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Compost spreading in Mediterranean shrublands has no strong short-term effect on Q. coccifera monoterpenes emissions at leaf level.

Keywords:
Monoterpenes
Biogenic emissions
Compost
Quercus coccifera
Nutrients

Abstract
Monoterpenes emissions of Quercus coccifera L. were repeatedly measured during the two years following the spreading of a sewage sludge compost at rates of 50 Mg ha⁻¹ and 100 Mg ha⁻¹, in a twelve-year-old post-fire Mediterranean shrubland. We also monitored the patterns of change in soil and leaf nutrient content, plant water potential, chlorophyll fluorescence, and plant growth. Compost spreading resulted in weak changes in leaf nutrient content and plant water status, and therefore no significant effect on monoterpenes emissions at leaf scale, except during one summer sampling, probably related to advanced leaf maturity with the highest compost rate. However, compost increased plant growth, particularly the leaf biomass. The results suggest that compost spreading in Mediterranean shrublands has no strong short-term effect on Q. coccifera monoterpenes emissions at leaf level, but may indirectly increase volatile organic compound fluxes at the stand scale, which may contribute to regional ozone pollution.

1. Introduction

Almost 50% of the plant species growing in Mediterranean type ecosystems produce and emit biogenic volatile organic compounds (BVOC) (Ross and Sombrero, 1991). Among these compounds, monoterpenes account for 11% of the global annual BVOC fluxes (Guenther et al., 1995). Terpene emissions are involved in atmospheric chemistry, notably by affecting the formation and degradation of tropospheric ozone in regions subjected to high solar radiation and in proximity to urban areas which are main sources of ozone precursors (Atkinson and Arey, 2003). These emissions are directly involved in air pollution (Varinou et al., 1999), particularly in the Mediterranean region, where critical ozone levels are recorded each year (European Environment Agency, 2009). BVOC emissions are also known to contribute to secondary organic aerosol formation (Guenther, 2002) and may increase the life span of greenhouse gases, such as CH₄, which are strongly involved in global change processes (Guenther et al., 1995).

A wide range of environmental factors have been shown to affect BVOC emissions. While the effect of climate-related factors is well documented, global eutrophication and soil nutrient impact on BVOC fluxes has received relatively little attention (Peñuelas and Staudt, 2010). For some years, studies have been carried out to assess the benefits and risks of using composted wastes to restore degraded natural soils, as an alternative to incinerating or dumping them (Gallardo-Lara and Nogales, 1987; Navas et al., 1999; Martinez et al., 2003; Larchevêque et al., 2006b, 2009). So far, the effects of compost on BVOC production have been investigated by a few studies which focused only on terpene content (Tanu et al., 2004; Hussein et al., 2006; Abdelaziz et al., 2007), but there have been none on emissions. The spreading of nutrient rich compost may directly affect the plant capacity to produce and emit BVOCs. Fertilization experiments have indeed been shown to increase or reduce foliar BVOC emissions depending on the type and dose of nutrients and experimental conditions (Harley et al., 1994; Lerdau et al., 1995; Litvak et al., 1996; Constable et al., 1999; Possell et al., 2004; Rosenstiel et al., 2004; Blanch et al., 2007; Fares et al., 2008).

In this context, our aim was to assess the effect of sewage sludge compost spreading, during two consecutive years, on the seasonal course of Quercus coccifera L. monoterpenes emissions, in a natural Mediterranean shrubland. Q. coccifera is an interesting model species for the Mediterranean area since this evergreen shrub covers a large surface of calcareous shrublands and is a pioneer species in post-fire lands (Konstantinidis et al., 2005). Furthermore,
little is known about its emission variability according to abiotic factors other than seasonality and drought. This species mainly features monoterpene emissions (Ormeño et al., 2007b) which are expected to behave in an "isoprene-like" way, since it does not have specialized structures to store terpenes (Hanssen and Seufert, 1996; Niinemets et al., 2002).

