound fluxes and concentrations in a California citrus orchard

S. Fares^{1,2}, J.-H. Park¹, D. R. Gentner³, R. Weber¹, E. Ormeño^{1,4}, J. Karlik⁵, and A. H. Goldstein^{1,3}

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Correspondence to: S. Fares (silvano.fares@entecra.it)

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¹University of California, Berkeley, Division of Ecosystem Sciences, Department of Environmental Science, Policy, and Management, USA

²CRA (Agricultural Research Council) – Research Center for the Soil-Plant System, Rome, Italy

University of California, Berkeley, Department of Civil and Environmental Engineering, USA
 Aix-Marseille University – Institut méditerranéen de biodiversité et écologie IMBE CNRS UMR7263, France

⁵University of California Cooperative Extension, Kern County, USA

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Orange trees are widely cultivated in Mediterranean climatic regions where they are an important agricultural crop. Citrus have been characterized as emitters of volatile organic compounds (VOC) in chamber studies under controlled environmental conditions, but an extensive characterization at field scale has never been performed using modern measurement methods, and is particularly needed considering the complex interactions between the orchards and the polluted atmosphere in which Citrus is often cultivated. For one year, in a Valencia orange orchard in Exeter, California, we measured fluxes using PTRMS (Proton Transfer Reaction Mass Spectrometer) and eddy covariance for the most abundant VOC typically emitted from citrus vegetation: methanol, acetone, and isoprenoids. Concentration gradients of additional oxygenated and aromatic compounds from the ground level to above the canopy were also measured. In order to characterize concentrations of speciated biogenic VOC (BVOC) in leaves, we analyzed leaf content by GC-MS (Gas Chromatography-Mass Spectrometery) regularly throughout the year. We also characterize in more detail concentrations of speciated BVOC in the air above the orchard by in-situ GC-MS during a few weeks in spring flowering and summer periods. Here we report concentrations and fluxes of the main VOC species emitted by the orchard, discuss how fluxes measured in the field relate to previous studies made with plant enclosures, and describe how VOC content in leaves and emissions change during the year in response to phenological and environmental parameters. The orchard was a source of monoterpenes and oxygenated VOC. The highest emissions were observed during the springtime flowering period, with midday fluxes above 2 nmol m⁻² s⁻¹ for methanol and up to 1 nmol m⁻² s⁻¹ for acetone and monoterpenes. During hot summer days emissions were not as high as we expected considering the known dependence of biogenic emissions on temperature. We provide evidence that thickening of leaf cuticle wax content limited gaseous emissions during the summer.

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Introduction

Oranges, Citrus sinensis L., are among the most economically important and widely cultivated crops in areas with Mediterranean climates, such as California, Italy, Spain, Morocco, and Israel, and areas of cultivation are often close to densely populated areas. All vascular plants including Citrus species emit biogenic volatile organic compounds (BVOC), with a global estimate of BVOC emitted from plants in the range of 1–1.5 Pg Cy⁻¹ (Guenther et al., 1995). In the presence of sunlight and nitrogen oxides (NO_x), the oxidation of BVOC can lead to formation of tropospheric ozone (Chameides et al., 1988; Papiez et al., 2009), a greenhouse gas with detrimental effects on plant health, crop yields, and human health (for a reference list, see EPA, 2011). BVOC are also precursors to atmospheric aerosol formation (Henze and Seinfeld, 2006), accounting for a significant fraction of secondary organic aerosol (SOA) produced in the atmosphere (Goldstein and Galbally, 2007).

The Mediterranean climates where citrus is cultivated are characterized by high summer temperatures. Emission of BVOC from Citrus is known to be temperature dependent due to volatilization from organs (ducts, glands) where BVOC are stored (Kesselmeier and Staudt, 1999), as shown in recent studies using branch enclosure techniques (Fares et al., 2011). In these studies, performed under controlled conditions and in the absence of any environmental stress, monoterpenes were the most abundant isoprenoids emitted by oranges. Sesquiterpenes were another important class of isoprenoids whose emissions depended primarily on temperature, but they are formed by different biosynthetic pathways than monoterpenes (for a review see Duhl et al., 2008). In addition to these reactive isoprenoids, oranges were shown to emit oxygenated volatile organic compounds (OVOC), which from biogenic sources are products of catabolism and depend mostly on temperature, but also on light conditions. Among OVOC, methanol is a plant volatile emitted to the atmosphere in large quantities from the demethylation of pectins in cell walls with global emissions estimated at 100-240 Tgy⁻¹ (Galbally and Kirstine, 2002; Jacob et al., 2005; Millet et al., 2008).

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Acetone and acetaldehyde are also important primary OVOC emitted from terrestrial ecosystems and oceans, but these are also produced in the atmosphere in large quantity as secondary compounds from oxidation of hydrocarbons of both anthropogenic and biogenic origin (Goldstein and Schade, 2000). The contribution of cultivated Cit-5 rus to VOC emissions in polluted or highly populated regions is poorly understood, but may be signficant considering that these compounds could notably influence the ozone-forming potential of the atmosphere, affect concentrations of HO_x and peroxyacetyl nitrates, and contribute to the formation of organic aerosol (Singh et al., 2001; Steiner and Goldstein, 2007).

The Central Valley of California is a region with both extensive agriculture and anthropogenic pollution. Its southern half, the San Joaquin Valley, experiences pollution from large nearby cities (e.g. Fresno, Bakersfield, Modesto, Stockton), as well as inflow of pollution from populated coastal regions (e.g. the San Francisco Bay area). We chose an experimental site in the southern half of the San Joaquin Valley where citrus. especially oranges and mandarins, are extensively cultivated. Here we report results of a one-year study at that site, specifically: (1) to quantify concentrations and fluxes of the main BVOC species "in-situ" using proton transfer reaction mass spectrometry (PTRMS) and the eddy covariance technique, along with in-situ gas chromatography; (2) to determine how fluxes measured in the field relate to previous studies made with plant enclosures; (3) to investigate how BVOC content in leaves and emissions change during the year in response to phenological and environmental parameters.

Material and methods

Site description

The measurement site was located in a private Valencia Orange orchard three km west of the University of California Lindcove Research and Extension Center near Visalia, California, USA. For a more detailed description of the site location, soil and plant **ACPD**

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characteristics, spacing of the plantation, irrigation treatments please refer to Fares et al. (2012). The site is characterized by a Mediterranean climate, with warm dry summers and cold wet winters, and a typical wind pattern which brings daytime air up the mountain slopes of the Sierra Nevada Mountains from the nearby urban area of Visalia, while at night a gentle downslope wind reverses the direction. For the measurement period, precipitation, daily and hourly averaged data of air temperature, photosynthetic active radiation (PAR) and turbulence (u^*) are shown in detail in Fares et al. (2012).

