Spatial and Temporal Patterns of CO$_2$, CH$_4$, and N$_2$O Fluxes at the Soil-Atmosphere Interface in a Northern Temperate Forested Watershed

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Abstract
Forest soils are recognized as sources and sinks of Greenhouse gases (GHG) (CO$_2$, CH$_4$, N$_2$O), but there are limited data quantifying the magnitude of GHG fluxes at the soil-atmosphere interface across a range of landscape hydrogeomorphic conditions. In our study, GHG fluxes were measured in a forested watershed across a range of hydrogeomorphic locations (wetlands, hillslopes, riparian zones, etc) and evaluated in relation to temperature, antecedent flow conditions, and stream chemistry to help develop strategies to scale GHG emissions from the point scale to the watershed scale. Mean study period CO$_2$ fluxes (0.61 to 2.89 gCm$^{-2}$d$^{-1}$) were positive at all sites, with larger fluxes occurring in well-drained soils. Negative fluxes (CH$_4$ sinks) were found at the hillslope and lowland sites, while the wetland was a large source of CH$_4$ emissions at the watershed scale. Mean CH$_4$ fluxes ranged from -2.53 to 330.34 mgCm$^{-2}$d$^{-1}$. Nitrous oxide fluxes were low relative to other GHG fluxes (in terms of CO$_2$ equivalent) and ranged between -0.72 to 0.70 mgNm$^{-2}$d$^{-1}$. Although carbon dioxide fluxes were positively correlated to soil temperature at all locations, CH$_4$ and N$_2$O fluxes were not significantly related to temperature, antecedent flow conditions, or stream chemistry at the watershed scale. However, strong differences in CO$_2$ and CH$_4$ fluxes related to landscape geomorphology were observed, and exceeded the magnitude of seasonal variations for CO$_2$ and CH$_4$ fluxes, suggesting that landscape hydrogeomorphology was likely a stronger predictor of GHG fluxes at the watershed scale than temperature and stream chemistry variables, at least within the confine of one watershed. In lieu of statistical approaches relying on environmental variables to predict GHG fluxes at the watershed scale, geomorphological approaches, potentially coupled with seasonal comparisons of GHG fluxes in each land class, might therefore be a promising research avenue to provide solid watershed wide estimates of GHG fluxes.

Keywords
Greenhouse gases, Forest soils, Carbon dioxide, Methane, Nitrous oxide, Geomorphology

Introduction

A comprehensive understanding of soil contributions to atmospheric greenhouse gas (CO$_2$, CH$_4$, and N$_2$O) concentrations is necessary to refine estimates of Greenhouse gas (GHG) budgets. Globally, CO$_2$ fluxes from soils are primarily driven by microbial and autotrophic respiration [1]. In contrast, CH$_4$ can be consumed through methane oxidation (under aerobic conditions) or produced via methanogenesis primarily in wetlands under anoxic conditions [2,3]. Globally, wetlands may account up to 15-40% of global CH$_4$ emissions or ~170 Tg CH$_4$ yr$^{-1}$, while aerobic upland soils are estimated to consume approximately 30 Tg CH$_4$ yr$^{-1}$ [4,5]. Nitrous oxide emissions from soils are also significant and are thought to account for up to two-thirds of the annual
global emissions annually (6.6 Tg N yr⁻¹; Anderson, et al. [5]). Nitrous oxide is the product of denitrification (dissimilatory) and nitrification (assimilatory); the former occurring under reducing conditions and the latter occurring under aerobic conditions [6]. However, in spite of the importance of GHG emission from soils at regulating GHG concentrations in the atmosphere, few studies have fully assessed the spatial and temporal variability of GHG fluxes from forest soils at the watershed scale, resulting in uncertainty in regional and global GHG budgets [7,8].

At the soil scale, the source/sink function and the magnitude of the GHG fluxes at the soil-atmosphere interface is mainly a function of physical controls on the biogeochemical processes of soil respiration, denitrification, nitrification, methanogenesis, and methane oxidation [9]. These biogeochemical reactions are strongly influenced by soil temperature, soil moisture, and the distribution of electron donors and acceptors in the soil profile [10,11]. For instance, soil temperature and soil water content are both directly positively related to rates of aerobic respiration in the soil profile, though these relationships can vary across space and time, and can change if temperature or moisture content become too high or too low [12-14]. Abundant soil organic matter coupled with anoxic conditions result in high rates of methanogenesis and associated CH₄ emissions [3]. The availability of organic carbon (i.e., organic matter) as an electron donor also often positively correlates with denitrification activity, while nitrate concentration and soil temperature often positively correlate with N₂O emissions [15].

