Seasonal Variations in Isoprene Emissions from a Boreal Aspen Forest

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Abstract

The primary objective of this study was to understand the environmental and seasonal controls over isoprene emissions from a boreal forest ecosystem whose isoprene source came from trees of the same species and age. A further objective was to establish an annual budget of isoprene emitted from a remote boreal forest and thus assess uncertainties associated with seasonal isoprene emission inventories. The onset of isoprene emissions occurred two weeks after the forest attained its maximum leaf area. Scaled to the foliage level, averaged isoprene fluxes approached $10^6$ nmol m$^{-2}$ s$^{-1}$ in the spring. During the middle of the growing season averaged isoprene emissions amounted to $28^64$ nmol m$^{-2}$ s$^{-1}$, whereas late summer values reached $16^62$ mmol m$^{-2}$ s$^{-1}$. These isoprene capacities were normalized to $25^8C$ and photosynthetically active radiation of 1000 mol m$^{-2}$ s$^{-1}$.

Given the strong seasonality observed in isoprene emissions, the authors propose to include seasonally adjusted emission rates to derive isoprene inventories for the entire foliage growing cycle. With an active biomass of 144 g m$^{-2}$, using a seasonally adjusted emission rate in a one-dimensional multilayered model it is estimated that during 1994 the boreal aspen forest emitted 32 mmol of isoprene per square meter. Such isoprene source strength represented approximately 1% of the photosynthetically fixed carbon by the aspen forest. In addition to the seasonal controls dictated by the inherent plant metabolic activity, low temperatures ($<10^8C$) strongly reduced the amplitude of diurnal isoprene emissions.

1. Introduction

Certain plant genera, especially many deciduous tree species, release large amounts of isoprene (2-methyl 1,3-butadiene $= \text{CH}_2=\text{C(\text{CH}_3)}=\text{CH}=\text{CH}_2$) to the atmosphere. At the global scale, the estimated phytogetic isoprene flux is approximately $5 \times 10^{14}$ g C yr$^{-1}$ (Guenther et al. 1995). Owing to its rapid gas-phase daytime reactions with ozone (O$_3$) and the hydroxyl radical (OH), isoprene can also impact both regional and global chemistry. By competing with greenhouse gases such as methane for OH, isoprene can influence the oxidation capacity of the troposphere. Reactions of isoprene with OH produce carbon monoxide and carbonyl compounds (i.e., vinyl ketone and methyl acrolein), which once in environments rich in nitrogen oxides under favorable meteorological conditions, can lead to the formation of ozone (Chameides et al. 1988; Fehsenfeld et al. 1992). Because isoprene is linked to the carbon fixation pathway in plants (Monson and Fall 1989; Sharkey et al. 1991), there is also scientific interest in discerning the fraction of photosynthetically fixed carbon entering the atmosphere in the form of isoprene. Therefore, understanding the magnitude and seasonal patterns of emissions is required to quantify and assess isoprene impacts on regional and global oxidant formation, and on atmospheric carbon cycling.

To investigate the role of isoprene in regional oxidant formation, biospheric modeling systems exist to derive isoprene inventories. These systems consider emission rates at the foliage level, active biomass, and microclimate characteristics within vegetated landscapes to integrate emissions to the canopy (Baldocchi et al. 1995; Fuentes et al. 1996; Lamb et al. 1996) and ecosystem (Geron et al. 1994; Guenther et al. 1994; Lamb et al. 1993) scales. At the canopy level, comparisons between modeled and measured isoprene fluxes show discrepan-
ancies ranging from 20% to 200%, with models overestimating emissions (Baldocchi et al. 1995; Fuentes et al. 1995; Geron et al. 1997; Guenther et al. 1996; Lamb et al. 1996). During the middle of the growing season, modeled isoprene emissions are more reliable and agree within 50% of measured quantities. Major uncertainties in inventories include the vegetation surveys incorporated in models and associated scaling of emission algorithms from leaves to trees of different species and ages. Another reason models may provide unrealistic scaling of isoprene emissions in mixed stands is due to the different “footprints” of isoprene emitting vegetation, the source strength varying with wind direction and atmospheric stability regime (Lamb et al. 1996). Moreover, in spite of isoprene emissions changing considerably over the growing season (Grinspoon et al. 1991; Fuentes et al. 1995; Monson et al. 1994), emission rates characterizing mid–growing season conditions are sometimes used with modeling systems to derive inventories for the entire growing season.

Canopy- and ecosystem-scale isoprene fluxes are needed to provide data for testing and improving inventory modeling systems. Existing forest canopy isoprene flux data come largely from short-term studies, representing conditions during the middle of the growing season (Baldocchi et al. 1999). Studies on seasonal isoprene emissions at the ecosystem level are rare. Therefore, in 1994 we conducted a field research project to determine isoprene fluxes from a deciduous forest in the southern boreal region of Canada. The primary objective of this study was to characterize the environmental and seasonal controls over isoprene emissions from a forest ecosystem whose isoprene source came from trees of the same species and age. A further objective was to establish an annual budget of isoprene emitted from a remote boreal forest and assess the uncertainties associated with seasonal isoprene emission inventories. The information presented here has important consequences relating to the isoprene content of the remote environment where it can alter the oxidation capacity of the atmosphere.