Measurements started in October 2009, and ended in November 2010. A shipping container was fitted with air conditioning and electrical wiring to serve as a field lab, and set in the orchard in September, 2009, to house analytical instruments in a temperature controlled environment. Sensors and inlet lines were attached to a telescoping tower, (Floatograph FM50 telescoping mast), 9.8 m tall, and located 5 m away from the field lab, on the same tree-line. The tower was equipped with meteorological sensors replicated at four measuring heights (9.18, 4.85, 3.76, 1 m). The height-replicated measurements included air temperature, relative humidity (RH), and wind speed. Other measured environmental variables included photosynthetically active radiation (PAR) (Li-Cor Inc., Lincoln, NE) and soil moisture (Campbell Scientific Inc., Logan, UT). A system of fine wire thermocouples was used to measure leaf temperatures (Omega Engineering, Precision Fine Wire thermocouples). All data were recorded at 1 min intervals using dataloggers (CR10x and CR3000, Campbell Scientific Inc., Logan, UT).

To be able to relate flux measurements to citrus leaf mass and leaf area, we harvested a "Valencia" orange tree from within the study block in August 2010. We measured five composite samples of fresh leaves with a Licor leaf area meter (mod. LI-3100 C), then dried the samples. All leaves were removed from the citrus tree, dried, and weighed. Leaf area for the whole tree was calculated using the mass-to-area conversion obtained from the leaf samples as applied to total leaf mass for the tree. We measured the number of trees per ha using Google Earth imagery and ground measurements. The leaf area index (LAI) for the orchard was 3.00 m⁻² leaves m⁻² soil. **ACPD**

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The average height of the trees surrounding the field lab and tower was 4.2 ± 0.23 m (n = 11).

PTRMS system for flux and gradient measurement

VOC mixing ratios were measured in situ by Proton-Transfer-Reaction Mass Spectrometer (PTR-MS), which has been described elsewhere in detail (Lindinger et al., 1998). During each hour air was sampled through five individual gas inlets made of Teflon with 4 mm internal diameter, each of which was protected by a Teflon filter (PFA holder, PTFE membrane, pore size 2 µm) 30 cm from the inlet. The filters were replaced every 2 weeks, a time interval considered adequate to avoid contamination or flow problems based on past research (Holzinger et al., 2005). One inlet was used to sample air at 4.85 m from 0 to 30 min for eddy-covariance flux measurements of methanol, acetone, isoprene, monoterpenes and an oxidation product, with m/z of 33, 59, 69, 81, 113, respectively. The measurement cycle duration for these 5 masses including water was 1.1 s. The sampling tube was 15 m long and heated at a constant temperature of 40 °C to avoid condensation inside the tubing. A sample flow of 10 lmin⁻¹ was generated with a diaphragm pump and maintained by a mass flow controller (MKS Instruments). Four additional inlets were used to sample vertical gradients at height-levels within (1.m, 3.76 m) and above (4.85 m and 9.18 m) the canopy sequentially for 6 min each during the second 30 min of each hour. In order to avoid different retention times of the air in the inlet lines, we used tubing with the same length for each inlet line (20 m). Table 1 lists the m/z monitored, the corresponding compounds, and the dwell time for each mass. Each sampling line was connected to a 3-way solenoid valve (TEQCOM Industries) controlled by a datalogger (mod. CR10x, Campbell Sci.). Air was continuously pulled through each sampling line to avoid memory effects of the air retained in the lines.

The instrument sampled from the main sampling line at 0.41min⁻¹ and was optimised to an E/N ratio of 128Td using a drift tube pressure, temperature, and voltage of 2.02 hPa, 45 °C, and 600 V, respectively. The reaction time was 100 µs and the

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count rate of H₃O⁺H₂O ions was less than 3% of the count rate of H₃O⁺ ions, which was $\sim 5 \times 10^6$ counts s⁻¹. Each measurement cycle lasted ~ 2 min, totalling 13 measured cycles per level for each hour. The first two cycles were discarded to prevent eventual error due to missed synchronization between the PTRMS and the datalogger clocks. The instrumental background signal was measured by directing the sample flow through a catalyst-based purifier for the first 3 min before starting the measurements in the second half of each hour, similar to Holzinger et al. (2005). The purifier consisted of a stainless steel tube filled with platinum-coated quartz wool (Shimadzu) heated to 350°C, which efficiently removed the VOC but not the water vapor from the sample. This is important because the background impurities may depend on the humidity of the sampled air.

Gas samples from gravimetrically-prepared standard cylinders (Apel and Riemer) of pure nitrogen with known mixing ratios (4-5 ppm) of methanol, acetaldehyde, acetone, isoprene, methyl-vinyl-ketone, benzene, hexenal, d-limonene, Δ-3-carene were automatically measured twice a day (at hours 02:00 and 16:00) by dynamically diluting with purified air to obtain concentrations in the range of 10-50 ppb, which were similar to those expected in the atmosphere. The count signal was then transformed to ppb after subtracting the averaged background levels and taking into account the measured sensitivities for each calibrated compound (i.e. counts/ppb, Davison et al., 2009). For concentrations of the other masses for which authentic standards were not available, we calculated normalized sensitivities (counts/concentration) based on theoretical proton transfer reaction rate coefficients and the instrument-specific transmission coefficient calculated from a transmission curve. This curve was determined at an array of masses from 33 to 219 m/z using our gas standards at concentrations of 100 ppb (Apel and Riemer).

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Hourly-resolved VOC concentrations were measured using an automated in-situ gas chromatograph (Agilent mod. 5890) equipped with a mass-selective detector (Agilent mod. 5971). The instrument was operated in-situ with a custom system that automated sample collection and analysis. Ambient samples were collected for the first 30 min of every hour via an inlet located at a height of 4 m. Ozone and particulate matter was removed at the inlet using 47 mm glass fiber filters (Pall, type A/E) that were coated in sodium thiosulfate according to the method vetted by Pollmann et al. (2005). After ozone and particulate removal, the sample travelled down a 1/4" heated Silcosteel line at ~1 Imin⁻¹ to a preconcentration system, where a ~600 ml sample was concentrated on a custom-made adsorbent trap (glass beads: Tenax TA: Carbopak B: Carbopak X held in place by glass wool on each end) and thermally desorbed onto a DB-624 capillary column (60 m × 0.32 mm × 1.8 µm), and then analyzed by the MSD. Daily calibration checks and blank runs were performed using gas-phase standards and a zero air generator. Calibrations were performed using gas-phase monoterpene standards and liquid standards for more reactive compounds (e.g. sesquiterpenes and unstable monoterpenes).

2.4 BVOC flux calculation

Wind velocity and sonic virtual temperature fluctuations were measured at $10\,\text{Hz}$ with a three-dimensional sonic anemometer (Applied Technologies, Inc., Boulder, CO) mounted on a horizontal beam with the air sample inlet attached inside the anemometer structure. The wind data were rotated according to the planar fit method (Wilczak et al., 2001). The lag interval between the instantaneous vertical wind velocity and the BVOC concentration measurement varied due to changes in clock synchronization between the PTRMS clock and the datalogger where sonic data were stored. To calculate this lag time, for each specific 30-min measurement period, vertical wind velocity and concentration were correlated in a $\pm 10\,\text{s}$ time window following the principle of the

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Fluxes of BVOC (F_c , nmolm⁻² s⁻¹, Eq. 1) were calculated using the continuous flow Disjunct Eddy Covariance method (Karl et al., 2002) in which fluxes are calculated from a subsample of the horizontal wind data corresponding to data collected with the PTRMS after subtracting the lag time (Δt):

$$F_{\rm c} = \frac{\sigma}{N} \sum_{i=1}^{N} w'(i - \Delta t/\Delta t_{\rm w})c'(i) \tag{1}$$

where σ is the air density (mol m⁻³), $w' = w - \overline{w}$ is the instantaneous deviation of the vertical wind speed (w) from its average, $c' = c - \overline{c}$ is that of the BVOC concentration (nmol mol⁻¹), $\Delta t_{\rm w}$ is the sampling interval in the wind measurements (0.1 s), and N is the number of PTR-MS measurement cycles (1680) during the flux averaging time (29.5 min).

