Volatile Organic Compound Emissions from *Miscanthus* and Short Rotation Coppice Willow Bioenergy Crops

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**Highlights**

- Miscanthus and coppice willow are increasingly important bioenergy crops.
- Above-canopy fluxes were measured using PTR-MS and virtual disjunct eddy covariance.
- Willow isoprene emission peaked at \(\sim 1 \text{ mg m}^{-2} \text{ h}^{-1}\) = \(20 \mu\text{g g}_{\text{dw}}^{-1} \text{ h}^{-1}\) standardised.
- Bioenergy crop species choice should consider their impact on regional air quality.
Abstract

*Miscanthus x giganteus* and short rotation coppice (SRC) willow (*Salix spp.*) are increasingly important bioenergy crops. Above-canopy fluxes and mixing ratios of volatile organic compounds (VOCs) were measured in summer for the two crops at a site near Lincoln, UK, by proton transfer reaction mass spectrometry (PTR-MS) and virtual disjunct eddy covariance. The isoprene emission rate above willow peaked around midday at ~1 mg m$^{-2}$ h$^{-1}$, equivalent to 20 µg g$_{dw}$ h$^{-1}$ normalised to 30°C and 1000 µmol m$^{-2}$ s$^{-1}$ PAR, much greater than for conventional arable crops. Average midday peak isoprene mixing ratio was ~1.4 ppbv. Acetone and acetic acid also showed small positive daytime fluxes. No measurable fluxes of VOCs were detected above the *Miscanthus* canopy. Differing isoprene emission rates between different bioenergy crops, and the crops or vegetation cover they may replace, means the impact on regional air quality should be taken into consideration in bioenergy crop selection.

**Keywords:** VOC, isoprene, bioenergy, *Miscanthus*, willow, eddy covariance
1 Introduction

Bioenergy crops are those grown specifically for energy production rather than food, as a means of mitigating two problems associated with the use of traditional fossil fuels: anthropogenic climate forcing and energy security (McKay, 2006). Such crops contribute to carbon neutrality since CO$_2$ produced during the combustion of the crop is offset by the CO$_2$ sequestered during growth. There is also potential for long-term storage of carbon via uptake by soil through plant roots (Grogan and Matthews, 2002). Consequently, cultivation of bioenergy crops is increasing rapidly. For example, power generators in the UK are required to increase to 15.4% by 2015/16 the energy derived from renewable sources (DTI, 2005), with biomass being acknowledged as a key resource in achieving this target.

Although bioenergy crops are perceived to be carbon neutral, full life-cycle analysis needs also to take account of changes in emissions of other potent greenhouse gases such as CH$_4$ or N$_2$O. Also, few studies have investigated volatile organic compound (VOC) emissions from bioenergy crops. Biogenic VOC emissions from vegetation (Steiner and Goldstein, 2007) are estimated as about 10 times greater globally than VOC emissions from anthropogenic sources (Guenther et al., 1995). The dominant BVOC is isoprene (Guenther et al., 2006), but other important compounds include oxygenated VOCs and terpenoids.

Emissions of VOCs are important for several reasons. Their rapid oxidation chemistry, particularly in the presence of NO$_x$, affects the oxidative capacity of the atmosphere, the generation of tropospheric ozone (Atkinson, 2000), of concern for human and plant health (Ashmore, 2005) and as a radiative forcing gas, and on formation of secondary organic
particles, which likewise affect human health (Dockery et al., 1993) and radiative forcing (Kulmala et al., 2004).

The potential for BVOC emissions from crops to have a significant impact on atmospheric composition has been demonstrated in the tropics (Hewitt et al., 2009). The aim of this study was to determine fluxes of BVOCs for two bioenergy crops grown in the UK and elsewhere: short rotation coppice (SRC) willow (Salix spp.), a woody crop grown in dense plantations of multi-stemmed plants and harvested every 3 years; and Miscanthus x giganteus, a perennial grass native to Asia, of the same taxonomic group as sugarcane, sorghum and maize (Naidu et al., 2003) but more resilient to lower temperature whilst maintaining high CO₂ assimilation and biomass conversion efficiency. The crop grows up to 3.5 m per year (Rowe et al., 2009), and is harvested annually between January and March. The chipped and dried biomass of both crops is used to fuel biomass burners or to co-fire existing coal-fired power stations.