At the watershed scale, forest soils are recognized as both a sink and a source of CO₂, CH₄, and N₂O [9,16], with landscape position (e.g., wetlands, hillslopes, and riparian areas) controlling much of the source/sink function at specific locations [13,17]. For instance, Riveros-Iregui and McGlynn [17] showed that soil respiration rates can be organized by the structure and morphology of the landscape (i.e., riparian zones versus hillslopes). Pacific, et al. [13] also showed that the landscape structure, along with groundwater dynamics and soil water content, influenced the rates of CO₂ flux in a forested watershed in Montana, USA. Other studies indicate that upland soil environments are often CH₄ sinks [18], that wetlands are large CH₄ sources [19], and that riparian areas can significantly contribute to N₂O fluxes [11,20]. However, in spite of the known importance of landscape position in regulating GHG emissions from forest soils, capturing the variability of gas fluxes at the soil-atmosphere interface at the watershed scale still remains a challenge due to the known variability in biogeochemical activity occurring throughout the landscape [11]. Therefore, further characterizing the relationship(s) between GHG fluxes across time, space, and landscape characteristics in northern forested watersheds is important to provide a more comprehensive understanding of the dynamics of these gas fluxes at the watershed scale [21].

Additionally, along with a detailed characterization of GHG fluxes across landscape hydrogeomorphic classes (wetlands, riparian zones, lower hillslopes, upper hillslopes), linking stream chemistry to GHG emissions at the watershed scale might be a promising research avenue to generalize GHG emissions at the soil-atmosphere interface at the watershed scale as stream chemistry is often a good indicator of dominant biogeochemical processes at the watershed scale [13,22]. Indeed, measuring soil moisture, temperature, and the distribution of various electron donors and acceptors at many locations in a watershed on a routine basis is not a reasonable approach to scale GHG emission at the watershed scale for logistical reasons (cost, man-hours…). Determining to what extent commonly measured stream chemistry parameters at the outlet of our study watershed could be used (or not) to constrain GHG emission at the watershed scale for scaling purpose therefore has the potential to significantly advance our ability to estimate GHG emission from soil for management purposes.

Our study therefore addresses the following questions: (1) How do CH₄, CO₂, and N₂O fluxes at the soil-atmosphere interface change over time and across hydrogeomorphically distinct locations in a northern temperate forested watershed? (2) Can stream discharge, temperature, and stream chemistry be used as predictors of soil CH₄, CO₂, and N₂O fluxes across locations in a watershed? The implications of our results for developing greenhouse gas budgets at the watershed scale are also discussed.

Materials and Methods

Study area

The Archer Creek Watershed (515 m above sea level, 43°59’32” N, 74°14’32” W) is a 135 ha forested catchment (1% water, 6% forested wetland, 93% forest) located in the Adirondack region of northern New York State (Figure 1). Climate is cool, moist, and continental. Between the years of 1941-2007, the mean annual temperature was 5.0°C, the mean annual precipitation was 1046 mm, and the mean annual snowfall was 303 cm [23]. Precipitation in 2011 was 47% greater than the 30-year average, 8% less than the 30-year average in 2012, and 12% greater than the 30-year average in 2013. Snowfall was less than the 30-year average for the entire study period with 3%, 52%, and 40% less snowfall occurring in 2011, 2012, and 2013 respectively. Precipitation and air temperature were recorded at the National Atmospheric Deposition
Program/National Trends Network Site (ID NY20) located in the Huntington Wildlife Forest approximately 2.0 km from the study sites (Figure 1).

From a geomorphological standpoint, the Archer Creek catchment has an average slope of 11%, and a total relief of 225 meters [23]. The surficial geology consists of glacial till composed of approximately 75% sand, and less than 10% clay [24]. Upland soils are sandy loam (Becket-Mundal series) approximately one meter thick with localized areas of deeper soil. Wetlands and valley-bottom soils are Greenwood Mucky Peats deposits 1 to 5 meters deep [25]. The lowland overstory in the Archer Creek Watershed is composed of eastern hemlock (Tsuga canadensis (L.) Carr), yellow birch (Betula alleghaniensis Britt.), speckled alder (Alnus incana (L.) Moench), red spruce (Picea rubens Sarg.), and balsam fir (Abies balsamea (L.) Mill). American beech (Fagus grandifolia Ehrh.), sugar maple (Acer saccharum Marsh.), red maple (Acer rubrum L.), yellow birch, and eastern white pine (Pinus strobus L.) dominate the upland environments [24].