2. Material and methods

2.1. Experimental set-up

The experiment took place on the plateau of Arbois (Southern Provence, France; 51°89′00″E – 43°29′01″N in WSG-84 Norm), an area of 6000 m², 240 m above sea level, under Mediterranean climatic conditions, with hot, dry summers and cool, wet winters. The soil is calcareous, classified as Rendoll (Soil Survey Staff, 1999), with a high percentage of stones (77%). This confers a very heterogeneous structure and soil depth. It shows low organic matter and available nutrient content (Larchevêque et al., 2006a). The vegetation is that of a typical Mediterranean shrubland, composed of evergreen shrubs (30 to 100 cm high) such as Quercus coccifera L., Cistus albidus L., Rosmarinus officinalis L. and Lilea porphyrotricha Pourr. This shrubland underwent a fire in June 1995, from which these pioneer species developed to form the current stand.

The compost tested in this study was made from green wastes, pine barks and municipal sewage sludge in identical proportions. This mature compost met the French legal standards (NF U 44-095) for pathogenic microorganisms, organic trace elements and trace metals. The nutrient composition of the compost was similar to that used in Larchevêque et al. (2009) (total N: 2% of dry matter (DM); total P: 3.2% DM; exchangeable K: 0.323DM; pH(H2O): 7.7). It was homogeneously hand-spread on the field to a depth of 10 cm, then covered with 2 cm of soil. Three blocks received 2.5 t ha⁻¹ (D50) and three others 100 t ha⁻¹ (D100) (fresh mass). The later spreading was performed in November 2008.

The compost was performed in November 2008. To achieve representative emissions from the soil and vegetation, three treatments between 10 a.m. and 14 p.m. (universal time). Emissions were measured every 3 min at 60 °C to eliminate trapped water. Preliminary tests showed no significant compound loss during this pre-flush. Trapped monoterpene emissions were then desorbed at 250 °C for 10 min. After injection, temperature was set to 40 °C, then increased by 3 °C per minute to 250 °C maintained for 5 min. Carrier gas flow (N2) was 1 ml min⁻¹. Calibration was done with internal standard (dodecane) in each analysis, together with the frequent injection of commercial terpene standards (Sigma–Aldrich®; Firmenich®). Peak identification was done using retention index (Adams, 1989) and comparison with standard retention times, and was confirmed by analyzing some samples in a GC (Hewlett Packard GC8900®). Agilent, Palo Alto, CA, USA) coupled to a mass selective detector (MSD, HP 5973N).

Because Q. coccifera is a BVOC emitting species which does not store monoterpene, its field emission rates can be normalized to standard temperature (30 °C) and PPFD (1000 μmol photons m⁻² s⁻¹) values using Guenther et al. (1993) for isoprene emissions. This algorithm allows a correction of emission rate embodying short-term effects of temperature and light:

\[ E_{r} = E_{1} \times \frac{C_{1}}{C_{T}} \]

where \( C_{1} \) and \( C_{T} \) are respectively light and temperature dependence coefficients derived from experimental measurements on several isoprene emitting species.

Parameters used to calculate these coefficients are those proposed in Guenther et al. (1993). Hereafter, normalized and un-normalized emission rates are referred to as \( E_{r} \) and \( E \) respectively.

2.4. Leaf and soil sampling and nutrient analyses

After BVOC sampling, leaves of each twig were cut off, lyophilized (Alpha 1+4 LD plus, Christ GmbH, Germany), and reduced to powder with a ball mill. Soil samples were taken a few cm under the litter-compost layer, taking care not to include compost fragments. Each sample was a mixture of three soil collections randomly performed on each plot. The soil was oven-dried at 60 °C for 48 h, sieved and ground in a ball mill.

Leaf and soil N content were analyzed with a Flash EA 1112 Elemental Analyzer (Thermo Finnigan, Milan, Italy). Leaf P and K contents as well as soil available P (Olsen method) and extractable K content were analyzed in the LCA (Laboratoire Centre Atlantique, La Rochelle, France) with an ICP (inductively coupled plasma) method.

2.5. Fluorescence measurement and xylem water potential

Diurnal chlorophyll fluorescence measurements were performed on the plants chosen for BVOC sampling, using leaves nearby those enclosed. For each individual, the fluorescence of three leaves was measured, using a portable Fluorescence Monitoring System (FMS 2, Hansatech®), Kings Lynn, Norfolk, UK). Each measurement provided a variable to maximum fluorescence ratio (Fv/Fm) value, after 30 min dark-adaptation with a leaf clip holder. Fv/Fm is a measure of the quantum efficiency when all Photosystem II (PSII) centers are open, i.e., the maximum yield of light energy conversion, in a given environmental situation (Krause and Weis, 1991).