Fluxes from this work are compared with those for conventional UK arable crops to assess the potential impact of this land-use change on atmospheric chemistry.

2 Methods

2.1 Sampling site

The field measurements were carried out from mid July to mid August 2010 near Lincoln, UK (53° 19’ N, 0° 35’ W). Figure 1 shows the layout of the site, which consisted of several fields of Miscanthus, willow and wheat, located within an area of predominantly flat arable fields separated by hedgerows and isolated areas of mixed deciduous woodland. Mean annual
rainfall at the site was 600 mm and the soil was a fine loam, overlying Charnmouth mudstone. The nearest settlement (population: 113), which had a relatively busy through-road, was ~0.7 km to the southeast. A minor road running east-west was situated 0.3 km to the south.

The *Miscanthus* plot (~11 ha, planted in spring 2006) was surrounded by the following vegetation types: hedgerow and wheat to the north; willow to the east; deciduous trees to the west; willow and wheat to the south. Crop height was typically 2.5 m. Sampling was carried out from 16th July to 2nd August 2010, near the NE corner of the plot, downwind of the prevailing wind direction, at an inlet height of approximately 4 m.

The willow plot (~6.5 ha, planted in 2001 with five different genotypes) was bounded as follows: a row of deciduous trees and a ploughed field to the north; *Miscanthus* to the west; mixed deciduous woodland to the south; wheat to the east. Typical canopy height was 4 m. Sampling was carried out from 5th to 13th August 2010 at an inlet height of 6.7 m on the north edge of the field. Trees were planted in pairs of rows 1.3 m apart, with 0.6 m spacing within each pair.

Flux footprints for both sampling sites were predicted using a simple parameterisation model (Kljun et al., 2004). Model results are shown in Supplementary Information Figure S1. For the *Miscanthus* measurements, the largest distance for 80% flux contribution over the range of friction velocities encountered (122 m) was within the area of the *Miscanthus* field throughout the south-westerly sector (180 – 270°). For the willow measurements, the largest distance for 80% flux contribution (185 m) meant there may have been some small flux contributions from outside the willow field when wind was from the west. Flux contribution was otherwise within the willow field for the whole southerly wind sector (90 – 270°). These sectors were
used for directional filtering of data prior to deriving diurnal averages. Directionally-filtered data comprised 23% and 71% of all data for Miscanthus and willow, respectively.

Harvesting activities in surrounding farms during this study may have affected results, particularly during Miscanthus measurements, and are discussed later.

2.2 Proton transfer reaction mass spectrometer (PTR-MS)

BVOC mixing ratios and fluxes above both crop canopies were measured using proton transfer reaction mass spectrometry (PTR-MS) (Blake et al., 2009) coupled with virtual disjunct eddy covariance (vDEC) (Karl et al., 2002; Rinne et al., 2001). PTR-MS is a ‘soft’ chemical ionisation method in which hydronium ions (H$_3$O$^+$) formed in a hollow cathode ion source pass into a drift tube subject to an electric field ($E$) into which the ambient air is also introduced. As most VOC molecules have a proton affinity greater than water, they react with H$_3$O$^+$ ions to form protonated products, predominantly the protonated molecular ion, but also fragments or clusters. The extent of fragmentation/clustering can be controlled by tuning the $E/N$ ratio ($N$ is the H$_3$O$^+$ ion density).

The PTR-MS used in this study (Ionicon Analytik, Innsbruck, Austria) was fitted with an extra turbopump connected to the detection chamber, and Teflon instead of Viton rings in the drift tube (Davison et al., 2009; Misztal et al., 2010). Pfeiffer turbopumps replaced the Varian equivalents. The drift tube conditions were held constant throughout (pressure 2 mbar, temperature 40 °C, voltage 572 V) to maintain an $E/N$ ratio of ~130 Td (1 Td = 10$^{-17}$ V cm$^2$).