**Watershed hydrology and stream water chemistry data**

Discharge data for Archer Creek (15 min. interval) were calculated for the outlet of Archer Creek watershed where stage height is recorded as part of the on-going long-term monitoring program of the watershed initially described in Mitchell, et al. [23]. One day (1dQ), 3-day (3dQ) and 7-day (7dQ) antecedent flow conditions at the outlet of the Archer Creek watershed were used as proxies for antecedent moisture conditions in the watershed.

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**Figure 1:** Location of Archer Creek Watershed in New York State, USA and of the 10 sampling sites in Archer Creek Watershed.
and calculated as the mean discharge 1, 3, and 7 days before each sampling date [26]. Stream Dissolved organic carbon (DOC) and nitrate (NO₃⁻) concentrations at the Archer Creek H-Flume were collected weekly as part of the on-going long-term monitoring program described above.

**Greenhouse gas sampling and analysis**

Instead of using a traditional statistically based sampling design (e.g. power analysis, random sampling) that would require sampling many areas of the watershed and render our study undoable for logistical reasons (access, cost), we used a geomorphologically driven sampling design (see below) based on the fact that previous studies have shown that such approaches can be suitable to generalize GHG emission estimates at the watershed scale [13, 17]. Ten sampling locations corresponding to a variety of landscape hydrogeomorphic classes (palustrine wetland, lowlands, riparian zones, upper and lower hillslopes, and headwater wetlands; Figure 1) were therefore identified throughout the Archer Creek Watershed for Greenhouse gas (GHG) sampling at the soil-atmosphere interface.

The lowland was located in a gently sloping area near the palustrine wetland. The riparian zone locations were located immediately adjacent to headwater streams, while the headwater wetlands were located near the source of the two streams used in this study (Figure 1). The upper hillslope sites were located on generally steep terrain (>20% slope) while the lower hillslope sites were located downstream of the headwater sites on more moderate slopes (<20% slope). At each of these sites, greenhouse gas (CO₂, CH₄, N₂O) sampling occurred 19 times during the 2011, 2012, and 2013 snow free seasons, with an additional three winter sampling days in January, February, and March of 2013. Gas samples for GHG flux calculations at the soil-atmosphere interface were collected using 6 static chambers per site, for a total of 60 chambers (Figure 1). Static chambers are commonly used for GHG flux estimates at specific times and places, and have the advantage of being disconnected from atmospheric flow, which is critical for accurate flux measurements at the soil-atmosphere interface [27]. Static chambers were made of PVC material, and consisted of two parts: a bottom section (37 cm height and 27 cm diameter) inserted 10 cm into the ground, and an airtight lid fitted with a gas sampling port [28]. At the time of sampling, soil temperature was recorded at each sampling location 5 cm below the soil surface using a portable Hanna® pH/ORP/temperature meter (Model HI 9125, Hanna Instrument Inc., Woonsocket, Rhode Island, USA). When sampling, the chambers were closed with a lid and headspace gas concentrations measured 3 times over 50 min. Air samples (~15 mL) were stored in evacuated vials (10 mL) fitted with grey butyl rubber septa. GHG fluxes (F) were computed as:

\[ F = \left( \frac{dC}{dt} \right) \cdot \left( \frac{V}{A} \right) \cdot k \]  

Where \( \frac{dC}{dt} \) is the rate of change in GHG concentration inside the chamber (mass GHG m⁻³ air per min), V is the chamber volume (m³), A is the area circumscribed by the chamber (m²), and k is a unit conversion factor (1,440 min/day) [27]. Winter sampling was carried out following the same protocol, except that snow was removed by hand from each chamber immediately before the airtight lid was placed onto the chamber, and the snow was returned into the chamber immediately after sampling. Samples were kept out of the light during transport and storage periods, and were usually analyzed within two weeks of collection. Carbon dioxide, CH₄, and N₂O concentrations were analyzed utilizing a Shimadzu GC-2014® gas chromatograph (Shimadzu Corporation, Kyoto, Japan) equipped with a flame ionizing detector and an electron capture detector interfaced with a CombiPal® autosampler (LEAP Technologies, Carrboro, North Carolina). The stationary phase consisted of a Porapak Q® precolumn (90 cm long) and HayeSep D® analytical columns (180 cm long). The GC was calibrated using standard gases obtained from Alltech (Deerfield, Illinois). To prevent moisture buildup in the GC columns, the GC oven was heated (150 °C) for 15 min for every 20 samples.

We tested for the importance of sampling time on daily soil GHG flux estimates by conducting four intensive sampling campaigns (5 hour intervals over a 24 hour period) on September 6 and October 20, 2012, and June 6 and July 16, 2013, at one chamber per site (for logistical reasons) at sites RZ1, LH1, RZ2, LH2, HW2 and UH2 (Figure 1). Carbon dioxide, CH₄, and N₂O fluxes varied over a 24-hour period, however, sampling time did not consistently affect flux estimates.