2. Site description and measurements

The data reported here were obtained during April to September 1994. This study was conducted as part of the Boreas Ecosystem Atmosphere Study (BOREAS) project, and therefore additional measurements exist on energy and trace gas fluxes for the aspen forest (Blanken et al. 1996; Black et al. 1996; Blanken et al. 1997; Sellers et al. 1995; Simpson et al. 1997).

a. Site description

Measurements were obtained at a forest stand of trembling aspen (Populus tremuloides), situated in the southern boreal forest of Canada (53.7°N, 106.2°W). Randomly situated throughout the landscape, the forest also included a small percentage (<8%) of balsam poplar (Populus balsamifera), white spruce (Picea glauca), and black spruce (Picea mariana). During the 1994 growing season the height of this 70-yr-old aspen forest averaged 22 m and had a final leaf area index (LAI) of 2.4 (Fig. 1). The canopy architecture featured an open trunk space between 2 and 15 m without foliage, and aspen leaves were distributed in the layer between 15 and 22 m. The stand density was approximately 900 stems ha⁻¹ and average tree diameter at the 1.5-m height was 17 cm. Between the ground and 2 m a hazelnut (Corylus cornuta) understory with an LAI of 3.2 (Black et al. 1996) existed throughout the landscape. To support measurements inside and above the canopy, a 37-m scaffold tower was established in 1993. From the tower site the fetch extended several kilometers in all directions. The site was flat and suited to make eddy covariance and gradient-type flux density measurements.

Final LAI of the aspen forest was determined by collecting the autumnal leaf-litter fall. Leaves were collected in baskets with upper diameter 0.5 m, lower diameter 0.4 m, and depth 0.3 m. Before leaves began to fall (20 August) 60 baskets were deployed along four transects, each extending 25 m, in the vicinity of the tower. The collected leaves were taken to the laboratory and soaked in water for several hours before measuring the planar area using an area meter (model LI-3000, LICor Inc., Lincoln, NE). Neumann et al. (1989) provided the details on this method of determining forest LAI. Black et al. (1996) determined seasonal LAI measurements using a plant canopy analyzer (model LAI-2000, LICor Inc.). The seasonal LAI for aspen foliage (Fig. 1) was used to estimate seasonal isoprene fluxes.

b. Microclimate and flux measurements

During April to September in 1994 both microclimate and flux measurements were accomplished by mounting instruments on the scaffold tower. Seasonal measure-
ments of wind speed and direction (R. M. Young ane-
ometer model 0571, Traverse City, MI), incoming so-
lar radiation (Model PSP Eppley pyranometer, Newport,
RI), net radiation (model S-1 net pyradiometer, Swis-
steco Instruments, Oberriet, Switzerland), photosyn-
thenetically active radiation (PAR, Model LI1900A, LICor
Inc.), and relative humidity (Model MP-100, Rotronic
Instrument Corp., Huntington, NY) were made above
the forest. Temperature of the forest crown (20–22 m
above the ground) was measured with an infrared ther-
mometer (model 2000A, Everest Interscience Inc., Ful-
lerton, CA). Air temperature measurements were made
as well at 12 levels above the ground using ventilated
and radiation shielded copper±constantan thermocou-
ples. Microclimate data were acquired using dataloggers
(model CS21XL, Campbell Scientific Inc., Logan, UT),
which provided half-hourly averaged quantities.

Eddy covariance fluxes for momentum ($\tau$), virtual
heat ($H_v$), carbon dioxide, and water vapor were deter-
mined throughout the growing season using three-di-
ensional sonic anemometers (model DAT-310, Kaijo
Denki Ltd., Tokyo, Japan) in combination with appro-
priate fast-response gas analyzers (model LI-6262, LI-
Cor Inc.). The data ($\tau$, $H_v$) were acquired at 100 Hz
and block averaged to 20 Hz. From the raw data, half-
hourly fluxes were calculated. For this study, such fluxes
were further reduced to hourly averages to obtain the
contextual eddy covariance data required to derive the
atmospheric eddy diffusivities used in the calculation
of isoprene fluxes. Black et al. (1996) provided more
details on the eddy covariance measurements.

To determine isoprene concentration gradients
($\partial X/\partial z$), air was drawn at fast flow rates (>20 L min$^{-1}$)
through cleaned Teflon tubing (0.5” ID) from two in-
takes placed at 27 and 39 m above the ground. Before
isoprene concentration gradient measurements com-
 menced, air of known isoprene concentration was passed
through the sampling lines to assess isoprene losses. The
detected isoprene losses were small, amounting to less
than 2%. Atmospheric samples from the two intakes
were collected in 3.2-L electropolished canisters. Iso-
prene concentrations were measured using gas chro-
matography (GC, model 5890, Hewlett Packward, Palo
Alto, CA) coupled with a flame ionization detector
(FID). Because atmospheric isoprene levels existed in
trace amounts, the analyte required a preconcentration
that was achieved by passing air samples through cryo-
genically cooled traps packed with glass beads. During
sample preconcentration, trap temperatures remained
constant at $-150^\circ$C and were controlled through com-
puter software. Samples of known flow rate and preset
volume passed the traps before concentrated analytes
were introduced to the chromatographic column (HP-1,
50 m $\times$ 0.31 mm ID, 1-$\mu$m film thickness) after thermal
desorption at 120$^\circ$C. Determined by replicate of sample
analysis, isoprene analytical precision was within 4% at
about 1.0 parts per billion on a volume basis (ppbv) and
7% at mixing ratios less than 0.2 ppbv. The detection
limit was 0.01 ppbv. Further details on GC–FID cali-
bration protocols are provided elsewhere (Fuentes et al.
1996). To characterize the most dominant anthropogenic
and biogenic volatile organic compounds from April to
September, ambient sampling was achieved within the
forest crown during 1200–1500 local time. Only am-
bient isoprene mixing ratios are presented here (Fig. 2)
to illustrate the seasonal patterns.

