Contribution of flowering trees to urban atmospheric biogenic volatile organic compound emissions

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Abstract. Emissions of biogenic volatile organic compounds (BVOC) from urban trees during and after blooming were measured during spring and early summer 2009 in Boulder, Colorado. Air samples were collected onto solid adsorbent cartridges from branch enclosures on the tree species crabapple (Malus sp.), horse chestnut (Aesculus carnea, “Ft. McNair”), honey locust (Gleditsia triacanthos, “Sunburst”), and hawthorn (Crataegus laevigata, “Pauls Scarlet”). These species constitute ∼65% of the insect-pollinated fraction of the flowering tree canopy (excluding catkin-producing trees) from the street area managed by the City of Boulder. Samples were analyzed for C10–C15 BVOC by thermal desorption and gas chromatography coupled to a flame ionization detector and a mass spectrometer (GC/FID/MS). Identified emissions and emission rates from these four tree species during the flowering phase were found to vary over a wide range. Monoterpene emissions were identified for honey locust, horse chestnut and hawthorn. Sesquiterpene emissions were observed in horse chestnut and hawthorn samples. Crabapple flowers were found to emit significant amounts of benzyl alcohol and benzaldehyde. Floral BVOC emissions increased with temperature, generally exhibiting exponential temperature dependence. Changes in BVOC speciation during and after the flowering period were observed for every tree studied. Emission rates were significantly higher during the blooming compared to the post-blooming state for crabapple and honey locust. The results were scaled to the dry mass of leaves and flowers contained in the enclosure. Only flower dry mass was accounted for crabapple emission rates as leaves appeared at the end of the flowering period. Total normalized (30°C) monoterpene emissions from

1 Introduction

Volatile organic compounds (VOC) play an important role in atmospheric chemistry, in particular in secondary pollutant formation (e.g., tropospheric ozone and secondary organic aerosols) (Andreae and Crutzen, 1997). Biogenic VOC emissions (BVOC) from vegetation comprise ∼90% of global terrestrial non-methane VOC emissions annually (Fuentes et al., 2000). BVOC emissions are highly dependent on vegetation activity and biomass, which are both governed by seasonal changes. BVOC flux estimates used in air quality models rely on emission rates, vegetation characteristics,
and weather conditions. Seasonal changes are not well understood and are thus difficult to account for. In particular, BVOC emissions from flowers during the blooming period are generally not considered. Ornamental vegetation with colorful and/or fragrant flowers is often planted to brighten up urban areas. This general vegetation group contributes to VOC concentrations to an extent that could potentially impact air quality. To our knowledge only three studies in the literature have addressed the possibility of increased atmospheric VOC during the blooming period (Arey et al., 1991; Ciccioli et al., 1999; Müller et al., 2002). Flower volatiles have been widely studied in botanical research for their role in plant-pollinator and plant-herbivore interactions. Therefore, an abundant body of literature on BVOC emissions from flowers is available. However, most studies on floral BVOC emissions report only qualitative results. A summary of this literature is presented in Table 1 in the Supplement. A few studies report emission rates, but it is difficult to incorporate them into quantitative models. For example, Baraldi et al. (1999) and Rapparini et al. (2001) report emission rates in $\mu$g per 100 flowers h$^{-1}$. Ibrahim et al. (2010) detected significantly higher isoprene and monoterpane emission rates from flower branches compared to vegetative branches of Yeheb tree (*Cordeauxia edulis*) and express these findings in $\mu$g (g dry weight)$^{-1}$ h$^{-1}$. Müller et al. (2002) reported terpene fluxes from flowering rape plants of 16–32 $\mu$g m$^{-2}$ (ground area) h$^{-1}$, and Rodriguez-Sanoa et al. (2011) observed a mean volatile production of 48.6 $\mu$g h$^{-1}$ per flower of blueberry.

The review of this literature demonstrates that there is a significant amount of BVOC emitted by flowers and that there is a high variability in emissions. There seems to be a common pattern in that there are higher VOC emission rates from flowering branches than vegetative branches (Arey et al., 1991; Ibrahim et al., 2010). These issues motivated us to study the floral emissions from urban trees in the City of Boulder and their contribution to the urban BVOC flux to assess the potential effect of flowering tree emissions on urban air quality.