**Data analysis**

Because of the high spatial heterogeneity of GHG emission within each site (e.g. RZ1, LH1, RZ2, LH2, HW2), CO₂, CH₄, and N₂O fluxes at each site (for each sampling date) were determined by taking the median flux of the six chambers at each site for each sampling date [20,29]; however, when multiple dates were aggregated to identify either annual or seasonal average, the arithmetic mean of GHG fluxes for each site and each date was used. For inter-site comparison purposes, a Kruskal-Wallis test was used to compare median fluxes of CH₄, CO₂, and N₂O among the 10 study sites. If a significant Kruskal-Wallis statistic (p < 0.05) was observed, pairwise differences among the 10 study sites were evaluated using the Two Sample Mann-Whitney test.
Spearman rank correlation analysis [30] was used to assess relationships between CO₂, CH₄, or N₂O flux at each site with soil temperature, stream nitrate (NO₃⁻), stream Dissolved organic carbon (DOC), and antecedent flow conditions (1dQ, 3dQ, and 7dQ).

**Results**

**Watershed hydrology and stream water chemistry**

Over the course of the study, the specific discharge at the Archer Creek outlet was 2.02 mm d⁻¹ (Figure 2). Most sampling dates (18 out of 21) occurred at time when stream flow was below 2.02 mm d⁻¹. On three occasions (D6, D15, D21), sampling occurred following intense storm events as observed by a sharp decrease in antecedent flow conditions in the days preceding sampling (i.e., 7dQ > 3dQ > 1dQ) (Figure 3).

Soil temperature varied seasonally with winter (January, February, March) temperatures below 4 °C, and spring (April, May, June) and autumn (October, November, December) temperatures between 4 °C to 14 °C. Summer soil temperatures (July, August, September) were higher and varied between 14 °C to 18 °C. During winter sampling, soils were not frozen on D17 (January 7, 2013) and were generally frozen at the surface on D18 (February 15, 2013) except at the HW1 site due to relatively warm discharging groundwater in the vicinity. The LH1, HW1, UH2, and LH2 sites remained unfrozen on D19 (March 8, 2013), while the soils at the other sites were frozen on this date.

Snowpack depth ranged from 20-61 cm with an average depth of 41 cm throughout the D17, D18, and D19 field sampling dates.

Over the study period, stream DOC concentrations...
at the Archer Creek outlet ranged from 264.0 to 1076.4 µmolCL⁻¹ (3.16 to 12.92 mgCL⁻¹), and NO₃⁻ concentrations ranged from 4.30 to 52 µmol NO₃⁻ L⁻¹ (0.06-0.73 mg N L⁻¹) (Figure 4). In spring, summer, and fall 2011 and spring and summer 2012, NO₃⁻ concentrations generally remained below 20 µmolL⁻¹ (0.28 mgL⁻¹). However, NO₃⁻ concentration increased through Fall 2012, and peaked during winter 2013 (D17, D18, and D19). The winter peak in NO₃⁻ concentrations (52.0 µmol NO₃⁻ L⁻¹ or 7.28 mg NO₃⁻ L⁻¹) coincided with a clear decline in DOC concentrations to 264 µmolL -1 (3.16 mgL -1) from a high of 1080 µmolL -1 (12.9 mgL -1) in September 2012 (D14). Av-

Table 1: Arithmetic mean and Relative standard deviation (RSD) for CO₂ (g C-CO₂ m⁻²d⁻¹), CH₄ (mg C-CH₄ m⁻²d⁻¹) and N₂O (mg N-N₂O m⁻²d⁻¹) fluxes at the Wetland, Lowland, RZ1-Riparian Zone, LH1-Lower Hillslope, HW1-Headwater Wetland, UH1-Upper Hillslope, RZ2 Riparian Zone, LH2-Lower Hillslope, HW2-Headwater Wetland (HW), and UH2-Upper Hillslope (UH) sites for the duration of the study period (June 2011-July 2013).