The isoprene fluxes were determined using the inte-
grated flux–gradient approach based on the Monin–
Obukhov similarity hypothesis (Monin and Obukhov
1954). In this method, it is assumed that the isoprene
flux can be characterized by the product of $\partial X/\partial z$ defined
between two levels (27 and 39 m above the ground)
above the isoprene source and an atmospheric eddy dif-
sivity ($K_z$), as shown in (1):

$$F_x = -K_z \frac{\partial X}{\partial z}.$$  (1)
The $K_v$ values were calculated based on hourly averaged, eddy-covariance measurements of momentum (which provided the friction velocity) and virtual heat fluxes (required in the diabatic correction functions for heat). Because the isoprene “footprint” source area had trees of the same species and age, the upper and lower intakes sampled air representing equal source strength and thus minimum uncertainties are expected in isoprene concentration gradients. Simpson et al. (1997) provided a flux source footprint analysis for the aspen forest. Due to uncertainties associated with determination of eddy diffusivities and measurements of isoprene concentration gradients, following Sinclair et al. (1975), the estimated uncertainty in isoprene fluxes is less than 30%. Further details on theoretical background and uncertainties associated with gradient diffusion fluxes are provided by Fuentes et al. (1996) and Simpson et al. (1997).

Following the measurement protocols outlined before (Fuentes and Gillespie 1992; Fuentes et al. 1996), leaf isoprene emissions were determined during 10 days in June to September to find what local plant species emitted isoprene and to establish diurnal cycles of isoprene emission rates. The plant species present within the flux footprint of the measurement tower (within a radius of $\sim$800 m) were trembling aspen, balsam poplar, and hazelnut. Hazelnut plants did not release isoprene, whereas trembling aspen and poplar trees exhibited similar isoprene emission rates defined at the leaf temperature of 25°C and photosynthetically active radiation (PAR) of 1000 $\mu$mol m$^{-2}$ s$^{-1}$.

3. Presentation and discussion of isoprene measurements

a. Seasonal ambient isoprene levels

Figure 2 includes isoprene mixing ratios measured within the forest crown during 1200–1500 LT. Because temperature influences isoprene emissions through the kinetics of the enzyme responsible for isoprene biosynthesis (Monson et al. 1992), averaged canopy temperature data are also shown (Fig. 2). Since isoprene is linked to carbon fixation (Monson and Fall 1989), we report averaged CO$_2$ fluxes to indicate the level of forest–atmosphere CO$_2$ exchange (the CO$_2$ fluxes were also reported by Black et al. (1996)). Isoprene mixing ratios varied considerably with growing season. Before leaf bud break (<day 120), isoprene mixing ratios stayed about 0.01 ppbv, which according to Young et al. (1997) represents the prevailing isoprene levels observed in the southern boreal forests of Canada during the wintertime. Leaves expanded rapidly and attained full expansion by day 150, with maximum LAI of 2.4. Leaf area remained constant until day 200 and thereafter leaves started to senesce and abscise (Fig. 1). In response to increasing emissions as foliage became fully developed (≈day 150) isoprene mixing ratios progressively increased with growing season. Maximum isoprene mixing ratios were measured during the middle of the growing season (days 200–210), with peak levels reaching nearly 16 ppbv. These maximum isoprene levels coincided with peak canopy temperature of 25°C. Isoprene emissions (Figs. 4, 5, and 6) and the resultant ambient levels were strongly modulated by canopy temperature. Thus, the low isoprene levels (<1 ppbv) measured during days 200 and 210 occurred in response to low emissions that were associated with low temperatures (Fig. 2). Ambient isoprene mixing ratio measurements stopped on day 268 when senescing leaves on trees were still releasing isoprene. As a result, toward the end of the growing season isoprene levels remained sufficiently above 0.1 ppbv. The aspen forest became a carbon sink after it became fully leafed (Figs. 1 and 2). Maximal rates of noontime CO$_2$ uptake exceeded $\sim$25 $\mu$mol m$^{-2}$ s$^{-1}$ during days 180–210. Details on the relationship between CO$_2$ and isoprene fluxes on a diurnal basis are provided below in section 3c.

b. Leaf isoprene emission rates

To illustrate the magnitude of isoprene emissions at the foliage level for the aspen trees, the leaf-based fluxes when PAR exceeded 200 $\mu$mol m$^{-2}$ s$^{-1}$ were grouped and correlated against temperature (Fig. 3). These data include the period during 16 July to 12 September 1994. The observed variability in isoprene fluxes likely resulted due to natural changes in emission rates among leaves of different biological vitality. As expected, isoprene emissions increased exponentially with foliage temperature as temperature is related to the energy required in the activation of isoprene biosynthesis (Kuzma and Fall 1993). With the exception of one data point, isoprene fluxes were less than 30 nmol (C$_5$H$_8$) m$^{-2}$ (leaf area) s$^{-1}$. These isoprene emission rates are within the range of reported values (<40 nmol m$^{-2}$ s$^{-1}$) for aspen leaves (Fuentes et al. 1995; Monson et al. 1994; Sharkey et al. 1991). When isoprene fluxes are scaled according
to the Guenther et al. (1993) emission algorithm at leaf temperature of 25°C and PAR of 1000 μmol m⁻² s⁻¹, the averaged emission rate was 26.6 ± 10.2 nmol m⁻² s⁻¹ (Fig. 3). Given that cuvette measurements precluded the leaf development and senescence periods, this emission rate may be taken to represent midgrowing conditions for the aspen forest. Below we discuss how the canopy isoprene fluxes changed with growing season.

c. Canopy isoprene fluxes

Canopy isoprene flux measurements commenced in spring (26 May) and continued until the beginning of the autumnal period (20 September) when aspen leaves started to abscise (Fig. 1). During spring, while leaves were growing, canopy isoprene fluxes were below levels of detection until two weeks after full leaf expansion.