The sampling inlet and 20 Hz sonic anemometer (WindmasterPro, Gill Instruments) were positioned above the canopy using a telescopic mast. Air was sampled at 30 L min$^{-1}$ through a
20 m PTFE inlet line (1/4” OD, 3/16” ID) with a T-piece for sub-sampling into the PTR-MS inlet at a rate of 100 mL min\(^{-1}\). Condensation of water vapour in the inlet line was prevented by wrapping with self-regulating heating tape (Omega, UK type SRF3-2C). Data were logged using a program written in LabVIEW (Version 8.5, National Instruments).

2.3 Determination of VOC mixing ratios and fluxes

The PTR-MS signal was calibrated explicitly for several VOCs using a mixed gas calibration cylinder (Apel-Riemer Environmental Inc., USA) containing 1 ppmv each of formaldehyde, methanol, acetonitrile, acetone, acetaldehyde, isoprene and 0.18 ppmv d-limonene. The calibration gas was diluted with VOC-scrubbed air to produce 6 samples with concentrations of 0.5, 1.0, 10, 20, 30 and 50% of the pure calibration gas standard. A relative transmission curve was then constructed to determine empirical calibration coefficients for other VOCs under study not present in the standard (Taipale et al., 2008). Calibrations were carried out in the lab before commencement of the field campaign, and on 22\(^{nd}\) July during the campaign. Concentrations of gases in the calibration cylinder were verified using GC-MS calibrated with its own independent standards (details given in Section 2.4).

The PTR-MS was run in multiple ion detection (MID) mode for two 25 min sampling periods per hour. During these periods only the targeted VOC ions listed in Table 1 were measured, with dwell times of 0.5 s, in addition to the primary ion \(\text{H}_3\text{O}^+\), and water cluster \((\text{H}_2\text{O})\text{H}_3\text{O}^+\), which had dwell times of 0.2 s. The sensitivities (ncps ppbv\(^{-1}\)) and limits of detection (ppbv) for the target ions for the Miscanthus and willow campaigns are also included in Table 1. LODs were calculated as the ratio of twice the standard deviation of the background ion counts for a particular \(m/z\) throughout the campaign divided by the sensitivity (Karl et al., 2003).
The remaining 10 min per hour were used for full mass scans in the range 21 – 206 amu at a dwell time of 1 s per amu. For one 5 min period, ambient air was scanned to allow information about the full VOC composition to be acquired. For a further 5 min per hour, ‘zero air’ was scanned to determine the instrument background. Zero air was achieved by sampling ambient air through a custom-made zero-air generator comprising a glass tube packed with platinum wool and a 50:50 mixture of platinum mesh and activated charcoal heated to 200°C. The background spectrum was subtracted in subsequent processing of data.

As the PTR-MS was run in MID mode, fewer data points were generated than required for direct eddy covariance due to the non-continuous manner in which the quadrupole mass analyser measures each \( m/z \). The set-up resulted in 30,000 wind speed measurements and up to 441 VOC measurements in each 25 min sampling period (depending on how many VOCs were being measured). The total lag time between PTR-MS and wind speed data was determined by examining the cross-correlation between vertical wind speed and VOC mixing ratio as a function of lag time (with 15 s window). Total lag includes residence time in the sampling inlet line but also lag associated with collection and data writing of a full cycle of analysis by disjunct sampling and the response of the PTR-MS. The median of the lag times for each 5 min sub-period was used to calculate the flux in that 25 min period (Misztal, 2010). For example, the average lag-time for isoprene above the willow was 9.58 s, with a standard deviation between 25 min periods of 1.41 s. This method produced less variable lag times than those derived using the prevalent MAX method in cross-correlation function (~72% lower sd), and has been shown to be a practical alternative (Taipale et al., 2010).
Quality controls were used to filter data for periods of low friction velocity ($u^* < 0.15 \text{ m s}^{-1}$), non-stationarity, large spikes in vertical wind speed or VOC concentration, and where $< 20000$ data points were acquired in a 25 min sampling period. Most discarded data occurred during night when turbulence was low. High frequency flux losses due to the relatively slow disjunct VOC sampling frequency (0.25 Hz, compared to 20 Hz sonic data capture) were estimated using empirical ogive analysis (Ammann et al., 2006) for each 25 min period. At least 78% of flux was captured for all individual 25 min data periods, and values were corrected accordingly. Standard rotations of the coordinate frame were applied to correct for sonic anemometer tilt for each 25 min period separately.