| Site                  | CO₂ flux | CH₄ flux | N₂O flux |
|-----------------------|----------|----------|----------|
|                       | Mean     | RSD (%)  | Mean     | RSD (%)  | Mean     | RSD (%)  |
| Wetland               | 0.61     | 111      | 330.34   | 127      | 0.07     | 152      |
| Lowland               | 2.74     | 62       | -0.54    | 376      | 0.01     | 2741     |
| RZ1-Riparian Zone     | 0.62     | 79       | 16.17    | 368      | 0.18     | 189      |
| LH1-Lower Hillslope   | 2.38     | 65       | -1.14    | 114      | 0.19     | 106      |
| HW1-Headwater Wetland | 1.41     | 66       | -0.92    | 105      | -0.72    | 606      |
| UH1-Upper Hillslope   | 2.89     | 61       | -2.53    | 57       | 0.03     | 697      |
| RZ2-Riparian Zone     | 1.1      | 55       | -0.59    | 135      | 0.07     | 408      |
| LH2-Lower Hillslope   | 2.12     | 57       | -2.13    | 105      | 0.16     | 728      |
| HW2-Headwater Wetland | 1.94     | 59       | -1.2     | 71       | 0.09     | 274      |
| UH2-Upper Hillslope   | 1.72     | 55       | 0.51     | 361      | 0.70     | 163      |
average stream DOC concentration over the study period was 578 µmolL⁻¹ (6.9 mgCL⁻¹).

**Variation of GHG fluxes across landscape positions**

During the duration of the study, CO₂ fluxes at the soil-atmosphere interface were generally positive (Table 1), although negative values were observed on specific dates/locations (Figure 5A). Mean CO₂ fluxes among the sampling sites during the winter sampling dates D17, D18, and D19 were significantly smaller and less variable (0.02 to 1.34 g C-CO₂ m⁻²d⁻¹) than during the remainder of the year (-0.77 to 6.87 g C-CO₂ m⁻²d⁻¹). When sites were compared, the lowland, LH1, UH1, and LH2 sites presented mean (across dates) CO₂ fluxes greater than 2 g C-CO₂ m⁻²d⁻¹ over the study period, while mean CO₂ fluxes at the wetland and RZ1 site were less than 1 g C-CO₂ m⁻²d⁻¹ (Table 1). Other sites (HW1, RZ2, HW2, and UH2) presented intermediary CO₂ flux values between 1 and 2 g C-CO₂ m⁻²d⁻¹. Overall, CO₂ fluxes varied significantly across the ten study sites (χ² (9, N = 219) = 73.43; p < 0.0001) with pairwise comparisons resulting in significantly different median CO₂ fluxes (using α =

![Figure 5](image-url)
0.05) for most site pairs. The only exceptions were the lowland, LH1, UH1, and LH2 sites for which no differences in CO$_2$ fluxes were observed (p < 0.05). The following sites also showed no significant differences (p < 0.05) between one another in term of CO$_2$ fluxes: the wetland and the RZ1 site, the LH2 and HW2 sites, and the HW2 and UH2 sites.

Methane (CH$_4$) fluxes at the soil-atmosphere interface were consistently positive at the wetland site (range: 2.2 to 1293 mg C m$^{-2}$ d$^{-1}$), with the largest fluxes occurring in the summer and early autumn (July through October). Relatively low fluxes below 15 mgCm$^{-2}$d$^{-1}$ at the wetland site occurred in the winter (D17, D18, D19), and each spring (D1, D4, D20). Positive CH$_4$ fluxes were also observed at the RZ1 site for 18 of the 22 sampling dates, with a temporal pattern similar to the one observed in the wetland site. At the other sampling locations, CH$_4$ fluxes were generally negative on most dates (Figure 5). When sites were compared, the average study period CH$_4$ flux at the wetland site (330 mgC m$^{-2}$ d$^{-1}$; RSD 127%) was two orders of magnitude higher than the average CH$_4$ flux at any other site. A Kruskal-Wallis test indicated that CH$_4$ fluxes differed significantly across the ten study sites ($\chi^2$ (9,219) = 130; p < 0.0001). Wetland CH$_4$ fluxes differed significantly from each of the other study sites (α < 0.05), while the lowland, LH1, UH1, and LH2 had similar median fluxes.

Soil N$_2$O fluxes generally ranged from -0.5 to 0.5 mg N m$^{-2}$ d$^{-1}$ for most of the study period (Figure 5). Both the highest (4.9 mg N m$^{-2}$ d$^{-1}$ at the LH2 site) and the lowest (-20.2 mg N m$^{-2}$ d$^{-1}$ at the HW1) observed values across all sites during the study occurred on the same date (D17-January 7, 2013). Over the study period, pairwise comparisons resulted in significantly different median N$_2$O fluxes (α = 0.05) between most locations, except between the RZ1, RZ2, HW1, and HW2 sites. The median wetland and lowland fluxes were also not statistically different from each other either.