A de-spiking routine was applied to exclude points clearly resulting from interferences. We considered outlier points where the difference from the averaged signal during the half hour was 10 times higher than the theoretical standard deviation (Bamberger et al., 2010). Measured fluxes were multiplied by a frequency response corrections factor compensating for the high frequency attenuation caused by the response time of the PTR-MS calculated using the formula and parameterization suggested by Horst (1997). We found that mid-day high-frequency attenuation was about 0.90. Flux values were discarded if at least one of the following conditions were met: (1) measured ambient concentration was close to the detection limit of the specific VOC. (2) Results from the stationary test for the various BVOC were above 60 % (Foken and Wichura, 1966). (3) the footprint area was outside the boundaries of the orchard (Hsieh et al., 2000). (4) turbulence was low ($u^* < 0.15$), a very frequent occurrence at our measuring site during night hours.

The uncertainty was measured according to the method proposed by Wienhold et al. (1994) determining the signal noise of the covariance by calculating the time

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shifts between vertical wind velocity and BVOC concentration far beyond the true lagtime. We used the same procedure recently adopted by Ruuskanen et al. (2011) using a lag window of 40 s and rated the 30-min flux data in three classes: (1) good quality, (2) low quality, (3) poor quality, as reported in Table 1. All data processing was computed using a Matlab routine.

For easier comprehension we grouped the data to obtain a continuous time series from day of year (DOY) 1 to DOY 365 by placing 2009 fall observations into the figures after the winter through summer measurements collected in 2010.

2.5 Analysis of BVOC concentrations in leaves and cuticle waxes

Storage of both leaf BVOC and waxes, which occurs in secretory cells and cuticles respectively, were extracted by suspending leaf dry matter (1 g) in hexane in a ratio of 1:5 w/v containing 0.0123 mg ml⁻¹ of dodecane (internal standard for quantitation) for 30–40 min with constant shaking at room temperature, according to the method in Ormeño et al. (2010).

Stored BVOC and waxes were analyzed by GC (Hewlett Packard GC 6890) coupled to a MSD (5973 Network) with an HP-5MS capillary column $50\,\mathrm{m} \times 0.20\,\mathrm{mm} \times 0.33\,\mathrm{\mu m}$. Four μ l of extracts were injected through an automatic injector (ALS 7683) at $280\,^\circ\mathrm{C}$ in split mode (5:1) with purge flow of $20\,\mathrm{ml\,min}^{-1}$ after 5 min. Helium carrier gas was used with constant flow rate of $1\,\mathrm{ml\,min}^{-1}$. The oven temperature program was $50\,^\circ\mathrm{C}$ for 2 min, increasing to $200\,^\circ\mathrm{C}$ at $4\,^\circ\mathrm{C}\,\mathrm{min}^{-1}$, then increasing to $320\,^\circ\mathrm{C}$ at $20\,^\circ\mathrm{C}\,\mathrm{min}^{-1}$ and then held constant for $15\,\mathrm{min}$. The MSD transfer line heater was maintained at $330\,^\circ\mathrm{C}$. Mass detector parameters were: ion source, $230\,^\circ\mathrm{C}$; quadrupole, $150\,^\circ\mathrm{C}$; EI, $70\,\mathrm{eV}$; EMV, $1530\,\mathrm{V}$; acquisition in scan mode from $40\,\mathrm{to}\,800\,\mathrm{amu}$.

Terpenes and waxes were identified by comparison of their retention index (RI) and mass spectra with those reported in published mass spectra libraries (Adams, 2007). Quantification was achieved based on relative response factors calculated for each single standard compound, when available.

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We report here the most extensive in-situ measurements of a suite of BVOC concentrations and fluxes observed to-date in a citrus orchard. For each measured compound reported in Table 1, we show the concentration at 4.85 m averaged every 30-min for the full measurement year (Fig. 1). We also show vertical gradients averaged over the diurnal cycle by interpolation of mean mixing ratios at the measurement heights of 1 m, 3.76 m, 4.85 m and 9.18 m (Figs. 2, 3), and the fluxes measured over the full year (Fig. 4). The year-long database allowed us to calculate the basal emission factors (BEF) of the temperature-dependent BVOC for the most important seasons (Table 2) using an exponential temperature dependence (Guenther et al., 1995; Tingey et al., 1991).

For all compounds, it was evident that the ambient concentration followed a diurnal cycle that was dependent on a combination of changes in emissions and changes in boundary layer dynamics, with additional influences from differences in regional photochemistry. The dynamics of the boundary layer in the Central Valley have recently been described by Bianco et al. (2011). During the day, when convective heat expands the boundary layer thus increasing the vertical mixing volume, ambient concentrations decrease. Lower mid-day concentrations are typical of plant ecosystems with both day and night emissions because the boundary layer is shallower at night causing the local concentration of emitted BVOC to increase. The boundary layer dynamic also influences ability to measure fluxes. Large gradients during the night were indicative of extremely slow vertical mixing, making vertical fluxes difficult or impossible to measure by eddy covariance. The daily dynamic of measured fluxes showed maximum peaks in the central hours of the day with minima at night (Fig. 5).

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3.1.1 Methanol

Methanol was the VOC observed with the highest mixing ratio, with peak values up to 35 ppbv (Fig. 1) during the spring-summer months suggesting high emissions from vegetation. This observation is consistent with previous results showing that increased emission occurs due to phenological modification of leaf tissues during leaf expansion, (Schade and Goldstein, 2002; Huve et al., 2007; Fall, 2003), and oxidative stress (Karl et al., 2001; Loreto et al., 2006), as a result of pectin demethylation when cell walls elongate during leaf expansion (Fall and Benson, 1996; Galbally and Kirstine, 2002), with plant growth recognized as the primary global source of methanol to the atmosphere (Galbally and Kirstine, 2002). In the diurnal cycle of gradient concentration shown in Fig. 2 for the winter and summer period, a slight gradient is visible in the morning between 09:00 a.m. and 11:00 a.m. during summer. The gradient is less visible during the central hours of the day, although these are the hours when maximum fluxes were recorded, rising up to a summer time average of 2 nmol m⁻² s⁻¹ (Fig. 5). Higher fluxes of methanol during mid-day hours have been previously described, with lightdependent emissions (Huve et al., 2007), and evidence of newly assimilated carbon reemitted as methanol exhibiting a temperature dependence (Folkers et al., 2008; Fares et al., 2011). Strong nocturnal gradients decreasing from above down to the canopy suggest that some deposition occurs at night. This may be explained by the presence of dew on leaves, which we measured using sensors for leaf humidity (data not shown). Previous research (Karl et al., 2004) showed that deposition on wet leaves can be responsible for a large percentage of total deposition, the latter enhanced by hydrolytic reactions (Jayne et al., 1992). Despite this deposition process at night, it was difficult to detect a negative flux at night with the eddy covariance system (Fig. 5) because of the low turbulence which caused systematic underestimation of nighttime fluxes. Therefore most of the night measurements have been discarded due to low atmospheric turbulence $(u^* < 0.15 \,\mathrm{m\,s}^{-1})$. Some positive gradients, however, were evident in the early