Because canopy isoprene fluxes were not continuously determined throughout the growing season, we segregated the data to obtain averaged fluxes for spring, summer, and late summer periods. Averaged quantities for canopy temperature, and CO₂ and isoprene fluxes, during 6–19 June are presented in Fig. 4a. Isoprene fluxes exhibited strong diurnal patterns in which emissions progressively increased after sunrise, reaching peak values during the 1500–1800 LT period, and rapidly declined after sunset (Fig. 4a). During spring, averaged maximum canopy isoprene emissions reached approximately 6 nmol (C₅H₈) m⁻² (ground area) s⁻¹, with peak values of 10 nmol m⁻² s⁻¹ being measured during the middle of June. Maximum isoprene fluxes occurred late in the afternoon and coincided with peak canopy temperature of 20°C (Fig. 4a). Despite the link between isoprene biosynthesis and foliage CO₂ uptake (Monson
isoprene fluxes reaching averaged peak values of approximately 40 nmol m$^{-2}$ s$^{-1}$ for oak and aspen forests situated in southern latitudes (Baldocchi et al. 1995; Fuentes et al. 1996; Geron et al. 1997; Guenther et al. 1996). However, during the middle of the growing season, the averaged peak isoprene fluxes of 28 nmol m$^{-2}$ s$^{-1}$ are lower than reported values (40–60 nmol m$^{-2}$ s$^{-1}$) for (aspen and oak) forests with comparable isoprene active biomass density distribution (for our boreal forest the active isoprene biomass density was 144 g m$^{-2}$).

During the middle of the growing season, ambient isoprene mixing ratios exhibited an unusual pattern with one peak occurring around 0900–1000 LT and a second one taking place at 1800–2000 LT (Fig. 4b). During these times the isoprene mixing ratios averaged 5 ppbv, with extreme high values approaching 16 ppbv. As in the spring, ambient isoprene levels rapidly decreased as evening approached likely in response to isoprene deposition and chemical processing. The lower isoprene mixing ratios measured during 1100–1800 LT can in part be explained by the temporal patterns of atmospheric turbulence and eddy diffusivities. In contrast to the springtime period (Fig. 4b), the eddy diffusivities in the summer period (Fig. 5b) exhibited a more pronounced diurnal variation. For example, the diffusivities remained low (<0.3 m s$^{-1}$; this form of eddy diffusivities already includes the effect of height) until 1000 LT and thereafter increased rapidly reaching peak values (~0.8 m s$^{-1}$) during 1200–1600 LT (data not shown). Other processes likely contributing to lower isoprene mixing ratios during the midday included photochemical reactions (Gao et al. 1993), particularly the ones with HO and O$_3$, which can be enhanced by warm air masses.

With the commencement of the autumnal foliage senescence, canopy fluxes progressively declined and averaged diurnal cycles exhibited similar characteristics to those measured in spring. Averaged isoprene fluxes reached nearly 5 nmol m$^{-2}$ s$^{-1}$ (Fig. 6a) with peak values approaching 8 nmol m$^{-2}$ s$^{-1}$. During 26 August–15 September, the maximum isoprene fluxes were measured in the period (1500–1800 LT) when the forest canopy was the warmest (~20°C), whereas maximum canopy CO$_2$ fluxes took place during the 1200–1500 LT period. In the fall, these uncoupled diurnal patterns of isoprene and CO$_2$ fluxes were observed at alpine aspen forests (Monson et al. 1994). Monson et al. (1994) ascribed the declining patterns in isoprene and CO$_2$ fluxes to the autumnal translocation of nitrogen out of leaves and the breakdown of foliage metabolic activity. However, at the canopy scale isoprene emissions decreased at a faster rate compared to the CO$_2$ flux trends. The data presented
in Fig. 6b show that ambient isoprene mixing ratios exhibited diurnal trends dissimilar to what was observed during the middle of the growing season. The double-peak cycle in isoprene diurnal variation was less discernible and peak levels occurred during the latter part of the afternoon when maximum averaged mixing ratios attained 1.5 ppbv (Fig. 6b).

Under comparable active biomass distribution, summertime averaged isoprene fluxes measured at the boreal forest are lower (e.g., 25 vs 50 nmol m\(^{-2}\) s\(^{-1}\)) than those from temperate and southern forest ecosystems. In part, the discrepancies result because southern forests experience a longer growing season under warmer climatic regimes. Southern forests can also undergo periodic droughts (Baldocchi 1997; Geron et al. 1997), which can induce foliage warming and thus increase isoprene emissions [at the research site, in 1994 conditions were 15% wetter than the 30-yr average; Black et al. (1996)]. In the boreal forest, other features of the climatic forcing variables on isoprene dynamics included frequent occurrences of low temperatures (<10°C), which may suppress isoprene emissions.

In the case of the aspen forest, low temperatures (<10°C) lasting as few as 3 h were sufficient to reduce the amplitude of daily isoprene emissions. Figure 7a includes an example of isoprene emissions before and after nighttime temperatures dropped below 10°C. Despite higher temperature and PAR levels, after the low temperature episode isoprene fluxes were lower compared to the previous day (Fig. 7a). In all cases examined, we observed a consistent suppression in emissions after the forest canopy experienced temperatures below 10°C. In total, four low temperature episodes were identified while measuring canopy isoprene fluxes. Figure 7b provides two additional cases of reduced isoprene fluxes after the exposure of low temperatures. The
history of low temperature exposure for both days was similar. In both cases the three preceding nights had temperatures around 5°C, whereas daytime maximum temperature reached 16°C. For Fig. 7b, the observed discrepancies in the diurnal isoprene fluxes likely resulted due to differences in PAR levels. Measured isoprene fluxes became large (>15 nmol m⁻² s⁻¹) 2–3 days after the low temperature occurrences (data not shown).

After exposure to low temperatures a general feature of diurnal isoprene emission patterns was that maximum fluxes did not necessarily coincide when temperature was highest (Fig. 7). At the leaf level, others studies give evidence of reduced isoprene emissions in response to low temperatures. For example, Sharkey and Loretto (1993) measured low isoprene emissions (~0.5 nmol m⁻² s⁻¹) when kudzu (Pueraria lobata) leaves were grown at 19°C, but emissions increased (~15 nmol m⁻² s⁻¹) when the growth temperature was raised to 24°C.