Predawn xylem water potential measurements were done using a pressure chamber (PMS Instrument Co. Corvallis, Oregon, USA) connected to a nitrogen tank. Measurements were performed on other individuals than those used for BVOC sampling to prevent any injury effect on the emissions rates. Three to five twigs were performed before sampling to minimize possible stress experienced by the leaves during bag installation. This pre-purging allowed the bag atmosphere to be renewed at least once completely. Afterwards, BVOCs were sampled during 15 min by passing a fraction of the bag air at 20 mL min⁻¹ through an adsorbent glass cartridge, by means of a void pump (Edwards RV8) and a bubble flowmeter (GPE, ‘MeTeRate’ 314-140/084 tube; Precision Engineering Ltd., Hemel Hempstead, UK). The adsorbant cartridge was filled with preconditioned Tenax TA (Varian®) and protected from sun-light with aluminium foil. During sampling, the bag remained over-pressurized, thus limiting contact between the leaves and the bag's inner surface. Inside the bag, temperature was measured near the leaves, while a small fan homogenized the atmosphere. Temperature within the bag – known to represent leaf temperature quite faithfully (Ortega and Helmg, 2008) – was generally 2 ± 1 °C higher than outside temperature. Photosynthetic photon flux density (PPFD) was measured outside bag enclosures using a portable photosystem plant and canopy transmission meter (EMS-7 Model; Sucechem® Marketing SDN, BHD, Kuala Lumpur, Malaysia). Preliminary tests showed that PPFD inside the bag over the upper leaves was not different from values recorded outside. After BVOC sampling, cartridges were protected and frozen using liquid nitrogen, then stored at −20 °C in the laboratory until analysis. Emissions were expressed on a leaf dry mass basis, determined after lyophilization of the sampled twigs.

Terpene analyses were conducted by gas chromatography (Pengcheng et al., 2008) coupled with a flame ionization detector (HP 5890 series II, Agilent, Palo Alto, CA, USA) equipped with a non-polar chromatographic column (Ultra 2, 50 m × 0.2 mm × 0.25 μm). Tenax TA cartridges were previously back-flushed during 3 min at 60 °C to eliminate trapped water. Preliminary tests showed no significant compound loss during this pre-flush. Trapped monoterpene emissions were then desorbed at 250 °C for 10 min. After injection, temperature was set to 40 °C, then increased by 3 °C per minute to 250 °C maintained for 5 min. Calibration was done with internal standard (dodecane) in each analysis, together with the frequent injection of commercial terpene standards (Sigma–Aldrich®; Firmenich®). Peak identification was done using retention index (Adams, 1989) and comparison with standard retention times, and was confirmed by analyzing some samples in a GC (Hewlett Packard GC8900®). Agilent, Palo Alto, CA, USA) coupled to a mass selective detector (MSD, HP 5973N).

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withdrawn (each twig representing a different plant) in each plot, corresponding to 9 to 15 measurements per treatment.

2.6. Growth measurement

As Kermes Oak makes one (exceptionally two) flush per year, depending on the climatic conditions (Pilar and Gabriel, 1998), growth measurements were performed on the three growth units of the longest axis, so that measurements integrate all the 2009 and at least part of the 2008 growth. We measured wood dry weight and dimensions on each of these growth units. Approximate wood volume was assessed by multiplying the basal diameter of the stem by its length. We counted the number of ramifications corresponding to the last flush (on the n-1 growth unit). On the last main growth unit (n) of the twig, we determined the number of leaves, measured their dry weight and calculated their Leaf Mass per Area ratio (LMA).

2.7. Statistical analyses

Repeated-measurements analyses (within subject ANOVA; von Ende, 2001) were performed with Statistica Edition 98 (StatSoft Inc., Tulsa, OK, USA) when comparing variables between non-independent individuals such as individuals from the same plots at different dates. It was used to test the effect of compost on E and k, and on leaf nutrient content. One-way ANOVA was performed using Statistical Graphics Plus 4.1 (Statpoint Onc, North Virginia, USA) to test the effect of compost on growth variables. Postulates for normality of distribution within each group as well as homoscedacity (variance homogeneity) were checked before ANOVAs, and data were transformed where needed to correct deviations from normality. When the data did not meet these requirements, as for soil nutrient, fluorescence and water potential, Kruskal–Wallis test was rather used, on Statistical Graphics Plus 4.1. Post hoc multiple range comparisons were done using Tukey test after ANOVA and Student–Newman–Keuls non-parametric tests after Kruskal–Wallis test.