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evening hours (07:00–11:00 p.m.), in particular in the summer period. At these hours leaves are not wet and a foliar emission may be generated since methanol is produced during tissue expansion. This positive gradient from the canopy to the atmosphere in the early evening was less visible for the winter period, when there was greater water condensation on leaf surfaces (data not shown), especially during the night hours, and decreased rates of biosynthesis in pathways for methanol formation (e.g. decrease in plant growth and photosynthesis) occurred. Flowering has been shown to produce a burst of terpenic and non-terpenic compounds (Arey et al., 1991; Ciccioli et al., 1999; Fares et al., 2011), and we observed an increase of ambient concentrations up to 35 ppb and emission fluxes above 3 nmol m⁻² s⁻¹, in agreement with branch enclosure studies (Fares et al., 2011), where in the latter case the BEF was about two times higher (Table 2), and there was a decoupling of emissions from photosynthesis. Both of these results clearly show an increase of emission during flowering, but it is difficult to compare the branch enclosure and field studies quantitatively because the abundance and stage of flowering varied.

3.1.2 Acetaldehyde and acetone

An estimate of global emission for acetaldehyde similar to that for acetone was recently reported by Millet et al. (2009). The good correlation of acetaldehyde vs. acetone (slope = 1.1, R^2 = 0.8) confirms a similar origin of these compounds, as previously observed (Karl et al., 2003; Schade and Goldstein, 2002). The physicochemical properties of these two organic compounds differ in terms of the reactivity in the liquid phase thus affecting their solubilities and Henry's law constants (Noziere and Riemer, 2003), with acetone being less reactive than acetaldeyde (Duncan et al., 1999). Acetaldehyde and acetone were also measured in concentrations up to 15 ppb during the flowering period (Fig. 1), with ambient concentrations of each of these compounds equal to about one half of methanol. The orchard was a source of acetone and acetaldehyde during summer, with a visible positive gradient in the early evening, and became neutral or a slight sink during winter (Fig. 5).

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Acetone is the most abundant ketone in the atmosphere (Koppmann and Wildt, 2007), released during senescence (de Gouw et al., 1999) and oxidative stress in plants (e.g. from ozone) (Cojocariu et al., 2005), with global emissions estimated at 95 Tgy⁻¹ (Jacob et al., 2002). Acetone is another OVOC emitted by citrus leaves that also forms in the atmosphere through oxidation processes. We directly measured acetone fluxes with eddy covariance (Table 1, Figs. 4-5). Flowering significantly increased acetone emission, as shown from the enhanced atmospheric concentrations and the hourly fluxes (Fig. 8), reaching levels up to 2 nmol m⁻² s⁻¹, a value about two times higher than the typical summer emissions. A minor deposition of acetone is evident as a small gradient in Fig. 2. Our results agree with previous research which found that rural areas can have significant sources of acetone (Goldan et al., 1995; Riemer et al., 1998; Ciccioli et al., 1999; Schade and Goldstein, 2001).

Acetaldehyde is emitted by leaves in large quantities during and after abiotic stresses (Fall et al., 1999; Loreto et al., 2006). Acetaldehyde in particular has been shown to be emitted by citrus plants especially during flowering (Fares et al., 2011), although this compound is also produced by atmospheric oxidation processes (e.g. photooxidation of linalool), as described by Ciccioli et al. (1999), and Smith et al. (1996). This last formation source may justify the high amount we recorded in the field especially relative to monoterpenes, in comparison with the minor amount measured in branch enclosures, where reactivity in the gas phase was minimized (Fares et al., 2011).

BEF variations between winter, spring (flowering season) and summer for methanol and acetone (Table 2), showed higher values during flowering, probably due to enhanced pectin demethylation (Galbally and Kirstine, 2002) during organ development and flowering. A seasonal variation of BEF was also observed for an oak forest (Geron et al., 2000), a hardwood forest (Karl et al., 2003) and a pine forest (Schade and Goldstein, 2006; Holzinger et al., 2006). In particular, Karl et al. (2003) observed a BEF for acetone higher in fall, ascribing this major emission to decaying plant material, and Schade and Goldstein (2006) saw enhanced emissions of acetone and methanol in the spring during budbreak and elongation of pine needles.

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3.2.1 Isoprene

Isoprene was measured in relatively low concentrations, rarely above 2 ppbv except during the flowering season, when nocturnal peak concentrations increased to 5 ppbv (Fig. 1). Isoprene fluxes were negligible in all seasons (Fig. 5), in agreement with previous findings showing that orange is not a high isoprene emitter (Winer et al., 1992; Ciccioli et al., 1999; Fares et al., 2011). During the winter period, the orchard was acting more as an isoprene sink based on our observations of the concentration gradients (Fig. 3). An evident deposition phenomenon is occurring at ~04:00 p.m. We hypothesize that isoprene was transported to the orchard through advection plumes from a source far away from our measuring footprint, likely oak trees in the foothills. Similar to isoprene, its primary oxidation products methyl-vinyl-ketone and methacrolein (MVK + MACR) follow the same pattern during the winter and in summer. Deposition of MVK + MACR has been recently observed by Karl et al. (2010) in a tropical forest, as a result of uptake and degradation inside leaves by enzyme activity. During summer, both isoprene and MVK + MACR followed the same dynamics, with a notable positive gradient suggesting emission from the ground level to above the canopy in the early evening hours. Despite this positive gradient, the low turbulence did not allow measurement of a significant flux during those hours, but the observed gradient suggests that a minimal production of isoprene occurs in the early night hours. This may be explained by the consumption at night of residual substrate pools (e.g. dimethyl-allyldi-phosphate) produced during photosynthesis in the light hours, although a strong post-illumination decay in isoprene emission has been described to happen in a few minutes (Rasulov et al., 2009) which does not correspond to the time delay observed in our study (2-3 h). The positive fluxes of MVK + MACR during the same hours are consistent with recent findings (Jardine et al., 2012) that isoprene oxidation products can be emitted directed from leaves as result of intercellular oxidation of isoprene with ROS (Reactive Oxygen Species).

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Monoterpenes were the isoprenoid emissions observed in the highest abundance. Both in winter and in summer, a positive gradient from the ground to above the canopy was detected (Fig. 3), although fluxes were quite small (<0.3 nmol m⁻² s⁻¹). Soil and litter may significantly contribute to monoterpene emissions, in part due to the organic matter degradation processes in soils, and in part due to biomass wounding and decay following harvesting or pruning operations. Consistent with this suggestion, in a study of limonene fluxes in a navel orange orchard, soils have been demonstrated to be a major source (Ciccioli et al., 1999).