2.4 Chromatographic analysis of ambient air samples

Ambient air samples were collected for chromatographic analysis, to confirm the identity of the VOC components measured by the PTR-MS, approximately hourly from 06:53 to 16:20 on 23 September 2010 above Miscanthus and from 06:32 to 17:30 on 11 August 2010 above willow (at ~1 m above the canopies). Sampling above Miscanthus was carried out at a later date because initial samples taken during the intensive campaign were lost due to GC-MS instrument failure. A mass-flow controlled Pocket Pump (210-1000 Series, SKC Inc.) was used to pump air at 100 mL min$^{-1}$ for 15 min through stainless steel adsorbent tubes (6 mm OD) packed with 200 mg Tenax TA and 100 mg CarboTrap (Markes International Ltd., UK). Prior to sampling, packed tubes were conditioned at 300 °C for 15 min with a flow of helium.

Analyses were carried out using a Hewlett-Packard 5890/5970 GC-MS with an automated thermal desorption unit (ATD 400, Perkin Elmer) connected via a 200 °C heated transfer line. Transfer of samples from the adsorbent tubes was performed in two steps: heat to 280 °C for 5 min at 25 mL min$^{-1}$ to desorb samples onto a Tenax-TA cold trap at −30 °C, followed by
transfer to the GC column at 300 °C for 6 min. Chromatographic separation utilised an Ultra-2 column (Agilent Technologies, 50 m × 0.2 mm ID × 0.11 μm film, 5% phenylmethyl silica) and a temperature program of 35 °C for 2 min, heat at 5 °C min⁻¹ to 160 °C, heat at 10 °C min⁻¹ to 280 °C, and hold for 5 min.

Calibration was carried out using a mixed monoterpe in methanol standard (10 ng μL⁻¹ α-pinene, β-pinene, α-phellandrene, 3-carene and limonene (Sigma Aldrich, UK)) and an isoprene in nitrogen gas standard (700 ppbv, BOC Gases, UK). Aliquots of the monoterpe standard (0, 1, 3 and 5 μL) were injected onto 4 adsorbent tubes with helium carrier gas. Tubes continued to be purged with helium for 2 min after the standard injection. Isoprene calibration tubes were prepared by slowly (over a period of about 2 min) injecting 0, 10, 30 and 50 mL of the gas standard onto 4 adsorbent tubes, while purging with helium. The limits of detection for isoprene, α-pinene and limonene were 0.16, 0.23 and 0.30 ng on column, corresponding to mixing ratios of 38, 27 and 35 pptv, respectively, for a 1.5 L sample.

2.5 Meteorological measurements

Photosynthetically active radiation (PAR), rainfall, temperature and relative humidity were available as part of long-term measurements at the site.

3 Results

3.1 Miscanthus

The time series of VOC fluxes above Miscanthus are shown in Figure 2 along with $u^*$ and sensible heat flux. Two periods of missing data 21st – 22nd and 25th – 27th July were due to
failure of the sampling pump. Data in the first few days were relatively noisy, showing no particular diurnal trend up to 20\textsuperscript{th} July. This was likely due to elevated $O_2^+$ impurities during transport of the instrument resulting in less reliable primary ion counts or higher LOD. Additionally, episodes of rainfall on 17\textsuperscript{th}, 18\textsuperscript{th}, 20\textsuperscript{th} and 22\textsuperscript{nd} July may have resulted in a reduction in mixing ratio of VOCs where emission is proportional to PAR.

Small net emissions of isoprene and MEK from \textit{Miscanthus} during daytime were just discernible, most noticeable on 18\textsuperscript{th} July when sensible heat flux was also at its maximum. However, in general, flux data were somewhat noisy for all VOCs measured, and mostly not significantly different from zero. The directionally-filtered diurnal averages of VOC fluxes are shown in Supplementary Information Figure S2. As described earlier, the relevant sector for the \textit{Miscanthus} measurements was south-west (180 – 270°). The mixing ratios of MVK+MACR (the first-generation oxidation products of isoporene) showed no diurnal pattern and were below LOD, so no data for these species are shown.