3. Results

3.1. Soil and foliar nutrients

Mineral soil total nitrogen was not significantly altered by compost treatments (Kruskal–Wallis tests, $W = 4.4, P > 0.05$; Fig. 1). Available phosphorus increased sporadically in March, June 2008 and July 2009. Most of the significant changes in mineral soil nutrients were observed on extractable K content which was quickly and durably enhanced by compost spreading.

Consistently with soil nutrient content, leaf nutrient content was poorly affected by the compost spreading (Fig. 2). No compost effect was observed on leaf N content throughout the survey period (Within subject ANOVA; $F = 1.6, P > 0.05$). Similarly to soil P, leaf P was affected only sporadically by the compost treatments. Leaf K was increased by both compost doses but only for the sampling done two years after spreading (July 2009). For both soil and leaf nutrient content, we observed relatively few differences between the two compost doses.

Fig. 1. Soil total nitrogen, available phosphorus and extractable potassium content through dates and treatments as a percentage of dry matter (DM). (○: control, ■: D50, ▲: D100). Error bars represent standard error (n = 3). * Denotes a significant difference from the control mean (P < 0.05) (Kruskal–Wallis test).

Fig. 2. Leaf total nitrogen, phosphorus and potassium content through dates and treatments as percentage of dry mass (○: control, ■: D50, ▲: D100). Error bars represent standard error (n = 3). Asterisks denote a significant difference from the control mean (*: P < 0.05) (Within subject ANOVA).
3.2. Growth

Despite the low enrichment of the mineral horizon in N and P, the compost spreading improved vegetation growth (Table 1). Compost spreading did not affect either LMA or the number of leaves on the last main growth unit, but increased total leaf dry weight by more than 40% in D50 and D100, resulting in larger leaves. D100 treatment raised by 87% the mean number of ramifications on the n – 1 growth unit. The diameter of the stems and their length was significantly increased in D100 in all growth units, leading to a total axial wood volume raised by 130% in D100.

3.3. Plant water status and leaf fluorescense

Climate data showed high summer irradiance levels and typical Mediterranean patterns of precipitation, with summer drought leading to xylem water potential reduction (Kruskal–Wallis tests, W = 60, P < 0.001; Fig. 3, A and B)). At week scale, mean summer temperature variations were sawtooth-shaped, given that increases or decreases of several degrees in a few days occurred frequently.

Predawn water potential monitoring showed that no BVOC sampling was done during the strongest water deficit period, in 2008. Early and late summer samplings were performed on plants with water potentials above –1.5 MPa. Compost supply had no significant effect on plant water status during drought periods (Fig. 3, B)); W = 1.38, P > 0.05).

In 2008 and 2009, daytime fluorescence measurements showed Fv/Fm values lower than 0.80 (Table 2). However, Fv/Fm tended to be higher in the compost treated plots than in the control plots and this difference was significant in early June 2008 (Kruskal–Wallis tests; W = 6.2, P < 0.05).

4. Monoterpene emissions

Mean total E varied widely with seasons, with the lowest values around 1 µg gDM⁻¹ h⁻¹ from late November to early June, and values more than four times higher in summer (4.1 to 9.4 µg gDM⁻¹ h⁻¹) (Fig. 3, C)). No compost effect on E was evidenced. Mean total E varied throughout seasons overall (Q. coccifera, the most commonly studied Mediterranean oak species, which is, like observed a strong increase in the total Eₑ (on average 3.6 times up), similar in each treatment. Late June/early July Eₑ were not significantly different from one year to another. Normalization enhanced 7 to 10 times emissions in November and overall increased data variances within treatments in November and February samplings. Compost spreading did not affect Eₑ throughout the survey period, except in early June 2008 where Eₑ was significantly enhanced by 2.5 in the D100 treatment (Fig. 3, D)).

The emitted compound, throughout the experimental period, was α-pinene (34%), followed by limonene (25%) and β-myrcene (16%), whereas β-pinene, 1,8-cineole, sabine, camphene, α-terpineol and γ-terpinene contributed less than 6% to the total emissions. A sesquiterpene (α-humulene) was identified but detected in negligible proportions (<1%), so it was not included in the data analysis.