In our study, both summer and winter fluxes of monoterpenes were quite low, with mid-day values of $\sim 0.4 \, \text{nmol} \, \text{m}^{-2} \, \text{s}^{-1}$. Much higher monoterpene fluxes were observed during the flowering period, when the obvious smell of terpenes permeated the whole region, consistent with previously reported findings in greenhouse-based studies with plant enclosures (Fares et al., 2011). We also measured changes in the ambient concentration of monoterpene species between spring (flowering) and summer periods (Fig. 6). During the flowering period, beta-myrcene, d-limonene, trans-beta-ocimene and sabinene were the most abundant monoterpene species, while monoterpenes in summer were dominated by d-limonene. Changes in emission blend during flowering were also observed in branch enclosure studies (Fares et al., 2011), confirming that certain biosynthetic pathways are triggered to attract pollinators. However for the enclosure studies, during flowering the dominant monoterpenes were the cis and trans forms of beta-ocimene, while during summer we measured prevalently beta-myrcene. These differences suggest that the measured monoterpenes in the atmosphere may have a biogenic signature but not necessary attributed to Valencia orange. Advective air masses from nearby cultivations of Murcot mandarin may have contributed to the measured monoterpene concentrations, and indeed we observed limonene emissions from Murcot mandarin in the enclosure studies. Another reason for the discrepancy may lie in the use of Navel orange for the enclosure studies, which may have different ACPD

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emissions than Valencia orange. We also want to point out that chemical reaction with ozone in the gas-phase is an important phenomenon in the California central valley, when ozone concentrations often exceed 100 ppb, and limonene chemical lifetime is longer than other monoterpenes (e.g. myrcene), as pointed out in previous studies (Fares et al., 2010), supporting the hypothesis that certain monoterpenes may have reacted in the atmosphere before being measured. This point is particularly important for sesquiterpenes, and is discussed in the next section.

The monoterpene content in leaves and the cuticular wax thickness may explain the lack of large seasonal change in monoterpene emissions between winter and summer. Terpene accumulation within leaves was highest in summer, followed by spring, with the lowest levels in fall and winter (Fig. 7a). Leaf waxes, represented by long-chain aliphatic alkanes, especially C₃₁H₆₄ and C₃₃H₆₈, showed the same seasonal pattern (Fig. 7b). A positive linear relationship was observed between the total monoterpene hydrocarbon concentration (y) and cuticle wax concentrations (x, represented by longchain aliphatic compounds) $(R^2 = 0.75, y = -9.27 + 3.72x)$ as well as between the total monoterpene hydrocarbons (y) and 1-dodecanol (x) (the only aliphatic alcohol detected) ($R^2 = 0.86$, y = 5.82 + 25.80x). These results suggest that despite higher synthesis and thereby accumulation of terpenes during the warmest months, cuticle waxes, which are known to accumulate during summer to minimize water losses, result in a limitation of terpene emissions to the atmosphere, as noted in Ormeño et al. (2011).

During flowering, ambient concentrations of monoterpenes reached their maximum with nocturnal values up to 10 ppb (Fig. 1), and fluxes reached the maximum annual levels in agreement with branch-level experiments in the greenhouse (Fares et al., 2011) and previous research (Ciccioli et al., 1999; Hansen and Seufert, 2003; Arey et al., 1991), showing values during the day close to 1.8 nmol m⁻² s⁻¹. In agreement with the enclosure experiment (Fares et al., 2011), BEF for monoterpenes were much higher during the flowering period than in summer (1.31 and 0.13 nmol m⁻² s⁻¹, respectively, Table 2), suggesting that in spring, the biosynthesis of monoterpenes is

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enhanced due to both emissions released by flowers and biotic damage to leaves. It has been previously shown that certain monoterpene species (e.g. ocimene, Fig. 6) are emitted in large amount during flowering, to attract pollinators (Dudareva and Pickersky, 2000) and due to insect damage (Pare and Tumlinson, 1999). The magnitude of BEF from enclosure experiments was almost five times higher than field-based BEF. This may be explained in part by a possible overestimation of canopy-scale fluxes obtained by simply scaling-up enclosure-derived BEF by LAI and considering shaded leaves to contribute as much to emissions as light-exposed leaves. Another reason for this discrepancy in flux magnitude could be that potted and well-watered trees in the greenhouse did not experience the high temperature and ozone stress which occurs in the Central Valley, with limitations in plant water as shown by vapour pressure deficit occurring even in presence of sufficient irrigation. Our results highlight the importance of calculating BEF in different seasons for a proper parameterization of emission models, as also suggested through measurements by Goldstein et al. (1998), Keenan et al. (2009) and Niinemets et al. (2010).

3.3 C-6 oxygenated compounds

Concentrations of aldehydes and alcohols containing 6 carbon atoms (hexenals and hexenols, respectively) were measured by the PTRMS at m/z 83. Hourly gradient concentrations for the winter and summer season are summarized in Fig. 8. The absence of a gradient from the ground to above the canopy during winter suggests that no significant emission took place during the cold period, but rather a strong deposition occurred in the afternoon at ~03:00 p.m., so that we can hypothesize transport of emissions from remote sources to the site. During summer, hexenals and hexenols had a diurnal pattern similar to monoterpenes, with gradients from the soil to above the canopy indicating emissions. Although we did not measure fluxes, we hypothesize that the orange orchards are a source of these so called "C-6 compounds" or "green leaf volatiles" to the atmosphere. C-6 compounds are formed of 6 carbon atoms and can originate from denaturation of membrane lipids, predominantly from unsaturated fatty

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acids, under the action of lipoxygenase and hydroperoxide lyase enzymes (Hatanaka, 1993). Emissions have been record in response to wounding (Fall et al., 2003; Loreto et al., 2007) and exposure to ozone (Heiden et al., 2003). Flowering and wounding due to harvesting apparently caused high concentrations (>4 ppb) recorded at night during the flowering and harvesting periods (Fig. 1).

3.4 Aromatic compounds

Concentrations of aromatic compounds have been measured with PTRMS at m/z 79, 93, 107, and could be likely ascribed to Benzene, toluene + p-cymene, and benzaldehyde + xylenes, respectively. Since a detailed analysis with GC/MS is not available for these compounds, we cannot discriminate between isobaric compounds (e.g. relative abundance of toluene vs p-cymene, and benzaldehyde vs. xylenes). These benzenoid compounds are typically assumed to originate from anthropogenic sources such as fuel use and combustion. Previous research in urban areas has shown how peaks of these compounds are recorded during high vehicular traffic, making these compounds good proxies for anthropogenic fuel emissions. Recent findings support the idea that plants can also produce and emit small amounts of benzenoid compounds. While these fluxes are very small and expected to be negligible in terms of their impact on atmospheric chemistry in a fairly polluted environment such as the San Joaquin Valley, the benzenoid emissions from plants are very interesting and may play a role in plant signalling and trophic interactions.