2 Experimental

2.1 Site description

The field site was located in the CreekSide Tree Nursery in Boulder, Colorado, USA. An enclosed trailer sitting within the tree nursery was used as a mobile field laboratory. Flowering trees were provided by the nursery during the measurement period corresponding to the blooming season. Four tree species were studied: crabapple (*Malus* sp.), hawthorne (*Crataegus laevigata*, “Pauls Scarlet”), horse chestnut (*Aesculus carnea*, “Ft. McNair”), and honey locust (*Gleditsia triacanthos*, “Sunburst”). Together, honey locust, crabapple and hawthorn represent ~65% of the insect pollinated, non-catkin-producing flowering tree canopy in the City of Boulder according to a municipal tree resource analysis report from the City of Boulder (City of Boulder, 2005; see Supplement Table 2), while horse chestnut represents less than 1%. Only those flowering species present in the urban tree inventory that produce conspicuous, showy floral structures and are assumed to be insect pollinated were considered for screening. The rationale for this sampling decision was that these tree species would be most likely to invest resources into floral BVOC production for the purpose of pollinator attraction. The remaining 35% of the insect-pollinated, non-catkin-producing flowering tree species in Boulder is comprised of 9 species, which were not screened during the course of this study, in part because of limitations in the number of branch enclosure experiments that could be operated simultaneously at the time of the study, and also because specimens were not available for screening at the tree nursery during the sampling campaign. All investigated trees were between 2 and 3 m tall and between 3 and 5 yr old. They were kept in their planting pots during the experiments.

2.2 Sampling

Sampling methods and materials used for the BVOC emission measurements followed the procedures recommended in Ortega and Helmig (2008). The studied vegetation was enclosed in a Tedlar bag with minimal contact of foliage and flowers with the bag. The bag was attached at the branch base with a Velcro strap. Ambient air was filtered for particles (respirator filter, Mersorb, Part no. 463532, Mine Safety Appliances Company, Pittsburgh, PA) and scrubbed of ozone (cartridge with MnO$_2$-coated screens, O.B.E. Corp. Fredericksburg, TX) and pumped at 251 min$^{-1}$ into the enclosure providing a slight overpressure inside the bag. The turnover time inside the enclosure was estimated as 2.0 to 2.5 min. A cold trap was used to reduce moisture and cool the air pumped into the enclosure. A small flow of a 5-component reference standard mixture, spanning a wide volatility range was doped into the purge air to serve as a reference for tracing compound recovery rates from the experiment (Ortega and Helmig, 2008). The standard was composed of 5 aromatic compounds including toluene, isopropylbenzene, tetrahydroxaphthalene, tri-isopropylbenzene and n-nonylbenzene, and concentrations were determined using a gas chromatograph calibrated with a multi-component-alkane gas standard as described in Helmig et al. (2003). The aromatic mixture was introduced in the main purge flow at a mass flow-controlled rate of 6.5 ml min$^{-1}$. The resulting mixing ratio of the five compounds ranged from 2.2 to 11.4 ppbv. Two automated sampling devices (Helmig et al., 2004a) were used for collection of emission samples, allowing for 10 to 20 samples to be collected sequentially. An ozone scrubber composed of a Na$_2$S$_2$O$_3$-coated glass-fiber filter (Helmig et al., 2006) was placed at the auto-sampler inlet. Samples were collected onto glass tubes filled with a
multi-adsorbent bed composed of half TenaxGR (Buchem BV, Apeldoorn, The Netherlands) and half Carboxen 1016 (Supelco, Bellefonte, PA, USA). A second adsorbent cartridge (breakthrough cartridge) was placed in series with every 10th sample cartridge. Analysis of the breakthrough cartridges was used to ensure that sampled compounds did not make their way through the primary adsorbent cartridge. Sampling flow was 200 ml min\(^{-1}\) for a typical sample volume of 121.

One tree of each species was selected for this study. For each tree, a single branch was chosen to be sampled repeatedly over the course of the study. The enclosure was installed as carefully as possible to minimize any disturbance effects. Figure 1 shows a picture of the enclosure setup for one of the crabapple experiments. Sampling on a branch enclosure started no sooner than six hours after the enclosure was installed. This delay allowed the plant and the enclosure to reach an equilibrium state, and also allowed for any stress-induced emissions caused by the installation of the enclosure to subside (Tarvainen et al., 2005). Sampling times varied from one hour during daytime to 2 h during nighttime. Two sets of enclosure equipment were used simultaneously when several trees were blooming at the same time. Approximately 400 samples were collected over the course of this study, from 16 April to 19 June 2009: 99 samples on hawthorn, 40 on crabapple, 145 on chestnut, and 70 samples on honey locust. In addition, 20 samples of the inlet purge air were collected to allow for a comparison of the aromatic reference standard recovery rates from the experiment. For each tree a second set of emission samples was collected once flowers had withered to obtain reference measurements of the foliar BVOC emissions from each species after the flowering period had ended.