4. Discussion

4.1. Seasonal emission variations and normalization effect

Q. coccifera monoterpene E varied throughout seasons overall following seasonal temperature variations, with higher emissions during the warm period than during colder periods, as commonly described (Boissard et al., 2001; Sabillon and Cremaudes, 2001; Staadt et al., 2002; Pio et al., 2005). Eₑ values ranged within the same order of magnitude as that previously found for this species (Llusia and Peñuelas, 2000; Ormeño et al., 2007a). After normalization, the two close June 2008 samplings still revealed a sharp rise of Eₑ with values multiplied by 3 within two weeks. This increase was probably due to the exceptionally rapid weather change, with an 8 °C difference between the mean temperature of the first and the last ten day period in June. This suggests that mid term past weather (in the order of days), mainly mid term temperature, which is not integrated by the G93 normalization, could be one of the main environmental variables involved in this rapid summer increase (Fuentes and Wang, 1999; Sharkey et al., 1999).

The sporadic seasonal increases recorded here corroborate the results of Llusia and Peñuelas (2000) for the same species under field conditions, with higher Eₑ in summer and autumn than during the rest of the year. In their survey, four monoterpene emitting plant species were sampled, including Quercus ilex, the two most commonly studied Mediterranean oak species, which is, like

| Table 1 |

Growth measurements performed on Q. coccifera twigs, in February 2010, on the last growth unit (n) and the two previous one (n – 1 and n – 2). Numbers in brackets represent standard deviation (n = 30). Different letters denote significant differences between treatments (One way ANOVA). ns: non-significant.

<table>
<thead>
<tr>
<th>Growth flush</th>
<th>Control</th>
<th>D50</th>
<th>D100</th>
<th>Post hoc Tukey test</th>
</tr>
</thead>
<tbody>
<tr>
<td>n Ø (mm)</td>
<td>2.05 (0.6)</td>
<td>2.66 (0.6)</td>
<td>2.65 (0.7)</td>
<td>a</td>
</tr>
<tr>
<td>Length (cm)</td>
<td>5.62 (2.7)</td>
<td>6.12 (2.4)</td>
<td>7.97 (3.1)</td>
<td>a</td>
</tr>
<tr>
<td>Number of leaves</td>
<td>8.3 (2.7)</td>
<td>10.07 (2.9)</td>
<td>9.67 (2.9)</td>
<td>a</td>
</tr>
<tr>
<td>Leaf dry mass (g)</td>
<td>0.16 (0.08)</td>
<td>0.25 (0.15)</td>
<td>0.23 (0.15)</td>
<td>b</td>
</tr>
<tr>
<td>Stem dry mass (g)</td>
<td>0.12 (0.09)</td>
<td>0.19 (0.15)</td>
<td>0.22 (0.15)</td>
<td>c</td>
</tr>
<tr>
<td>LMA (ng cm⁻²)</td>
<td>94 (7.18)</td>
<td>95 (9.80)</td>
<td>93 (4.93)</td>
<td>ns</td>
</tr>
<tr>
<td>n – 1 Number of ramifications</td>
<td>3.67 (2.34)</td>
<td>5.17 (3.05)</td>
<td>6.87 (3.01)</td>
<td>a</td>
</tr>
<tr>
<td>Ø (mm)</td>
<td>2.78 (0.73)</td>
<td>3.45 (0.70)</td>
<td>3.68 (0.93)</td>
<td>a</td>
</tr>
<tr>
<td>Length (cm)</td>
<td>4.19 (2.4)</td>
<td>5.53 (3.0)</td>
<td>6.87 (3.1)</td>
<td>b</td>
</tr>
<tr>
<td>Leaf dry mass (g)</td>
<td>0.21 (0.22)</td>
<td>0.35 (0.25)</td>
<td>0.56 (0.44)</td>
<td>c</td>
</tr>
<tr>
<td>n – 2 Ø (mm)</td>
<td>3.59 (0.96)</td>
<td>4.11 (0.89)</td>
<td>4.43 (1.14)</td>
<td>a</td>
</tr>
<tr>
<td>Length (cm)</td>
<td>3.84 (2.7)</td>
<td>4.59 (2.8)</td>
<td>5.34 (3.9)</td>
<td>b</td>
</tr>
<tr>
<td>Stem dry mass (g)</td>
<td>0.39 (0.48)</td>
<td>0.54 (0.4)</td>
<td>0.76 (0.89)</td>
<td>b</td>
</tr>
<tr>
<td>Sum Total length (cm)</td>
<td>13.65 (6.42)</td>
<td>16.24 (5.81)</td>
<td>20.18 (8.2)</td>
<td>a</td>
</tr>
<tr>
<td>Total stem dry mass (g)</td>
<td>0.72 (0.68)</td>
<td>1.08 (0.63)</td>
<td>1.54 (1.38)</td>
<td>a</td>
</tr>
<tr>
<td>Total stem volume (cm³)</td>
<td>1.10 (1.2)</td>
<td>1.69 (1.1)</td>
<td>2.57 (2.6)</td>
<td>b</td>
</tr>
</tbody>
</table>
Kermes Oak, a non-storing monoterpenoid emitter. Interestingly, they observed that Kermes Oak was the only species displaying delayed seasonal emission tendencies while _Q. ilex _rather showed the highest emission rates in spring and the lowest in autumn. The increase in emission rate we observed in autumn may be related to some phenological events such as second growth flushes occurring with the first rain episodes in late summer. Fischbach et al. (2002) showed that monoterpenoid-synthesizing activity was 67% higher in newly developed leaves of _Q. ilex _than in one-year-old leaves. Hence, June and November could correspond to periods of strong emissions due to a higher monoterpene synthase activity in recently-developed leaves.