The time series of VOC mixing ratios above \textit{Miscanthus} are shown in Supplementary Information Figure S3. For the period 27\textsuperscript{th} July to 2\textsuperscript{nd} August, mixing ratios of all measured VOCs had maxima at night except for isoprene whose mixing ratios were elevated in late afternoon. The average diurnal VOC mixing ratios above \textit{Miscanthus} are shown in Figure S4. Methanol, acetaldehyde, acetone, acetic acid and MEK had similar diurnal profiles in mixing ratio. All showed a minimum mixing ratio around midday. The isoprene mixing ratio peaked around midday consistent with observation of a possible small isoprene flux above \textit{Miscanthus} (Figure S2). No isoprene or monoterpenes were detected in the GC-MS analysis of adsorption tube sampling above the \textit{Miscanthus} canopy.
3.2 Short rotation coppice willow

The time series of VOC fluxes, and $u^*$ and sensible heat, measured above willow are shown in Figure 3. Missing data on 10th and 12-13th August were due to failure of the mobile power supply. Data were directionally filtered to include only those from over the willow field (90 – 270°) before diurnally averaging (Figure 4). Willow showed a distinct diurnal pattern of isoprene flux, peaking at ~1 mg m$^{-2}$ h$^{-1}$ around midday and decreasing to zero overnight, driven by the strong dependence of isoprene emissions on temperature and PAR. All other VOC measured showed positive and negative fluxes throughout the day, with no significant net positive or negative daily flux overall.

Supplementary Information Figure S5 shows the time series of VOC mixing ratios and temperature above SRC willow. The time series showed clear diurnality in mixing ratios of all VOCs measured, except for methanol. The directionally-filtered diurnal averages of mixing ratio over the willow are shown in Figure 5.

Isoprene had a dominant maximum mixing ratio in early afternoon (~1 ppbv), temporally coincident with the temperature profile, and low mixing ratios at night. Figure 5 also plots the isoprene mixing ratios determined by adsorption tube sampling and GC-MS analysis. There was good agreement. Small quantities of the monoterpenes α-pinene and limonene were also detected by GC-MS, but no diurnal patterns were discernable in these data.

Acetic acid, acetaldehyde and MVK+MACR also showed diurnal profiles with maxima in the afternoon and minima around 6 am, closely mirroring daily temperature variation. The amplitude of the daytime maximum of MEK mixing ratio was considerably less. Acetone
exhibited low diurnal variability but with the small maximum in early morning similar to the observation of *Miscanthus*.

As isoprene oxidation is the only known source of MVK and MACR, the ratio of MVK+MACR to isoprene mixing ratios can be used to examine the degree of isoprene oxidation (Figure 6). The [MVK+MACR]:[isoprene] ratio peaked around midnight with an average value of about 0.8 when isoprene was not being emitted and MVK+MACR were not undergoing photochemical loss or dispersion. At dawn there was a rapid decline in the ratio as the canopy responded to increasing PAR and temperature hence isoprene emissions increased, and the boundary layer depth also increased. The minimum ratio of ~0.1 occurred for several hours around midday. The ratio rose in late afternoon as isoprene emissions declined but isoprene oxidation continued.

The average measured daytime [MVK+MACR]:[isoprene] ratio of 0.24 is comparable with those from other northern latitude studies. A daytime ratio of 0.23 was measured in a rural Canadian forest clearing (Biesenthal et al., 1998), 0.12 was reported for a mixed forest in Michigan, USA (Apel et al., 2002) and 0.4 – 0.8 in a deciduous forest in Pennsylvania, USA (Martin et al., 1991).

### 3.3 Standardised isoprene emission

As isoprene emission from plants is strongly influenced by light and leaf temperature, the canopy-level emission, $F$, was recalculated as a standard emission factor ($\varepsilon$) normalised to a standard leaf temperature of 303 K and PAR flux of 1000 µmol m$^{-2}$ s$^{-1}$, as described by the G95 algorithm (Guenther et al., 1995),
\[
\varepsilon = \frac{F}{D} \gamma
\]  
(1)

where \(D\) is foliar density (g dry weight m\(^{-2}\)) and \(\gamma\) is a non-dimensional activity adjustment factor to account for the effect of light and temperature:

\[
\gamma = C_L C_T.
\]  
(2)

The light dependence, \(C_L\), is defined by

\[
C_L = \frac{\alpha c_L \beta Q}{\sqrt{1 + \alpha^2 Q^2}}
\]  
(3)

where \(\alpha\) (0.0027) and \(c_L\) (1.066) are empirical coefficients and \(Q\) is PAR flux (\(\mu\)mol m\(^{-2}\) s\(^{-1}\)).