Samples collected at the field site were brought back to the laboratory and stored in a freezer for 1 to 2 days before being analysed. Thermal desorption using a Perkin-Elmer ATD 400 allowed the sample to be transferred from the cartridge to a GC (Model 5890, Hewlett-Packard). Separation was achieved on a 0.32 mm i.d., 50 m long, 5 µm film thickness DB-1 capillary column (Agilent). Analyte identification and quantification were performed using a mass spectrometer (Model 5970, Hewlett-Packard) and flame ionization detector (FID) after splitting the column flow. Similar instrumentation and calibration procedures have been described previously (Helmig et al., 2004b; Ortega et al., 2008). Compound identification was achieved using relative retention times, along with comparing mass spectra with the NIST database and Adams (1989). All quantifications were performed using the FID signals. The typical limit of detection of the system during this study was 50 pptv and the typical accuracy was \(\sim 10\%\). Isoprene (which previously has not been reported as a major floral emission) was not quantified as it was not sufficiently retained under the chosen GC conditions. Quantitative results are reported in mass of carbon (C).

![Fig. 1. Branch enclosure experiment setup on a crabapple tree during its flowering period.](image-url)
ambient air temperature. A thermocouple was placed in each bag to monitor the temperature inside the enclosure. BVOC and particularly sesquiterpenes are very reactive. It was therefore important that oxidant levels inside the enclosure were kept low. Ozone concentrations in the enclosure were measured regularly, and were usually below 2 ppbv. Environmental data were acquired at 1 s and averaged and recorded every 5 min using a Campbell Scientific CR10X datalogger (Campbell Scientific, Logan, UT, USA).

2.5 Normalized emission rate calculations

Temperature dependencies for emissions of monoterpenes, benzyl alcohol and benzyl aldehyde, oxygenated monoterpenes, and sesquiterpenes were observed in this study. There was no obvious light dependency in observed emissions. Normalized emission rates, referred to here as basal emission rates (BER), were calculated for conditions of $T_i = 30\,^\circ\mathrm{C}$ using the algorithm of Guenther et al. (1993); $E = \text{BER} \times \text{exp}(\beta(T - T_i))$, where $E$ is emission rate and $T$ is leaf temperature. $\beta$ was derived from experimental results for each tree. The results for the flower dry weight and leaf biomass determination were used to normalize emission rates to the amount of biomass inside the enclosure. Since leaves had just started to develop on the crabapple by the end of the flowering period, only flower petal dry weight was considered for normalizing emission rates. For the control experiments conducted after the flowering had subsided, the leaf biomass inside the enclosure was used to normalize emission rates.

2.6 Modeling

The Model of Emissions of Gases and Aerosols from Nature version 2.0 (MEGANv2.0) was applied for conditions in Boulder, Colorado, USA (latitude 40.0° N, longitude 105.3° W, elevation 1600 m), which is located at the border of the Rocky Mountain and Central Plains ecoregions. Floral and foliar BVOC emissions were estimated using a single grid version of MEGAN (Guenther et al., 2006), which is driven by both land cover data and environmental conditions. Land cover inputs included the City of Boulder tree inventory and an urban tree cover fraction estimate. Tree cover was estimated by analyzing high resolution land surface imagery using the approach described by Duhl et al. (2011), whereby the Boulder area was divided into 900 cells (in this case, we selected a cell size of 10 s by 10 s; $\sim 0.73 \, \text{km}^2$). 350 cells located in residential, commercial/industrial, and city park land use classes within the city limits were randomly evaluated; average tree cover fraction was found to be 0.15. Tree species information for Boulder was derived from a municipal tree resource analysis report by Boulder foresters (City of Boulder, 2005). Since there were no data available for the tree species composition on private property, for the purpose of this study it was assumed to be the same as on city property and BVOC emissions were assumed to be the same on public and private land throughout the city. Monoterpene emissions were calculated for April to June 2009. Temperature and solar radiation inputs for MEGAN were obtained from a University of Colorado weather station (http://foehn.colorado.edu/weather/atoc1/). Observed floral emission rates were included in the model for honey locust and crabapple using the experimental results of this study, as together these two species represent $\sim 64\%$ of the fraction of insect-pollinated flowering street trees in Boulder that produce conspicuous flowers, and $> 13\%$ of total street trees (by number) according to the tree inventory. Hawthorn and horse chestnut trees did not exhibit significantly different emissions during flowering as compared to non-flowering emissions and were therefore excluded from the simulations.