We show vertical gradient concentrations of m/z 79, 93, 107 (Fig. 9), for winter, summer, and the flowering period. In all seasons, we identify in the early evening a clear positive gradient indicative of emissions, similar to that observed for isoprenoids, which causes us to hypothesize biogenic sources for these compounds. This emission pattern is guite evident for the summer season, and the figure includes a significant number of averaged days (N > 80). During the flowering period, concentrations were higher, in agreement with previous finding reporting that during flowering the biosynthetic pathways for benzenoid compounds is triggered by up-regulating genes of the shikimate

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pathway (van Schie et al., 2006). Benzenoic compounds can play a role, in addition to certain isoprenoids (e.g. linalool, ocimenes), in attracting pollinators (Dudareva et al., 2004). Interestingly, a source of m/z 107 seems to be located in the upper part of the canopy based on the observed vertical gradients, just where a higher density of flowers was observed, thus suggesting a floral origin. Future research is needed to assess whether biogenic emissions of these compounds are significant compared to anthropogenic emissions, and to better determine the biological significance of these emissions.

Evidence for unmeasured additional BVOC emissions

The PTRMS technology employed in this study allowed us to measure fluxes of major atmospheric VOC components for one full year. Together with fluxes of selected compounds, the PTRMS measured concentrations and vertical gradients of other important VOC which we attribute to a biogenic origin, described in the sections above. We want to mention that during an intensive summer campaign, a PTR-TOF-MS (Proton Transfer Reaction Time of Flight Mass Spectrometer) was deployed at the field site, allowing for one month of flux measurements for an extended suite of compounds. While the PTR-TOFMS showed excellent agreement with the measurement of the compounds observed with PTRMS, it allowed us to measure a multitude of additional masses which significantly increased the observed total biogenic emissions from the orchard. Thus, the fluxes reported here can indeed explain the seasonality of the major BVOC emitted from vegetation, although we are confident the reported compounds do not represent the totality of emissions.

Among the unmeasured compounds are sesquiterpenes. These compounds are a very important class of isoprenoids which have been identified when emissions from oranges were measured with branch enclosures (Fares et al., 2011; Ciccioli et al., 1999). Sesquiterpenes are very reactive with tropospheric ozone (Atkinson and Arey, 2003) and therefore have very short atmospheric lifetimes. β -caryophyllene is the main sesquiterpene emitted from citrus (Fares et al., 2011; Ciccioli et al., 1999), and we **ACPD**

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estimated an atmospheric lifetime of ~30-80s when ozone concentrations are between 40 and 100 ppb, and peak ozone was often measured at the orchard at levels from 70 to almost 120 ppb in summer during the study. We tried to minimize the residence time of the air in the sampling line (\sim 2.2 s), but the high reactivity with ozone, the poor transmission efficiency of β -caryophyllene in the PTRMS, and likely losses in our sampling lines resulted in very low concentration measurements for these compounds (0.05 ppb during summer mid-day), and these conditions preclude us from providing a quantitative analysis of these emissions. A discrepancy in magnitude between β caryophyllene measured in branch enclosures and in the field at the whole canopy scale was also reported by Ciccioli et al. (1999), with enclosure fluxes being very high, even higher than monoterpenes, similar to what we found in our greenhouse enclosure experiments (Fares et al., 2011). In the field, Ciccioli et al. using the relaxed eddy accumulation (REA) technique observed low fluxes of β -caryophyllene in comparison to the enclosure measurement, and justified this by the high estimated resident time (360–480 s) of the molecule in the air space between the soil and the sensor above the canopy. Turbulence at our site was low, similar to that observed by Ciccioli et al.; therefore, it is reasonable to hypothesize a similarly long residence time for β -caryophyllene relative to its atmospheric lifetime. Our results therefore suggest that models to predict sesquiterpene emissions from BEF estimated using leaf/branch enclosure data from the controlled greenhouse experiments may be a better and more quantitative approach than what we could achieve in the field, where oxidant (ozone) levels were high and sesquiterpene lifetimes were very short.

Conclusions

We measured concentrations and fluxes of an important fraction of BVOC emitted from a Valencia orange (Citrus sinensis) orchard continuously for one year. This Citrus species is extensively cultivated in California and in many other regions of Mediterranean climate. Our observations were focused on the most abundant BVOC which we

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previously identified using enclosures of plants grown in greenhouses under optimal conditions.

Oxygenated VOC (methanol, acetaldehyde, and acetone) and monoterpenes were the major BVOC emissions we observed from oranges in the field. We found gradients in concentrations from the soil to above the canopy, especially at night, when the atmospheric boundary layer is low and vertical turbulence is minimized, suggesting that fluxes occur even at night, but cannot be quantitatively described in our study due to methodological limitations of the eddy covariance technique when turbulence is low. We found high concentrations of monoterpenes stored inside leaves during summer coinciding with lower emissions and high cuticular wax content; thus, we hypothesize limitations to gaseous emissions because of the enhanced physical barrier.

Isoprene and its oxidation products, methyl-vinyl-ketone and methacrolein, were shown to be emitted in very small amounts in the late afternoon, and occasionally deposited at night (again in small amounts) based on observation of their vertical profiles. Our observations of a positive gradient of these oxidation compounds supports recent findings (Jardine et al., 2012) that a fraction of these compounds may be formed inside leaves from reaction of isoprene with reactive oxygen species (ROS).

In this study, we reported atmospheric concentrations of other important volatile compounds which may have a biogenic source. Both aromatic and C6 oxygenated compounds had positive concentration gradients from the ground to above the canopy, demonstrating that orchards emit small amounts of these compounds. Although further research is necessary to determine the importance of biogenic emissions of these compounds for atmospheric chemistry in comparison with anthropogenic emissions in this environment, they may be important for pollinator attraction and other ecological interactions.

The full year of measurements allowed us to compare BEFs for different seasons. We found that during flowering BEFs were consistently higher for most BVOC studied. This seasonality of BVOC emissions from crops should be considered in global and statewide emission models, and the largest annual fraction of emissions from Central **ACPD**

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Valley crops that flower in springtime is likely to occur during that period. Our results should be useful in atmospheric chemistry models to estimate whether BVOC emitted from these crop species play a significant role in regional photochemistry, especially when Citrus plantations are close to urban areas as in the Central Valley of California, where BVOC emissions can combine with anthropogenic emissions to contribute to ozone and secondary aerosol production.