The temperature dependence \(C_T\), is defined by

\[
C_T = \exp \left( \frac{c_{T1} (T - T_2)}{RT_2 T} \right) \cdot \frac{1}{1 + \exp \left( \frac{c_{T2} (T - T_M)}{RT_2 T} \right)}
\]  
(4)

where \(T\) is leaf temperature (K), \(T_2\) is leaf temperature at standard conditions (303 K), \(R\) is the universal gas constant (8.314 J K\(^{-1}\) mol\(^{-1}\)), and \(c_{T1}\) (95000 J mol\(^{-1}\)), \(c_{T2}\) (230000 J mol\(^{-1}\)) and \(T_M\) (314 K) are empirical coefficients.

Values of above-canopy PAR and temperature, and of isoprene flux (from Figure 4), at hourly intervals during the willow campaign were used for the calculation of \(\gamma\) and \(F\) respectively. Foliar density \(D\) was estimated at 150 g\(_{dw}\) m\(^{-2}\) for Salix spp. (Karl et al., 2009). Hourly emission factors \(\varepsilon\) were then determined for isoprene from willow, and were found to have a peak value of 25 \(\mu\)g g\(_{dw}\)\(^{-1}\) h\(^{-1}\) at 10:00. A mean midday value of 20 \(\mu\)g g\(_{dw}\)\(^{-1}\) h\(^{-1}\) for 12:00 \pm 2 h was determined to allow comparison, in Table 2, with mean values from other studies.

Table 2 shows that the emission factor from this study was within the range of values derived previously for Salix spp. The slightly lower measurements derived in this work and in the
other above-canopy study (Olofsson et al., 2005) were canopy-averaged emissions factors which included leaves which were in shade as well as those in direct sunlight. It was therefore expected that these measurements would result in lower standard emission factors than from individual branch or leaf-level experiments.

4 Discussion

In the context of SRC willow as a bioenergy crop, the significant isoprene emission factor could potentially impact local and regional air quality by affecting tropospheric ozone production and SOA formation. Conventional agricultural crops are regarded as being low emitting species. For example, wheat and oats are estimated as having isoprene emission factors in the range 0 - 0.5 µg g$_{dw}^{-1}$ h$^{-1}$ (Karl et al., 2009; König et al., 1995), while those for rapeseed, rye and barley are zero (Karl et al., 2009; Kesselmeier and Staudt, 1999). Replacement of conventional crops with SRC willow would therefore result in increased isoprene emission. A recent study examined the impact of SRC crop cultivation in Europe (Ashworth et al., 2012). It was concluded that monthly mean increases in ozone and BSOA (+1% and +5% respectively) from low level planting scenarios were significant enough to affect regional air quality and therefore warrant consideration in short term local impact assessments, as well as life cycle analysis of bioenergy crops.

At the end of 2009, total UK plantings of Miscanthus and SRC willow were 12,700 and 6,400 ha respectively (DEFRA, 2009). A government report stated that there is potential in the UK to increase bioenergy crop cultivation substantially by a further 350,000 ha by 2020, accounting for ~6% of total UK arable land (DEFRA, 2007), with the assumption that 70%
would be *Miscanthus* and SRC willow (Rowe et al., 2009). In the case of 70% being converted solely to SRC willow, then a UK-wide increase of up to 7.35 t h\(^{-1}\) in emissions of isoprene would result (assuming zero isoprene emissions from the land prior to conversion to willow, 150 g\(_{dw}\) m\(^{-2}\) willow, and an isoprene standard emission rate of 20 µg g\(_{dw}\) h\(^{-1}\) determined here). The annual increase in isoprene would require calculation using PAR and temperature data across the likely planting sites in the UK, and a whole-canopy model.