3 Results and discussion

Blooming occurred at different times for each tree species and lasted between 7 and 18 days. Figure 2 shows the flowering period (shaded) along with the sampling schedule for each species.

3.1 Emission results

BVOC emission rates varied significantly between vegetation species. Identified floral emissions showed a strong dependence on temperature and consequently were much higher during the day. The following section provides results for each of the species investigated.

3.2 Crabapple

Crabapple flowers appeared first, in late April, at a time when leaves were still in the bud stage. Benzyl alcohol and benzaldehyde were the only volatiles identified from branch enclosure samples. Figure 3 shows emission rate results for a typical day for each compound and Fig. 4 depicts the temperature dependence of both compounds.

The averaged contribution of benzyl alcohol and benzaldehyde to the overall emissions was 94 % and 6 %, respectively. The averaged total normalized emission rate during the flowering period was 93 $\mu$gC g$^{-1}$ h$^{-1}$ (standard deviation $\sigma$, determined by repeated experiments on the same branch, was 30 $\mu$gC g$^{-1}$ h$^{-1}$). The emissions of both compounds exhibited clear temperature dependencies as presented in Fig. 4, which is similar to the dependencies observed for foliage terpenoid emissions. Emissions of both compounds dropped to near detection limits after the flowers had withered (Fig. 5). This finding shows that VOC emissions from crabapple are dominated by compounds released during the blooming stage. The emissions recorded from crabapple during the flowering period are among the highest BVOC emission rates reported from vegetation, i.e. these rates are on the same order as isoprene emission rates from
3.3 Horse chestnut

Compounds identified in the horse chestnut branch enclosure samples included thujene, \( \alpha \)- and \( \beta \)-pinene, camphene, \( \beta \)-myrcene, \( \alpha \)- and \( \gamma \)-terpinene, o-cymene, \( \beta \)-phellandrene, terpinolene, 4-terpineol, methyl salicylate, \( \alpha \)-terpineol, and limonene.

Figure S1 in the Supplement shows emission rate results for a typical day for the major compounds emitted. Normalized emission rates were calculated using an empirically-derived \( \beta \)-factor of 0.20 for the flowering period and 0.21 after flowering (Fig. 6). The average total basal BVOC emission rate during flowering was 9.1 \( \mu \text{gC g}^{-1} \text{ h}^{-1} \) \( (\sigma = 3.0 \mu \text{gC g}^{-1} \text{ h}^{-1}) \), whereas after the flowering period emissions reached 12 \( \mu \text{gC g}^{-1} \text{ h}^{-1} \) \( (\sigma = 6.3 \mu \text{gC g}^{-1} \text{ h}^{-1}) \) (Fig. 7). \( \alpha \)-pinene, \( \beta \)-pinene, and limonene were the most abundant individual compounds emitted. The data in Figs. 6 and 7 show no significant change in the emission rates of these compounds during the blooming stage as compared to afterwards. However, the compounds terpinolene, 4-terpineol, methyl salicylate and \( \alpha \)-terpineol show post-blooming increases.
3.4 Honey locust

Forty samples were collected during the flowering period of the honey locust tree and twenty more samples were taken after flowering. Monoterpenes and oxygenated monoterpenes were identified, including α-thujene, α- and β-pinene, β-myrcene, α-cymene, β-phellandrene, limonene, γ-terpinene and α-terpineol. The average normalized total MT emission rate during the flowering period was 5.3 µgC g⁻¹ h⁻¹ (σ = 3.5 µgC g⁻¹ h⁻¹) with α-pinene, β-pinene, and limonene being the dominant BVOC (Fig. 8). Figure S2 in the Supplement shows the temperature dependence of the emissions. After flowering, average emission rates dropped to 1.2 µgC g⁻¹ h⁻¹ (σ = 0.85 µgC g⁻¹ h⁻¹) (Fig. 9); γ-terpinene, β-phellandrene, and limonene were the major BVOC emitted at this time.