The high _E_ observed in autumn may also be due to the high uncertainty of the G93 algorithm under low temperatures (Simon et al., 2001). Indeed, normalization increased the emission variability among replicates measured in the cold season, while it decreased the emission variability in the warm season. Boissard et al. (2001) evaluated the accuracy of the G93 model on isoprene emissions from _Ulex europaeus_ through the seasons and revealed an overestimation of _E_ in the cool periods from early November to late January. The normalization with G93 is more accurate when light and temperature are close to the standard conditions (30 °C and 1000 μmol m⁻² s⁻¹). The rise in emission rate we observed in November after normalization was also noticed by Sabillon and Cremades (2001) on _Q. ilex_, while the initial field emission rate they measured was relatively stable throughout the cold season.

### 4.2. Compost effect on soil and leaf nutrients

Due to its strong nutritive properties, compost was expected to improve soil fertility in the spread plots. However, only mitigated changes in soil nutrient content were observed, two years after spreading (n.b., on the same experimental site. N and P soil content were found to have durably increased only from 7 years after spreading; Olivier et al., in prep.). Consistently with Larchevêque et al. (2006a), we did not observe any increase in soil N content during the two years following soil amendment and therefore no enhancement in leaf N content. The compost used was a mix of sewage sludge and other organic wastes with high C/N ratios (such as green wastes) which generally reduce the rate of nitrogen mineralization and provide slow release to the plants (Epstein _et al._, 1978). Besides, Kermes Oak has a relatively deep root system (Baquedano and Castillo, 2007) and we did not observe any fine root development in the surface compost layer. Therefore, nutrients had to percolate to deep mineral soil layers to be available for plants. Only sporadic enrichments in available phosphorus were found within the soil profile, probably resulting from rainy episodes favoring its leaching. In contrast to nitrogen and phosphorus, potassium percolation was much quicker and enabled a durable enrichment of the mineral soil layer, but led to enhancement of the leaf potassium content from June 2009 only.