**Relationships between CO$_2$, CH$_4$, and N$_2$O fluxes and watershed conditions**

Over the study period, CO$_2$ flux was positively correlated with temperature both on a site-by-site basis ($r = 0.47$ to 0.83; all p < 0.05; Figure 6) and across all locations ($r = 0.50$, n = 203, p < 0.001). With respect to NO$_3^-$, it was only when all sampling dates were grouped together (n = 203) that CO$_2$ fluxes were significantly correlated to stream NO$_3^-$ ($r = -0.24$, n = 203, p < 0.05) and DOC ($r = 0.28$, n = 203, p < 0.05). No significant correlation was observed between antecedent flow conditions and CO$_2$ fluxes at any of the sites. As for CH$_4$ and N$_2$O, temperature was significantly correlated with CH$_4$ flux at the wetland site ($r = 0.51$, n = 20, p < 0.05), and to N$_2$O ($r = 0.47$, n = 20, p < 0.05) at the HW1 site. However, no significant correlations were observed between N$_2$O and temperature (except at the HW1 site) or 1dQ, 3dQ, and 7dQ at any of the sites. Antecedent flow conditions were also generally not significantly correlated to CH$_4$ except at the UH2 site ($r = -0.46$ with 1dQ, $r = -0.54$ with 3dQ) and the RZ2 site ($r = 0.46$ with 3dQ). Methane fluxes were only significantly correlated to NO$_3^-$ at the RZ2 site ($r = -0.47$) and to DOC at the wetland ($r = 0.47$, n =

**Figure 6:** Spearman correlation coefficients of soil temperature, Archer Creek nitrate (NO$_3^-$) concentrations, Archer Creek Dissolved organic carbon (DOC) concentrations, and 1-day (1dQ), 3-day (3dQ), and 7-day (7dQ) antecedent flow conditions in Archer Creek with CO$_2$ (a), CH$_4$ (b), and N$_2$O (c) fluxes.
203, p < 0.05) and RZ2 (r = 0.42, n = 22, p < 0.05) sites. Nitrous oxide flux was significantly correlated with NO$_3^-$ concentrations only at the LH2 site (r = 0.45 n = 21, p < 0.05), but was not related to DOC stream concentrations.

Discussion

Impact of landscape position on overall CH$_4$, CO$_2$, and N$_2$O fluxes at the soil-atmosphere interface

The CO$_2$ fluxes reported across the range of hydrogeomorphic locations studied here are consistent with other studies. For instance, CO$_2$ fluxes reported here are comparable to those reported in a northern Rocky Mountain watershed (2.7 to 18.07 gCm$^{-2}$d$^{-1}$) [17] and similar to those reported during the growing season in the Harvard Forest, MA, USA (0.84 to 2.88 gCm$^{-2}$d$^{-1}$) [31]. On a site by site basis, CO$_2$ emissions from the hillslope sites are consistent with those reported by Ullah and Moore [16] for well-drained soils in a deciduous forest in eastern Canada (1.4 to 2.9 gCm$^{-2}$d$^{-1}$). From a biogeochemical process standpoint, the riparian zone and wetland soils had the smallest average fluxes of CO$_2$ across the study period, which is consistent with depressed aerobic respiration compared to the other sites owing to high soil moisture at these sites, and CO$_2$ consumption via methanogenesis [12,19,32]. In contrast, the hillslope and lowland soils consistently had the largest amounts of CO$_2$ per unit area, which is consistent with aerobic, yet moist conditions, observed at these sites over the duration of the study. Fluxes of CO$_2$ from headwater wetlands, which were at times either saturated or near saturated (field observations), presented intermediate CO$_2$ values. These moderate fluxes in headwater wetlands may be attributed to a combination of periods of low moisture during which soils were drier and aerobic (higher flux) and periods of high moisture conditions when soils were saturated and anaerobic (lower flux).

Local hydrologic and geomorphic conditions also played an important role at regulating CH$_4$ fluxes across the forest landscape. For instance, wetlands and seasonally saturated soils were CH$_4$ sources, while upland and well-drained soils were CH$_4$ sinks, which is consistent with our current understanding of the impact of landscape geomorphology on CH$_4$ emission and results published in other studies [3,16,33]. More specifically, the wetland site exhibited an average CH$_4$ flux of 330.34 (RSD 127%) mgCm$^{-2}$d$^{-1}$, which is approximately three times the average global flux estimated by Turetsky, et al. [33] from a study of boreal, temperate and subtropical wetlands, but lower than CH$_4$ fluxes (>5,000 mgCm$^{-2}$d$^{-1}$) reported for temperate peatlands in the same study. The CH$_4$ fluxes observed at the RZ1 site (16.17 mgCm$^{-2}$d$^{-1}$) where organic rich mucky soils dominated are consistent with values reported by Morse, et al. [3] in restored agricultural wetlands (range: 16.2 to 21.6 mgCm$^{-2}$d$^{-1}$). Methane fluxes at sites other than the Wetland or RZ1 sites were negative (-2.53 to -0.54 mgCm$^{-2}$d$^{-1}$), which is consistent with fluxes reported in forested hillslope sites in eastern Canada (-1.7 ± 0.1 mgCm$^{-2}$d$^{-1}$) [16] and with the aerobic conditions and associated CH$_4$ oxidation often observed in well drained forested soils [18]. The only exception is the upper hillslope 2 site (UH2), which had a slightly positive CH$_4$ flux (0.51 mgCm$^{-2}$d$^{-1}$).