The standard emission factor for isoprene from SRC willow measured in this study was 26.5 g C ha\(^{-1}\) h\(^{-1}\). This is an order of magnitude higher than was determined for total emission of VOCs from the biofuel crop switchgrass (Eller et al., 2011), where emissions were dominated by oxygenated VOCs and isoprene contributed less than 8%.

For *Miscanthus*, the near-zero values of flux at night were in contrast to the increase in mixing ratios of oxygenated VOCs (Figure S3 & S4). Since reliability of the eddy covariance technique depends on friction velocity, the greater boundary layer stability at night (hence low friction velocity) resulted in unreliable night time flux measurements. It may therefore be possible that night time fluxes were non-zero. A more likely scenario is that increasing VOC mixing ratios at night were affected by sources in the wider area. Towards the start of the measurement period, several of the surrounding fields were subject to harvesting and subsequent ploughing activities, which are known to be a source of oxygenated VOCs (Karl et al., 2001). Mixing ratios of methanol, acetaldehyde and acetone were comparable to those measured in previous field experiments of crop cutting (Warneke et al., 2002). The enhanced mixing ratios towards dusk, and at night can be explained by reduced radical sink chemistry, together with accumulation within a shallower nocturnal boundary layer from reduced vertical transport and mixing, as demonstrated by the lower wind speed and \(u^*\) at night (Figure 2).
During willow measurements, atypical increases in mixing ratios of methanol, acetone and acetic acid on 8th August may also have been caused by further harvesting activity in the wider area.

5 Conclusions

Measurements of above-canopy fluxes and mixing ratios of VOCs revealed significant emissions of isoprene from short rotation coppice willow, with a standard emission factor of 20 µg g_{dw}^{-1} h^{-1}. No significant emissions were measured from Miscanthus. This is the first field study of bioenergy crops in the UK and shows that a change in land use from conventional to bioenergy crops could result in increased isoprene emissions. Bioenergy crop species choice should therefore include consideration of their impact on regional air quality. Further work could include measurement of VOC emissions from Miscanthus and SRC willow during senescence and harvesting.

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Tables

Table 1: Compounds measured during the field campaign including dwell times, sensitivities and limits of detection.

<table>
<thead>
<tr>
<th>m/z [amu]</th>
<th>Contributing compound(s)</th>
<th>Formula</th>
<th>Dwell time [s]</th>
<th>Sensitivity [ncps ppbv⁻¹]</th>
<th>Limit of detection [ppbv]</th>
</tr>
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<tr>
<td></td>
<td></td>
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<td></td>
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<td>Miscanthus</td>
</tr>
<tr>
<td>21</td>
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<td>-</td>
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<td>33</td>
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<td>-</td>
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<tr>
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<td>acetaldehyde</td>
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<td>acetone propanal</td>
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<td>isoprene furan methyl butenol fragment</td>
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<td>3.5</td>
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<td>methyl vinyl ketone (MVK)</td>
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<td>methyl ethyl ketone (MEK)</td>
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Table 2: Comparison of standardised emission rates of isoprene from willow. REA = relaxed eddy accumulation

<table>
<thead>
<tr>
<th>Species</th>
<th>Standard emission rate / µg g&lt;sub&gt;dw&lt;/sub&gt; h&lt;sup&gt;-1&lt;/sup&gt;</th>
<th>Measurement type</th>
<th>Reference</th>
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<td>Salix spp.</td>
<td>20</td>
<td>Canopy-scale, PTR-MS</td>
<td>This study</td>
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<td>Salix spp.</td>
<td>28</td>
<td>Branch enclosure</td>
<td>(Owen and Hewitt, 2000)</td>
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<td>Salix alba</td>
<td>18</td>
<td>Branch enclosure, lab conditions</td>
<td>(Pio et al., 1993)</td>
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<td>Salix alba</td>
<td>37.2</td>
<td>-</td>
<td>(Karl et al., 2009)</td>
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<td>(Winer et al., 1983)</td>
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Figures