The documented cases of reduced fluxes after the occurrences of low temperature were too few to examine whether changes occurred in the basal isoprene emission rates because of low temperature exposure. To further investigate the influence of low temperature on isoprene emissions, the noontime ambient isoprene mixing ratios during and after the low temperature episodes were segregated. Furthermore, to account for influences of atmospheric mixing on ambient isoprene levels, isoprene mixing ratios were multiplied by the friction velocity (u*). The resulting quantities (in units of ppbv m s⁻¹) serve as surrogate isoprene fluxes. To include the effects of both temperature and PAR, the calculated quantities were further analyzed within the framework of the iso-
Isoprene emission algorithm (Guenther et al. 1993) detailed in section 4 [Eqs. (2)−(4)]. Regression analysis (Fig. 8) was done between temperature and light modulating functions \( f(T_L) f(\text{PAR}) \) and surrogate fluxes. Although the correlation was low \( R^2 = 0.24 \), subsequent to the low temperature episodes the slope of the regression line declined by a factor of nearly 3 compared to the case with no low temperature influence. The results (Figs. 7 and 8) indicate that prolonged exposure to low temperature may modify the basal emission rate regimes.

To determine the amount of aspen forest carbon assimilation that was emitted as isoprene, during 1000–1600 LT (the time period when \( \text{CO}_2 \) fluxes were most reliable) we calculated the percentage of carbon fixed in gross canopy photosynthesis and reemitted to the atmosphere as isoprene. In the calculations (Fig. 9) we considered that the hazelnut understory accounted for 32% (Black et al. 1996) of the gross canopy photosynthesis measured at 39 m above ground. The percentage of carbon assimilation lost in isoprene emission increased with temperature (Fig. 9). At the foliage level, Monson and Fall (1989) and Harley et al. (1996) reported similar relationships for aspen and sweetgum \( (\text{Liquidambar styraciiflua}) \) leaves, respectively. For the aspen forest, the amount of carbon lost in isoprene emission was between 0.1% and 3.2% of that assimilated through canopy photosynthesis. When all data (Fig. 9) were combined, the carbon introduced to the atmosphere in the form of isoprene amounted to 0.7 ± 0.6% but for the temperature range 20°–25°C the value increased to 1.3 ± 0.4%. These values are within the range (<8%) of previously reported data for aspen leaves (Fuentes et al. 1996; Monson and Fall 1989; Sharkey et al. 1991). Under the environmental conditions experienced during 1994, canopy isoprene fluxes did not represent substantial carbon losses from the aspen boreal forest.
4. Modeling isoprene emissions

A central goal of this study is to assess uncertainties associated with seasonal isoprene emissions. With the data presented above, we determined isoprene emissions throughout the growing season by applying a one-dimensional, multilayered canopy model. We employed the algorithm developed by Guenther et al. (1993) to integrate leaf-based isoprene emission rates ($E_L$) to the canopy scale. At the leaf level, isoprene emission rates are obtained using a standard emission factor ($E_S$) and modulating $E_S$ using leaf temperature [$f(T_L)$] and PAR [$f(PAR)$] functions as shown in Eq. (2):

$$E_L = E_S f(PAR) f(T_L),$$

where $E_S$ is defined at a specified leaf temperature ($T_L$) and intercepted PAR. The modulating functions, $f(PAR)$ and $f(T_L)$, are nondimensional and adjust the emission rate according to variations of leaf temperature and intercepted PAR. Guenther et al. (1993) developed Eq. (3) to describe the light influence on emissions and assumed a value of 1.0 at $PAR = 1000 \, \mu\text{mol} \, \text{m}^{-2} \, \text{s}^{-1}$:

$$f(PAR) = \frac{\alpha C_L PAR}{\sqrt{1 + (\alpha PAR)^2}},$$

where $R$ is the universal gas constant ($= 8.314 \, \text{J} \, \text{K}^{-1} \, \text{mol}^{-1}$), $T_L$ is the leaf temperature (K), and $T_{opt}$ is the optimum temperature (K). The $C_T$ and $C_L$ coefficients represent the activation and deactivation energies (J mol$^{-1}$), respectively.

To derive isoprene emissions at the forest canopy level, PAR and leaf temperature must be known with canopy depth. For this study, the leaf temperature profile was not measured. Therefore, to apply $f(T_L)$ inside the canopy, $T_L$ in each layer was determined from ambient temperature profile measurements. As confirmed from infrared thermometer and profile measurements, the trembling aspen foliage and air had similar temperatures under daytime conditions. The PAR levels inside the forest canopy were obtained using a bi-Lambertian radiative transfer model, which is described below.

a. Estimating photosynthetically active radiation

To determine PAR distribution within the canopy, we used a two-stream approximation model originally developed by Dickinson (1983) and recently revised by Gu (1998). The two-stream model is used here because it provides robust analytical solutions for diffuse and direct beam radiation separately. The model treats the forest as an active medium in partitioning the incoming radiation stream into intercepted, reflected, transmitted, and absorbed components. Such processes are incorporated in the radiation propagation Eq. (5) (Dickinson 1983):

$$f(T_L) = \frac{\exp\left(\frac{C_{T1}(T_L - T_S)}{RT_L T_S}\right)}{1 + \exp\left(\frac{C_{T2}(T_L - T_{opt})}{RT_L T_S}\right)},$$

where $R$ is the universal gas constant ($= 8.314 \, \text{J} \, \text{K}^{-1} \, \text{mol}^{-1}$), $T_L$ is the leaf temperature (K), and $T_{opt}$ is the optimum temperature (K). The $C_{T1}$ and $C_{T2}$ coefficients represent the activation and deactivation energies (J mol$^{-1}$), respectively.