Unlike CO$_2$ and CH$_4$, no relationship could be established between N$_2$O fluxes and landscape position in our study (Table 1). Such heterogeneity in N$_2$O fluxes has been highlighted by others and tied to the fact that both nitrification (under aerobic conditions) and denitrification (under anaerobic conditions) can lead to N$_2$O production [19,34]. Our results are nevertheless consistent with values reported from mixed deciduous forests in the eastern USA [27], and an infrequently flooded riparian forest in Indiana, USA [35].

Impact of seasonality on CH$_4$, CO$_2$, and N$_2$O fluxes at the soil-atmosphere interface

Higher CO$_2$ and CH$_4$ fluxes during the summer, and lower fluxes during winter and spring, are consistent with prior studies that indicated the importance of temperature-mediated controls on respiration and methanogenesis [16,34]. In the rare cases when relatively low CO$_2$ fluxes occurred in summer during warm conditions (>15°C), those were concomitant with relatively dry periods (low antecedent flow), suggesting that moisture could become a limiting factor to CO$_2$ production in summer. This is consistent with results reported by Rive ros-Iregui, et al. [36] in a forested watershed in Montana showing that moisture can become a limiting factor for CO$_2$ flux rates by regulating the amount of microbial activity occurring in the soil profile. With respect to CH$_4$, highest CH$_4$ fluxes occurred on summer/fall dates with high antecedent flow conditions (D7, D12, and D13), which again is consistent with other studies showing that warm temperature along with high antecedent moisture conditions are both needed for very high CH$_4$ emissions to occur [3,29,34].

Unlike for CO$_2$ and CH$_4$, no seasonal patterns were observed for N$_2$O fluxes, which may be due to fine-scale temporal variability in the watershed conditions affecting both nitrification and denitrification [37]. Vidon, et al. [34] also observed a lack of clear seasonal patterns in N$_2$O flux in a restored wetland in Indiana, USA. However, unlike several studies that found increases in N$_2$O flux after intense re-wetting events [3,34,35], we observed no significant statistical or consistent qualitative relationships between N$_2$O fluxes and antecedent flow conditions across the sites.
Impact of temperature, stream discharge, and stream chemistry on CH$_4$, CO$_2$, and N$_2$O fluxes

The soil temperature-CO$_2$ relationship over the study period ($r = 0.50$, $n = 203$, $p < 0.001$) indicated in our results is consistent with the CO$_2$ flux-temperature relationships found at the Harvard forest in Massachusetts, USA [12], and in a mixed hardwood forest in Korea [38], suggesting that temperature correlations with CO$_2$ flux can be useful in assessing possible behaviors of forest soils in response to changing temperature conditions. The stronger CO$_2$-temperature correlations observed at some sites (e.g. $r = 0.83$, $n = 21$, $p < 0.05$ at the RZ2 site) are consistent with a study in Montana showing that the explanatory power of a CO$_2$-temperature regression analysis is often muted when considered across the watershed scale compared to individual sites [13]. The lack of significant correlation between CO$_2$ and 1dQ, 3dQ or 7dQ at any location suggests that, at least for our study period, moisture was unlikely to be a strong limiting factor for aerobic respiration. However, the significant correlations between CO$_2$ fluxes across all sites over the study period and DOC ($r = 0.28$, $n = 203$, $p < 0.05$) and NO$_3^-$ ($r = -0.24$, $n = 203$, $p < 0.05$) suggest that stream water quality does have an impact of overall CO$_2$ fluxes at the watershed scale.