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Table 1. BVOC species measured during the field campaign in 2010. Bold indicates compounds (m/z) for which fluxes were measured. n.a. = not available.

m/z	Compound	Formula	Dwell time (s)	Norm. sensitivity (ncps ppbv ⁻¹)	Detection limit (ppbv)	Mid-day mixing ratio ³ (ppbv)	Mid-day flux ⁴ (nmol m ⁻² s ⁻¹)	mean 95 % Cl ⁵	N (tot 30-min flies)	N (after filtering)	Quality test (1, 2, 3)
33	methanol	CH ₃ OHH ⁺	0.2, 0.5	5.9 ± 0.02	1.15 ± 0.24	2.59, 6.2, 9.02	0.26, 2.74, 1.48	0.04, 0.48, 0.11	5003, 5141	2427, 5104	1076, 189, 1699
42	acetonitriie	C ₂ H ₃ NH ⁺	1	11.2 ± 0.04	0.06 ± 0.01	0.07, 0.09, 0.18	n.a.		n.a.	5141 2804	n.a.
45	acetadehyde	C ₂ H ₄ OH ⁺	0,5	10.6 ± 0.04	0.36 ± 0.09	0.99, 1.4, 2.6	n.a.	n.a.	5135	4293	n.a.
59	acetone	C ₃ H ₆ OH ⁺	0.2, 1	13.8 ± 0.04	0.14 ± 0.04	0.98, 2.1, 3.5	0.03, 0.57, 0.31	0.02, 0.12, 0.05	5003, 5135	2427, 3482	1076, 189, 1699
69	isoprene ¹	C ₅ H ₈ H ⁺	0.2, 1	8.7 ± 0.03	0.03 ± 0.02	0.22, 0.20, 0.3	0.05, 0.06, 0.04	0.01, 0.01, 0.01	5003, 5135	2427, 4776	1076, 189, 1699
71	MVK+MCR ²	C ₄ H ₆ OH ⁺	1	10.07 ± 0.03	0.04 ± 0.02	0.07, 0.18, 0.27	n.a.	n.a.	5135	4669	n.a.
79	benzene	C ₆ H ₆ H ⁺	1	9.6 ± 0.03	0.03 ± 0.01	0.12, 0.07, 0.06	n.a.	n.a.	5135	4529	n.a.
83	hexanal, hexenols	$C_6H_{10}H^+$	1	10.5 ± 0.03	0.02 ± 0.02	0.14, 0.25, 0.15	n.a.	n.a.	5135	4742	n.a.
87	2-methyl-3-butene-2-ol	C ₅ H ₉ OHH⁺	1	9.5 ± 0.03	0.05 ± 0.02	0.03, 0.04, 0.07	n.a.	n .a.	5135	4707	n.a.
93	toluene, p-cymene	C ₇ H ₈ H ⁺	1	12.6 ± 0.04	0.02 ± 0.01	0.06, 0.05, 0.05	n.a.	n.a.	5141	4904	n.a.
99	hexenals	C ₆ H ₁₀ OH ⁺	1	10.6 ± 0.03	0.04 ± 0.01	0.03, 0.04, 0.06	n.a.	n.a.	5141	4674	n.a.
107	benzaldehyde, xylenes	$C_8H_{10}H^+$	1	10.5 ± 0.03	0.04 ± 0.01	0.03, 0.05, 0.02	n.a.	n.a.	5141	4754	n.a.
111	oxidation product	n.a.	1	10.4 ± 0.03	0.02 ± 0.02	0.00, 0.01, 0.00	n.a.	n.a.	5141	3537	n.a.
113	oxddation product	n.a.	0.2, 1	10.3 ± 0.03	0.02 ± 0.02	0.01, 0.02, 0.04	0.00, 0.00, 0.00	0.00, 0.00, 0.00	5003, 5141	2427, 4088	1076, 189, 1699

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Table 1. Continued.

m/z	Compound	Formula	Dwell time (s)	Norm. sensitivity (ncps ppbv ⁻¹)	Detection limit (ppbv)	Mid-day mixing ratio ³ (ppbv)	Mid-day flux ⁴ (nmol m ⁻² s ⁻¹)	mean 95 % Cl ⁵	N (tot 30-min flies)	N (after filtering)	Quality test (1, 2, 3) ⁶
81, 137	monoterpenes	C ₆ H ₉ ⁺	0.2, 1	10.5 ± 0.03	0.03 ± 0.02	0.16, 0.52, 0.19	0.15, 0.61, 0.17	0.01, 0.04, 0.03	5003, 5135	2427, 4.815	1076, 189, 1699
139	oxidation product	n.a.	1	8.8 ± 0.03	0.02 ± 0.02	0.01, 0.01, 0.00	n.a.	n.a.	5141	3409	n.a.
149	methyl-chavicol	C ₁₀ H ₁₂ OH ⁺	1	8 ± 0.02	0.02 ± 0.02	0.01, 0.02, 0.00	n.a.	n .a.	5141	3563	n.a.
151	oxidation product	n.a.	1	7.9 ± 0.02	0.02 ± 0.02	0.01, 0.01, 0.00	n.a.	n.a.	5141	3094	n.a.
155	oxygenated monoterpenes	C ₁₀ H ₁₈ OH ⁺	1	7.6 ± 0.02	0.02 ± 0.02	0.01, 0.01, 0.00	n.a.	n.a.	5141	2830	n.a.
205	sesquiterpenes	C ₁₅ H ₂₄ H ⁺	5	2.4 ± 0.00	0.03 ± 0.03	0.17, 0.06, 0.00	n.a.	n.a.	5141	2572	n.a.

Furans and methyl-butenol fragment are also a minor contributor to m/z 69 from intercomparison with GC-MS.

² Sum of methyl-vinyl-ketone and methacrolein;

³ The limit of detection (LOD) is calculated setting a minimum acceptable signal to noise ratio equal to 2.

⁴ Median concentration values at 4.85 m above ground in winter, flowering, summer periods, respectively, in the central hours of the day: 12:00 to 14:0;

⁵ Median values in winter, flowering, summer periods, respectively in the central hours of the day: 12:00 to 14:00 for the 30-min files which were assigned to quality category 1 and 2;

⁶ Measure of the Confidence Interval as the mean noise for the 30-min fluxes which were assigned to quality category 1 and;

⁷ Number of flux observations with 1 = good quality (distinct maximum in the covariance, $R^2 > 0.2$), 2 = low quality (slightly visible maximum in the covariance), 3 = poor quality (or no visible maxima).

Table 2. BVOC basal emission factors (BEF, nmol m⁻² s⁻¹) of valencia Orange for winter, flowering, summer periods, respectively. The β value calculated from the Tingey (T) algorithm is reported below for each BVOC species. Data \pm standard deviations refer to basal conditions of Temperature = 30 \pm 2 °C extrapolated from the observations. BEF for the 2010 field experiment were calculated using β values from the greenhouse experiment using plant cuvettes because in the greenhouses the environmental conditions were close to basal condition thus providing a more robust dataset for β calculation. β from summer 2008 was used to calculate BEF for winter period. Leaf-scale BEF measured using enclosures were up-scaled to canopy level by multiplying by Leaf Area Index (LAI = 3 m² (leaf) m⁻² (ground)).