Fig. 9. The percentage of the aspen forest photosynthetically fixed carbon introduced to the atmosphere in the form of isoprene as a function of temperature.

at temperature $T_L$ (303 K) and is given by Eq. (4) (Guenther et al. 1993):

$$f(T_L) = \frac{\exp\left(\frac{C_{T1}(T_L - T_S)}{RT_L T_S}\right)}{1 + \exp\left(\frac{C_{T2}(T_L - T_{opt})}{RT_L T_S}\right)},$$
where \( F^\uparrow \) and \( F^\downarrow \) represent the upward and downward diffuse radiative fluxes, respectively, normalized to the incident PAR measured above the forest; \( K \) is the light extinction coefficient; \( \omega \) denotes the PAR leaf scattering coefficient and equals the sum of the leaf reflection \( (\alpha) \) and transmission \( (\Psi) \) coefficients; \( L \) is the downward cumulative leaf area index; \( \mu \) is the average inverse diffuse optical depth per unit leaf area; and \( \beta \) and \( \beta_0 \) are the upscatter parameters for the diffuse and direct beam, respectively. The diffuse upscatter parameter \( \beta \) can be obtained from the analysis of Norman and Jarvis (1975), and is defined as

\[
\beta = \frac{[\alpha + \Psi + (\alpha - \Psi)\mu_\perp]}{2\omega},
\]

where \( \mu_\perp \) is the cosine of the leaf inclination angle and the overbar denotes the ensemble average for the canopy. The direct beam upscatter parameter \( \beta_0 \) is derived in Dickinson (1983):

\[
\beta_0 = \frac{1 + \mu K}{\omega K} a_\perp(\mu),
\]

where \( a_\perp(\mu) \) is the single scattering albedo for a semi-infinite canopy and is determined from

\[
a_\perp(\mu) = \omega \int_0^1 \frac{\mu \Gamma(\mu, \mu')}{\mu G(\mu') + \mu' G(\mu)} d\mu',
\]

where \( \mu \) is the cosine of solar zenith angle; \( G(\mu) \) represents the relative projected area of leaf elements in the direction of \( \mu \) and equals 0.5 for spherically distributed leaf canopies (Ross 1981). The azimuthally averaged area scattering function, \( \Gamma(\mu, \mu') \), defines the fraction of the radiant energy in the direction of \( \mu \) that is scattered by a unit area of leaf elements into the direction of \( \mu' \). According to Myneni et al. (1989), \( \Gamma(\mu, \mu') \) can be expressed as

\[
\Gamma(\mu, \mu') = \int_0^\mu g_\perp(\mu_\perp)[\alpha \Pi^+(\mu, \mu', \mu_\perp)
+ \Psi \Pi^-(\mu, \mu', \mu_\perp)] d\mu_\perp.
\]

In Eq. (9), \( g_\perp(\mu_\perp) \) is the leaf orientation distribution function and equals 1 for spherically distributed leaf canopies (Ross 1981), and

\[
\Pi^+(\mu, \mu', \mu_\perp) = \frac{1}{4\pi^2} \int_0^{2\pi} \int_0^{2\pi} (\Omega_2 \cdot \Omega')(\Omega_1 \cdot \Omega_2 \cdot \Omega') d\varphi_2,
\]

where \( \Omega_1, \Omega_2, \) and \( \Omega' \) represent the solar beam direction (\( \mu, \varphi \)), leaf normal orientation (\( \mu_\perp, \varphi_\perp \)), and scattered direction (\( \mu', \varphi' \)), respectively; \( (\Omega_1, \Omega) \) and \( (\Omega \cdot \Omega') \) represent the cosine of the angle between \( \Omega_1 \) and \( \Omega \), \( \Omega \), and \( \Omega' \), respectively. The average inverse diffuse optical depth per unit leaf area, \( \bar{\mu} \), was obtained from (Sellers 1985)

\[
\bar{\mu} = \int_0^1 \frac{\mu}{G(\mu)} d\mu.
\]

The light extinction coefficient \( K \) is defined as the ratio of \( G(\mu) \) to \( \mu \). Finally, the analytical solution to Eq. (5) is

\[
\begin{align*}
F^\downarrow &= \alpha_1 e^{-\mu} + \alpha_2 e^{-\alpha \mu} + \alpha_3 e^{\alpha \mu}, \\
F^\uparrow &= \alpha_4 e^{-\mu} + \alpha_5 e^{-\alpha \mu} + \alpha_6 e^{\alpha \mu},
\end{align*}
\]

where \( \alpha_1, \alpha_2, \ldots, \alpha_6 \), are coefficients derived from algebraic combinations of parameters in Eq. (5). Expressions for \( \alpha_1, \alpha_2, \ldots, \alpha_6 \), can be found in Sellers (1985). For the incident diffuse radiation, the direct radiative term on the right-hand side of Eq. (5) should be dropped and the solution can be found in Sellers (1985).

The two-stream approach has been extensively applied to study radiative transfer in plant canopies (Dickinson 1983; Sellers 1985; Kimes et al. 1987; Dickinson et al. 1990; Joseph et al. 1996). However, Gu (1998) identified two shortcomings in the original two-stream model and made relevant revisions. The original two-stream model cannot deal with foliage clumpiness. Most plant canopies have leaves grouped into crowns, branches, and shoots, and effectively alter the radiation regime (Chen et al. 1997). The clumping of foliage reduces the optical depth and thus results in more radiation transmission. Therefore, for a clumped canopy, it is the effective LAI, not the actual LAI, that determines the radiation transmission. The effective LAI is given as the product of the actual LAI and the clumping factor, a characteristic parameter for a given canopy (Chen and Cihlar 1995; Chen et al. 1997). To account for the effects of clumpiness, the effective LAI is used in this study to replace the actual LAI in the two-stream radiation equations. In this study, the seasonal clumping factor (whose value changed from 0.70 to 0.85) was derived from the relationship developed by Chen et al. (1997) for the aspen forest.