At sites where CH$_4$ fluxes were negative, temperature was not significantly correlated to CH$_4$ fluxes, but temperature was significantly correlated to CH$_4$ at the wetland site ($r = 0.51$, $n = 20$, $p < 0.05$). This suggests that in locations where large CH$_4$ fluxes due to methanogenesis dominate (e.g. peatlands), CH$_4$-temperature regression curves might be a suitable tool to help generalizing CH$_4$ emissions at the landscape scale, but that such relationships are less likely to help constrain CH$_4$ emissions in complex landscapes where both methane oxidation and methanogenesis actively occur. Similarly, although our analysis of seasonality illustrates the need for both high temperature (summer) and high moisture conditions (wetlands) to co-occur for the highest CH$_4$ fluxes to occur, antecedent moisture conditions were not a strong predictor for CH$_4$ fluxes across the watershed. The lack of significant correlation between N$_2$O and either temperature or antecedent moisture conditions is consistent with other studies showing somewhat erratic N$_2$O flux patterns due to fine-scale temporal variability in the watershed conditions affecting both nitrification and denitrification [37]. These results suggest that much more research is needed to identify landscape scale variables capable of predicting N$_2$O fluxes at the watershed scale in a variety of landscapes.

Implications for GHG budget estimation at the watershed scale and conclusions

Our study indicates that estimating N$_2$O fluxes at the watershed scale remains a challenge, as neither landscape geomorphology nor environmental conditions (temperature, water chemistry, and antecedent flow conditions) appear to be strong predictors of N$_2$O emissions. However, our results indicate that temperature for CO$_2$, and to some extent stream NO$_3^-$ and DOC for CO$_2$ at the watershed scale, were significantly correlated to CO$_2$ fluxes. Similarly, in the wetland (high organic carbon content, anaerobic conditions), the significant positive correlation between CH$_4$ and soil temperature suggests that temperature could be used to constrain CH$_4$ emissions at such locations. However, data suggest that landscape geomorphology based approaches likely hold more promise than temperature or stream chemistry based approaches as tools to scale up CO$_2$ and CH$_4$ point flux measurements for GHG budget development at the watershed scale. Indeed, as indicated in the results, strong differences in CO$_2$ and CH$_4$ fluxes related to landscape geomorphology were observed (e.g. wetlands vs. other locations for CH$_4$; low riparian and wetland CO$_2$ fluxes compared to those at other locations, etc), and the magnitude of these differences exceeded the magnitude of seasonal variations for CO$_2$ and CH$_4$ fluxes (Figure 5 and Table 1), and this, in spite of the potential importance of temperature at regulating CO$_2$ across the watershed and CH$_4$ fluxes in the wetland (Figure 6). This indicates that landscape hydrogeomorphology is likely a stronger predictor of GHG fluxes at the watershed scale than environmental variables (temperature and stream chemistry), at least within the confines of one watershed. As we reach this conclusion, it is important to note that we understand that on a site-by-site basis, alternative approaches such as measuring soil moisture, temperature, and the distribution of various electron donors and acceptors at individual locations where GHG fluxes are measured may allow for the development of predictive relationships on a case-by-case basis. However, such approaches are not suitable for scaling purposes for logistical reasons (cost, man-hours…). We therefore propose that landscape based approaches aimed at identifying the relative importance of land classes (e.g. wetland versus riparian zone versus hillslope), potentially coupled with long-term analyses of the influences of seasonality on GHG emissions in each land class, are likely to lead to the best results in determining the long-term contributions of forest soils to GHG emissions at the watershed scale. In a companion paper [39], we therefore explore how GHG budgets can be developed for a whole watershed based on a geomorphic classification of the landscape (beyond the scope of this paper), so this study investigating the relative importance of landscape position/geomorphology and environmental variables (temperature, stream chemistry) at predicting GHG emission at the watershed scale can significantly advance our ability.
to scale up point measurements of GHG emission to a whole watershed. In that paper, we also demonstrate that the \( \text{N}_2\text{O} \) fluxes presented here only represent a couple percentage points of total \( \text{N}_2\text{O} \), \( \text{CO}_2 \), and \( \text{CH}_4 \) emissions (in \( \text{CO}_2 \) equivalent) at the watershed scale, and that our limited ability to solidly predict \( \text{N}_2\text{O} \) fluxes at the landscape scale is therefore unlikely to affect overall estimates of GHG production (in \( \text{CO}_2 \) equivalent) at the soil-atmosphere interface at the watershed scale.

Acknowledgements

This project was supported by a USDA McIntire-Stennis Formula Grant (award # NYZ:2611-20-001) to P. Vidon, M. Mitchell, and C. Beier. The support of NYSERDA in providing funds for the monitoring at the Arbutus Watershed and Huntington Forest is also greatly appreciated. The authors would also like to thank Pat McHale, SatishSerchan, Cheryl Glor, and the staff of the Adirondack Ecological Center in Newcomb, NY, for help in the field and laboratory.

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