### 4.3. Compost did not have direct effect on monoterpene emissions

Compost supply is generally expected to induce changes in soil nutritive and water status (Gallardo-Lara and Nogales, 1987; Martínez _et al._, 2003). Various fertilization effects on BVOC emissions have been described in the literature (Litvak _et al._, 1996; Possell _et al._, 2004; Blanch _et al._, 2007; Fares _et al._, 2008), but all of them are related to alterations in leaf nutrient concentrations. In the time frame of our study, going up to two years after spreading, little change in leaf nutrient concentrations was induced by the compost, which explains why no clear alteration of the foliar BVOC emissions was observed.
In the Mediterranean region, summers are often stressful periods for plants due to high temperature and irradiance levels, and strong water deficit. Low water availability is known to negatively affect monoterpene emissions from *Q. ilex* if drought is very severe (Staudt et al., 2002; Lavoir et al., 2009). In our case, predawn xylem water potential remained clearly above −2 Mpa, considered as a critical threshold (Lavoir et al., 2009) for BVOC emissions from *Q. ilex* under natural conditions. Compost did not induce any visible change in the plant’s summer water status (*P* > 0.05), as observed by Larchévéqué et al. (2009) at the same study site, one year after spreading. Mineral soil probably benefited from enhanced water retention in the first centimeters only, while deep soil layers penetrated by Kermes Oak root system may have remained unaffected by the compost spreading. Indeed, even though compost probably limited soil water evaporation, water uptake by long-root species such as Kermes Oak depends more on precipitation and water stored at depth than on surface water evaporation (Baquedano and Castillo, 2007). Consequently, the positive effect of compost on soil water retention was probably too weak to induce clear changes in BVOC emissions.

The increase of *E*₀ with compost observed in early June 2008 was probably related to better photosynthetic performance. Indeed, in early June 2008, the increase of *Fv/Fm* with compost was significant and comparable to that of monoterpene emissions. Emissions of monoterpenes from non-storing species were shown to be correlated with photosynthetic parameters (Niinemets et al., 2002). Compost may have induced earlier leaf development in April–May, resulting in a higher photochemical efficiency at the beginning of June 2008, while leaves from control treatments could have been mature but not yet fully photosynthetically active. Even if the whole photosynthetic machinery is already present in the leaf, it may take several weeks before this leaf reaches its maximum photosynthetic capacity (Both and Brüggemann, 2009).

On the other hand, potassium, the only element significantly supplied by the compost so far, could also explain the *Fv/Fm* and *E*₀ increase in early June 2008. Potassium is known to have essential functions in osmotic regulation but also favorable effects on CO₂ assimilation (Mengel and Kirkby, 2001). Moreover, it was shown to enhance protection against high solar irradiance levels and increase plant resistance to drought (Ismail, 2005), both conditions being typical of summers under Mediterranean climate.

### 4.4. Indirect effects of compost spreading on monoterpene emissions

Although compost did not affect leaf nutrient content, it enhanced plant growth in most of the investigated variables. This biomass increase could also explain why leaf nutrients were not much enriched in the treated plots. The higher nutrient pools available to the plants were probably used in higher biomass production (Chapin, 1980). Over two years, compost increased the wood biomass, thus allocating nutrients to non-emitting organs which were not investigated in this study. Moreover, spreading enhanced the number of ramifications and the leaf dry mass, resulting in larger leaves, since LMA was not affected. As a consequence, at plot scale, the first effect of compost on Kermes Oak BVOC emissions appeared indirectly through this annual increase of emitting biomass. This indirect effect has to be taken into account since its impact on regional-scale emissions is independent of climatic conditions and constant all the year long.

### 5. Conclusion

Emissions of monoterpene from *Q. coccifera* were assessed in a Mediterranean shrubland after compost spreading. Variable normalized emission rate was observed, depending on the season, with higher rates in the heart of summer and in autumn than during the rest of the year. Our data showed that emission normalization with Guenther’s algorithm does not always reduce measure variability, notably in autumn and winter, seasons for which the model seemed to be inappropriate.

Two years after spreading, compost did not directly affect BVOC emissions at leaf scale, but indirectly enhanced annual emissions by increasing emitting leaf biomass. Further measurement on the experimental site will allow assessment of the longer-term effect of this amendment, and the expected plant nutrition enhancement with the advanced mineralization of the compost.

Even if compost spreading in natural lands can be an interesting recycling outlet for sewage sludge production, the impact of this practice on BVOC emitting species has to be considered since it indirectly contributes to regional scale ozone pollution. Biogenic emissions are now recognized as one of the major sources of tropospheric ozone. This is particularly relevant in the Mediterranean region which is, each summer, one of the European zones most exposed to ozone pollution. In this season and at this location, several of the factors contributing to ozone accumulation are combined: BVOC emitting species, high temperatures increasing emissions, high irradiance levels contributing to ozone formation by allowing atmospheric photochemical reactions, and high vehicle traffic — in a region that is a frequent holiday destination — leading to enhanced production of ozone precursors.

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