Figure 1: Aerial view of the *Miscanthus* and willow plantations. The white dots denote the measurement locations at the NE corner of the *Miscanthus* field and the N edge of the SRC willow field. (Map attributable to: ©2001 DigitalGlobe, GeoEye, Getmapping plc, Infoterra Ltd & Bluesky, TerraMetrics. Map data ©2011 Google).
Figure 2: Time series of VOC fluxes measured above Miscanthus. Dashed gridlines denote midnight. Note the variable flux scales.
Figure 3: Time series of VOC fluxes measured above willow canopy. Dashed gridlines denote midnight. Note the variable scales.
Figure 4: Average diurnal profiles of VOC fluxes above willow, and of sensible heat flux, when wind direction was between 90° and 270° (i.e. from over the willow field). Note the variable scales. Grey areas show variability calculated as ±1 sd of the averaged half-hourly values of all measurements.
Figure 5: Average diurnal profiles of VOC mixing ratios above willow, and of temperature, when wind direction was between 90 and 270° (i.e. from over the willow field). Note the variable scales. Dashed lines denote LOD. Grey areas represent variability calculated as ±1 sd of the averaged half-hourly values of all measurements.
Figure 6: Average diurnal profile of [MVK+MACR]:[isoprene] ratio above the willow canopy. Grey areas represent variability calculated as ±1 sd of the averaged half-hourly values of all measurements.
Supplementary Information

Volatile Organic Compound Emissions from Miscanthus and Short Rotation Coppice Willow Bioenergy Crops

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The main paper reports on field measurements of VOC concentrations and fluxes above Miscanthus and short rotation crop (SRC) willow crops.

Average diurnal profiles were derived from the time series of measurements using data filtered to examine only those times when the wind direction and flux footprint was over the relevant crop field. The flux footprint for the range of friction velocities encountered was estimated as described in the main text, and the model output is shown here in Supplementary Information Figure S1. For measurements over Miscanthus and willow, directional filtering was undertaken for the sectors 180 – 270° and 90 – 270°, respectively. (See also the layout of the field site shown in Figure 1.)

Time series and directionally-filtered diurnal averages of both fluxes and mixing ratios of VOCs above both Miscanthus and SRC willow comprise too many figures to present in the main text. The most salient are presented in the main paper, and are mainly concerned with the measurements above SRC willow since measurements above Miscanthus showed essentially no significant VOC fluxes. For completion, the directionally-filtered diurnal averages of fluxes above Miscanthus are presented here in Figure S2, and the time series and directionally-filtered diurnal averages of the VOC mixing ratios above Miscanthus are presented in Figures S3 and S4, respectively. The time series of VOC mixing ratios above willow are shown in Figure S5.
Figure S1: Modelled flux footprints for Miscanthus and SRC willow measurements. The following parameters were used. Miscanthus: measurement height $z_m$ 4 m; roughness length $z_0$ 0.25 m (estimated as 1/10th of the canopy height, 2.5 m); boundary layer height $h$ 1000 m. SRC willow: $z_m$ 6.7 m; $z_0$ 0.4m; $h$ 1000 m. Footprints were calculated for 90th percentile (P90), median and minimum values of $u^*$ (1 sd of the vertical wind speed, $\sigma_u$, shown in brackets) as indicated on the graphs. The distance at which maximum contribution can be expected, and at which 80% of the flux is contained, are given as $X_{max}$ and $X_r$, respectively.
Figure S2: Average diurnal profiles of VOC fluxes above Miscanthus when wind direction was between 180 and 270° (i.e. from over the Miscanthus field). Note the variable scales. Grey areas represent variability calculated as ±1 sd of the averaged half-hourly values of all measurements. A profile for MVK+MACR is not included due to insufficient data.
Figure S3: Time series of VOC mixing ratios, and of temperature, measured above Miscanthus. Dashed gridlines denote midnight. Note the variable mixing ratio scales.
Figure S4: Average diurnal profiles of VOC mixing ratios above *Miscanthus*, and of temperature, when wind direction was between 180 and 270° (i.e. from over the *Miscanthus* field). Note the variable scales. Dashed lines denote LOD. Grey areas represent variability calculated as ±1 sd of the averaged half-hourly values of all measurements.
Figure S5: Time series of VOC mixing ratios, and of temperature, measured above willow. Dashed gridlines denote midnight. Note the variable mixing ratio scales.