Another weakness of the original two-stream model is that it assumes isotropic scattering by leaf elements while typical values of leaf reflection coefficients are nearly twice as much as those of leaf transmission coefficients in both the visible and near-infrared regions. This unrealistic assumption can cause significant errors in the estimation of canopy reflectance and transmission. Better model performance was found when the bi-Lambertian leaf scattering model (Ross and Nilson 1968; Myneni et al. 1989) replaces the isotropic leaf scattering model in the two-stream model (Gu 1998). We use the bi-Lambertian leaf scattering model in this study.

For the boreal aspen forest we assumed that the canopy has a spherical leaf orientation distribution (Chen...
et al. 1997). We also assumed that the vertical LAI density exhibits a triangular LAI distribution with maximum foliage at the 18-m height and no foliage below 15 m and above 22 m. Throughout the modeling period, for PAR the leaf reflection coefficient \((a)\) and leaf transmission coefficient \((\Psi)\) were considered constant with the values of 0.09 and 0.06, respectively (Oke 1987, p. 117). We tested the modified two-stream radiative transfer model with spatially averaged radiation data obtained below the forest canopy. Radiometers mounted above the forest gave incoming radiation levels. The dataset used in the mode test included primarily middle of the growing season conditions, canvassing a wide range of radiation regimes that included cloudless and overcast conditions. Model outputs gave close agreement with measurements, with an overall correlation coefficient \((R)\) of 0.90 (Gu 1998).

b. Determining isoprene emission rates

In order to model the seasonal pattern of isoprene emissions at this site, we need to know how the emission factors changed over the season. Because isoprene records are more complete at the canopy scale we determined \(E_s\) based on measured canopy fluxes, modeled PAR, and estimated leaf temperature from measurements of air temperature profiles. Thus, given the isoprene active biomass in the forest (represented by the LAI), canopy-based emissions \((E_c)\) can be expressed as

\[
E_c = \int_0^{LAi} E_L dL = \int_0^{LAi} E_s f(PAR) f(T_c) dL, \tag{13}
\]

where \(L\) denotes the downward accumulated LAI, starting from the canopy top. The \(f(PAR)\) and \(f(T_c)\) functions are defined in Eqs. (3) and (4). If we further assume that \(E_s\) remains constant with canopy depth, then \(E_s\) can be obtained through

\[
E_s = \frac{\int_0^{LAi} E_c f(PAR) f(T_c) dL}{\int_0^{LAi} f(PAR) f(T_c) dL}. \tag{14}
\]

Other studies (Harley et al. 1996; Geron et al. 1994) indicate that \(E_s\) varies with canopy depth, particularly in forests where substantial amounts of foliage remain under shade. In this study we did not systematically examine how \(E_s\) varies with canopy depth. On the basis of similar specific leaf area (SLA) measurements made at the crown (20 m) and bottom (16 m above ground) of the canopy (SLA = 0.0192 ± 0.0004 m² g⁻¹) by the end of June 1994, we assume here invariant \(E_s\) with canopy depth [this forest allowed ~30% of the incoming light reach the top of the hazelnut understory; Chen et al. (1997)]. To characterize the required PAR and \(T_c\) inside the forest, the canopy was divided into 48 layers, each one having a \(dL\) value of 0.05 m² m⁻².

To assess whether the method outlined above provides reliable outputs, the \(E_s\) values derived from daily canopy fluxes were compared with emission rates obtained from cuvette measurements (section 3b), which were also normalized at leaf temperature of 25°C and PAR = 1000 μmol m⁻² s⁻¹ [i.e., Eqs. (3) and (4)]. The two independent methods of deriving \(E_s\) yielded acceptable results, with \(R^2 = 0.60\) (Fig. 10). Cuvette-based emissions exhibited greater variation but on average were only 8% greater than canopy-based \(E_s\) estimates. Accepting this level of uncertainty, we now proceed to determine seasonal isoprene emission rates from the more complete record of canopy fluxes.

c. Seasonal isoprene emissions

Using the method described above (section 4b), the \(E_s\) values over the growing season were determined from available canopy fluxes and the results were expressed as a function of day of the year. At this boreal forest, isoprene emissions commenced two weeks after the forest became fully leafed (Figs. 1 and 11). Delays in emissions during spring are consistent with observations at alpine (Monson et al. 1994) and temperate (Fuentes et al. 1995) aspen forests, and are ascribed to reduced enzymatic activity as developing foliage translocates substrates to carry out other physiological functions (Grinspoon et al. 1991; Kuzma and Fall 1993). Maximum emissions occurred during days 200–210 (Fig. 11), with maximal rates of emissions approaching 35 nmol m⁻² (leaf area) s⁻¹. These emission rates agree with reported emissions (~30 nmol m⁻² s⁻¹) for aspen foliage (Monson et al. 1994). After the onset of autumnal leaf senescence, isoprene emissions steadily declined in response to lower magnitude of climatic forcing variables and diminished isoprene-emitting substrate. Measurements were suspended before complete leaf fall. As a result, substantial isoprene fluxes (Fig.
11) were determined by the middle of September when nearly half of the foliage still remained on trees (Fig. 1). Because we did not continuously measure canopy isoprene fluxes, the interactions between low temperature and emissions are not incorporated in the normalized isoprene emission rates (Fig. 11).

d. Modeled canopy isoprene emissions

Before the canopy model can be applied to derive seasonal fluxes, it is necessary to understand whether it provides realistic isoprene scaling to the canopy dimension. Thus, using leaf-based emission rates (Fig. 10), the model was applied to determine isoprene emissions for the days when both canopy and leaf fluxes were available. The results (Fig. 12) showed that calculated isoprene flux densities compared closely with measurements ($R^2 = 0.89$). The model yielded credible results that on average were 14% greater than measured isoprene fluxes. As reported in previous studies (Baldocchi et al. 1999; Geron et al. 1997; Guenther et al. 1996; Lamb et al. 1996), we attribute this close agreement between modeled and measured fluxes to the proper description of both isoprene active biomass and microclimate characteristics inside the forest canopy. Based on these results (Fig. 12), we conclude that under the growing conditions experienced at our aspen boreal forest (but precluding the influence of low temperatures) the integration of isoprene emissions from leaf to canopy dimensions can be realistically achieved.