	2008 Cuvette	2010 Field (using beta from 2008)
Monoterpenes β	n.a., 6.29 ± 1.73, 2.01 ± 1.37 n.a., 0.15 ± 0.016, 0.14 ± 0.014	0.87 ± 0.49 , 1.31 ± 1.51 , 0.13 ± 0.13
Methanol β	n.a., 7.11 ± 1.11 , 3.87 ± 1.61 n.a., 0.59 ± 0.04 , 0.59 ± 0.011	0.66 ± 1.2 , 3.98 ± 3.21 , 0.22 ± 0.23
Acetone β	n.a., 1.34 ± 0.17 , 0.64 ± 0.24 n.a., 0.12 ± 0.001 , 0.078 ± 0.009	0.11 ± 0.37 , 21.11 ± 0.66 , 0.35 ± 0.41

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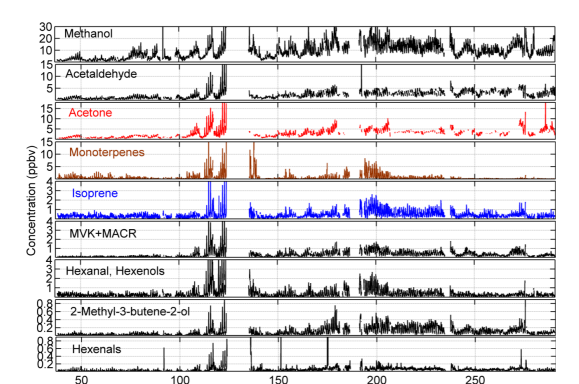


Fig. 1. Concentration of the major BVOC species measured hourly by PTRMS at 4.85 m above ground at the Citrus site between February and November 2010.

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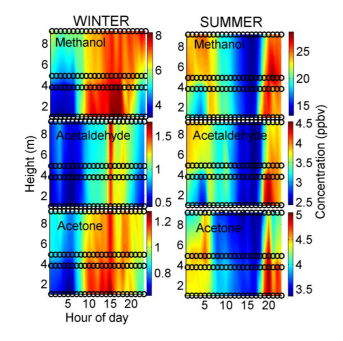


Fig. 2. Hourly average winter and summer vertical profiles of the major OVOC species measured by PTRMS (methanol, acetaldehyde, acetone) within (1.m, 3.76 m) and above (4.85 m and 9.18 m) the orange orchard canopy (circles indicate measurement heights).

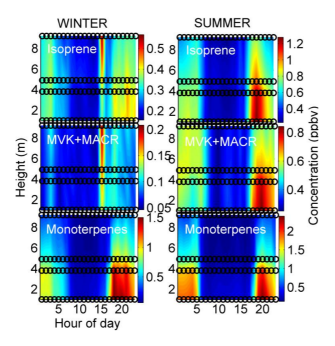


Fig. 3. Hourly average winter and summer vertical profiles of the major isoprenoid species 816 measured by PTRMS (isoprene, sum of isoprene oxidation products methyl-vinyl-ketone and 817 metachrolein, and sum of monoterpenes) within (1. m, 3.76 m) and above (4.85 m and 9.18 m) 818 the orange orchard canopy (circles indicate measurement heights).

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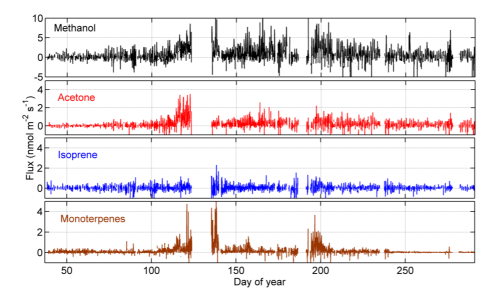


Fig. 4. Fluxes of the major BVOC species measured hourly by PTRMS Eddy Covariance at 824 the Citrus site between February and November 2010.

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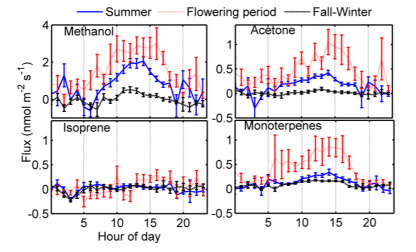


Fig. 5. Hourly average fluxes of BVOC species measured by PTRMS at the Citrus site during 829 winter (black dashed line), flowering (red dotted line) and summer (blue line) periods.

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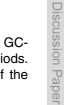
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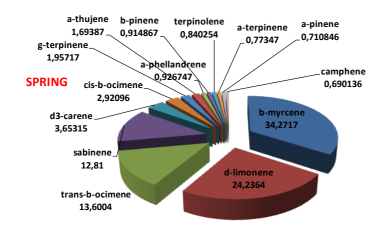
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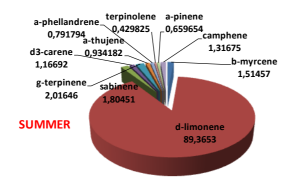
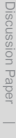


Fig. 6. Pie chart showing the percent distribution of monoterpene species measured with GC-MS during the spring flowering (25 April to 5 May) and summer (11 to 24 August) periods. Percents were calculated as the total fraction of the mass observed over the period of the measurements.



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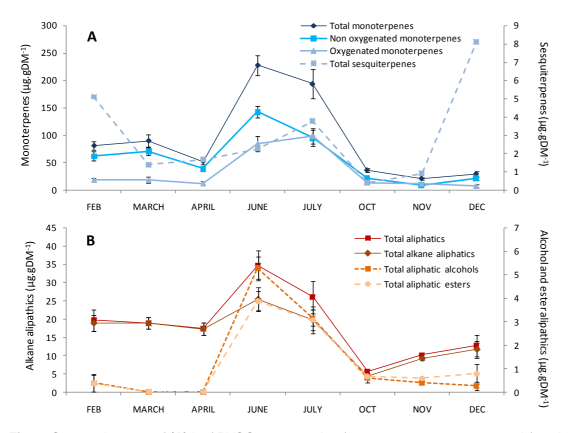


Fig. 7. Seasonal course of (A) leaf BVOC concentration (80 % monoterpene compounds) and (B) cuticle waxes (88% alkane-alipathic compounds). Values shown are the average of ten trees each month and bars denote the standard error.

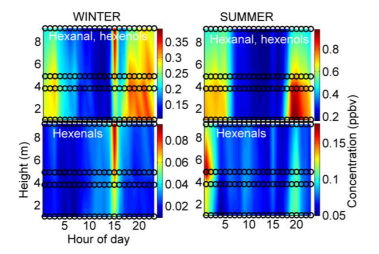


Fig. 8. Hourly average winter and summer vertical profiles for two C-6 compound classes measured with PTRMS at m/z 83 (hexenal, hexenols) and m/z 99 (hexanals) within (1.m, 3.76 m) and above (4.85 m and 9.18 m) the orange orchard canopy (circles indicate measurement heights).

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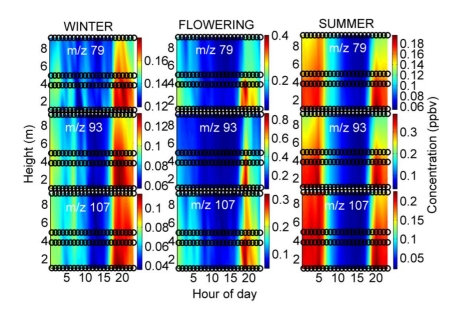


Fig. 9. Hourly average winter, flowering, and summer vertical profiles for the major aromatic compounds measured with PTRMS at m/z 79, 93, 107, corresponding to benzene, sum of toluene and p-cymene, and the sum of benzaldehyde and xylenes, respectively. Measurements were made within (1.m, 3.76 m) and above (4.85 m and 9.18 m) the orange orchard canopy (circles indicate heights).