3.5 Hawthorn

Two sesquiterpenes, β-caryophyllene and humulene were identified in the hawthorn emissions during flowering (Fig. S3, in the Supplement). Averaged BER for β-caryophyllene and humulene were 15.1 ngC g⁻¹ h⁻¹ (σ = 18 ngC g⁻¹ h⁻¹) and 4.4 ngC g⁻¹ h⁻¹ (σ = 5.6 ngC g⁻¹ h⁻¹), respectively (Fig. S4, in the Supplement).

These compounds were also identified in the later foliage emissions. The post-flowering average emission rates were 4.0 ngC g⁻¹ h⁻¹ (σ = 6.9 ngC g⁻¹ h⁻¹) and 0.5 ngC g⁻¹ h⁻¹ (σ = 0.8 ngC g⁻¹ h⁻¹) for caryophyllene and humulene, respectively. A third sesquiterpene, α-farnesene, and the monoterpane limonene, neither of which were observed during flowering, were detected in the hawthorn emissions after flowering.

3.6 Modeling results

Figure 10 presents the results of the MEGAN model simulations performed for the 1 April–30 June 2009 simulation period. For the MT time series, emissions were calculated using default MEGAN species-specific MT emission rates for all tree species in the City of Boulder tree inventory. The floral time series incorporates BER as determined in the present study. Floral emissions can be substantial during periods of flowering. Overall, floral BVOC emissions are equivalent to 11 % of MT emissions integrated over the three months simulation period. During the crabapple blooming period, emissions from flowers were as high as 116 % of the estimated
non-flower (foliage) MT emissions (Fig. 11). Honey locust flowers appear later in the season (Fig. 12); during that time these emissions represent 36% of MT emissions.

The simulated estimates have high associated uncertainties due to assumptions made, such as using the city-maintained tree inventory to represent tree species composition across the entire domain. It should also be recognized that the contribution of floral emissions to total biogenic emissions within other urban areas may be quite different from what was observed in Boulder. As a comparison, in the southern US city of Riverside, California, flowering trees represent 40% of total street trees with more than 10 different flowering tree species according to the City of Riverside urban tree inventory. It should also be noted that floral emissions within a city may be clustered in some regions, such as city parks and specific neighborhoods, and so could make a higher contribution in some areas and a lower contribution in others. Finally, we would like to emphasize that although in the present analysis a direct comparison of MT emissions from honey locust flowers against simulated leaf MT emissions is made, this is not the case for crabapple, in which the dominant BVOC emitted (benzyl alcohol and benzaldehyde) are oxygenated compounds. Species-specific emission factors for oxygenated compounds have a high degree of uncertainty, and as such, we felt it prudent to instead compare our simulated floral emission estimates for these compounds with simulated MT emissions, which have substantially lower uncertainties associated with their emission factors.

4 Conclusions

This study represents an attempt to assess the significance of urban tree floral BVOC emissions at an urban scale. The findings demonstrate that floral BVOC emissions may constitute a significant seasonal source of BVOC in areas containing high proportions of flowering trees, with floral emissions approaching and possibly even exceeding non-isoprene BVOC emissions during the peak flowering period. These emissions estimates are based on the survey of ~65% of the insect-pollinated flowering tree species present in the tree inventory and did not include catkin-producing species. Therefore, reported emissions should be considered lower-bounds of the likely actual emissions from urban trees within the study domain. These estimates suggest that floral tree emissions should be more closely examined and possibly considered in emissions models. To that end, efforts should be made to better understand whether municipal tree inventories are representative of the overall species compositions present in a given urban area.

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composition in urban settings. Additional screening experiments of important flowering tree species should also be conducted and the temporal, species-specific blooming patterns of these trees should be characterized to better quantify the significance of floral emissions in locations which contain substantial fractions of flowering species. The timing of many phenological processes (such as bud-burst, flowering, etc.) may shift in response to global climatic changes. In addition to ecological repercussions associated with climatic shifts (such as chemically mediated biological interactions between plants and their pollinators or even herbivores), changes and shifts in floral BVOC emissions might occur. The potential also exists for the timing of these floral emissions bursts to coincide with the Northern Hemisphere springtime ozone maximum, a scenario with implications for urban air quality (due to the reactive nature of many BVOC chemical species); all of these possibilities highlight the need for further research in this area.

Supplementary material related to this article is available online at: http://www.biogeosciences.net/9/3777/2012/bg-9-3777-2012-supplement.pdf.

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