Given that the canopy model calculated realistic fluxes on a daily basis, we then investigated uncertainties associated with seasonal isoprene emissions. We applied the model to examine three emission scenarios. One scenario involved the use of a seasonally adjusted emission factor [$E_s(t)$] as established from the normalized fluxes reported in Fig. 11. This case was contrasted with model outputs using a constant emission factor. For the latter, we incorporated in the model the seasonal averaged isoprene flux [$E_s = 15$ nmol m$^{-2}$ (leaf area) s$^{-1}$]. Compared to seasonally adjusted values, the use of a constant emission rate gave higher isoprene fluxes during early and latter parts of the growing season, whereas close agreement resulted in the middle of the season (data not shown). Using a constant emission rate, the seasonally integrated isoprene emissions were underestimated by 12% (Fig. 13). Since the discrepancy is within measurement uncertainties, this level of agreement is adequate for most applications of isoprene emission inventories. However, the difficulty of choosing an appropriate averaged emission factor remains unresolved. To illustrate how unreliable isoprene inventories become, we considered a third modeling scenario using an emission factor that represented averaged isoprene emissions ($E_s = 28$ nmol m$^{-2}$ s$^{-1}$) measured during the middle of the growing season. This emission factor is similar to the rate (=26 nmol m$^{-2}$ s$^{-1}$) recommended for plants of the genus Populus (Guenther et al. 1994).

![Figure 11](https://example.com/image11.png)

**Fig. 11.** Seasonal pattern of isoprene emission rates (○) during the 1994 growing season. Emission rates were normalized to 25°C and photosynthetically active radiation of 1000 μmol m$^{-2}$ s$^{-1}$. The bars denote ±1 std dev from the mean (●) values, and the solid line represents the best fit to data points.

![Figure 12](https://example.com/image12.png)

**Fig. 12.** Comparison between measured and calculated (○) flux densities of isoprene (in units of nmol m$^{-2}$ s$^{-1}$) for the aspen forest during selected days (16 Jul–12 Sep) of the 1994 growing season.

![Figure 13](https://example.com/image13.png)

**Fig. 13.** Cumulative isoprene fluxes during the 1994 growing season using seasonally adjusted (-----) and constant emission rates. For the latter, the data denoted by the solid line (——–) used an emission factor of 15 nmol m$^{-2}$ s$^{-1}$, whereas the thin line (——) incorporated an emission rate of 28 nmol m$^{-2}$ s$^{-1}$. Cumulative daily fluxes are in the units of mmol m$^{-2}$.
Applying the emission rate of 28 nmol m\(^{-2}\) s\(^{-1}\), we calculated a 65% overestimate of seasonally integrated isoprene emissions (Fig. 13). With this emission rate the model estimated realistic isoprene fluxes during the middle of the growing season but failed to describe the seasonal emission patterns established with the seasonally adjusted emission rates. Although in the opposite direction, larger deviations would be expected if we used the emission rates (~5 nmol m\(^{-2}\) s\(^{-1}\)) incorporated in the modeling systems to derive inventories from forests in southern Canada (Fuentes et al. 1995). Based on the integrated, observed seasonal canopy isoprene emissions we calculated that the boreal aspen forest with an active biomass of 144 g m\(^{-2}\) during 1994 produced 34 mmol of isoprene per square meter.

5. Summary and conclusions

The information reported here represents a unique dataset of seasonal isoprene fluxes from a forest ecosystem whose isoprene source came from trees of the same species and age. Such seasonable isoprene fluxes can be used to test and evaluate inventory modeling systems. At this boreal forest, the onset of isoprene emissions occurred two weeks after the forest attained its maximum leaf area. Representing emissions during spring and scaled to the leaf level, averaged isoprene fluxes approached 10 ± 5 nmol m\(^{-2}\) s\(^{-1}\). During the middle of the growing season averaged isoprene emissions amounted to 28 ± 4 nmol m\(^{-2}\) s\(^{-1}\), whereas late summer values reached 16 ± 2 nmol m\(^{-2}\) s\(^{-1}\). These isoprene capacities were normalized to 25°C and PAR of 1000 μmol m\(^{-2}\) s\(^{-1}\).

In addition to the seasonal controls dictated by the inherent plant metabolic activity, low temperatures (<10°C) strongly modulated isoprene emissions. The frequent low temperatures observed at the boreal forest caused substantial reductions in the diurnal amplitude of isoprene emissions. Even after the forest had attained its maximum capacity to emit isoprene, following the occurrence of low temperatures, isoprene emissions were suppressed despite favorable temperature regimes to promote large fluxes. Reduced isoprene emissions resulted likely due to diminished enzymatic activity in leaves.

Using a multilayered canopy model, we confirmed that realistic isoprene emissions can be obtained for forests if emission rates, active biomass, and influencing environmental variables are properly characterized as they vary with depth in the canopy. Given that emissions substantially change with growing season, it is necessary to determine appropriate emission rates to derive inventories for either short- or long-term temporal scales. In this study, we found that with an averaged emission rate (~15 nmol m\(^{-2}\) s\(^{-1}\)), characterizing prevailing emissions throughout the season, the calculated fluxes exceeded the measured quantities by 12%. However, emission rates representing the middle of the growing season yielded poorer results and the uncertainties exceeded 65%. These results demonstrate the inadequacies of using a single emission rate to characterize seasonal isoprene emissions. To obtain improved emission inventories, modeling systems need to incorporate seasonally adjusted emission factors. In the form of isoprene, the boreal aspen forest lost approximately 1% of the carbon assimilated through photosynthesis. On average, this did not represent a significant amount of carbon lost but values exceeded 3% when the temperature was higher than 20°C.

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