1	Nitrous oxide consumption potentials of well-drained forest soils in
2	southern Québec, Canada
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21	denitrification; N <sub>2</sub> O fluxes

#### 22 ABSTRACT

- 23 To establish the major controls on N<sub>2</sub>O consumption by forest soils, we conducted
- 24 laboratory incubations of sixteen samples from four soil types, two organic and two
- 25 mineral, varying in overlying forest vegetation (sugar maple, American beech and eastern
- 26 hemlock). The fastest potential consumption of N<sub>2</sub>O occurred under anoxic conditions
- 27 with little soil nitrate and under elevated headspace N<sub>2</sub>O concentration. Potential N<sub>2</sub>O
- 28 consumption rates were fastest in organic soils under hemlock and beech trees (111 and
- 29 75 ng N<sub>2</sub>O-N  $g^{-1} d^{-1}$ , respectively) compared to mineral soils under beech and maple trees
- 30 (45 and 41 ng N<sub>2</sub>O-N  $g^{-1} d^{-1}$ ). Organic soils showed faster N<sub>2</sub>O consumption rates than
- 31 mineral soils, possibly due to larger organic C levels and higher C:N ratios. Acetylene
- 32 treatment confirmed that denitrification was the process underlying N<sub>2</sub>O consumption.
- 33 These results suggest that soils regularly consume N<sub>2</sub>O with varying magnitude, most
- 34 likely in anoxic microsites throughout the soil profile and that the potential for  $N_2O$

35 consumption is larger in organic than in mineral forest soils.

#### 37 INTRODUCTION

Soils emit nitrous oxide (N<sub>2</sub>O), a greenhouse gas, into the atmosphere and account for 10 of the 16 Tg nitrogen (N) of the total N<sub>2</sub>O released into the atmosphere each year (IPCC 2001; IPCC 2007). Approximately 4 Tg comes from agricultural soils, thus of anthropogenic origin, while the remaining 6 Tg are attributed to emissions from soils under natural ecosystems (IPCC 2001; IPCC 2007). Although forest soils are net sources of N<sub>2</sub>O to the atmosphere, there is evidence that soils may also consume atmospheric N<sub>2</sub>O (Arah *et al.* 1991).

45 The capacity of soils to act as sources or sinks of N<sub>2</sub>O is the result of dynamic 46 microbial processes of consumption and production occurring within the soil profile 47 (Chapuis-Lardy et al. 2007). Denitrification and nitrification are the two dominant 48 mechanisms of  $N_2O$  production; other biological and abiological processes (such as 49 assimilatory and dissimilatory nitrate reduction and chemodenitrification) are thought to 50 contribute < 1% of N<sub>2</sub>O emissions (Chapuis-Lardy *et al.* 2007). The mechanisms of N<sub>2</sub>O 51 consumption in soils are less well studied and both atmospheric N<sub>2</sub>O and locally 52 produced N<sub>2</sub>O can be taken up by soils and reduced to N<sub>2</sub> as the last step in the 53 denitrification process, owing to the N<sub>2</sub>O reductase enzyme (named Nos; Chapuis-Lardy 54 et al. 2007). Nitrifiers have also been shown to play a role in  $N_2O$  consumption by 55 reducing NO<sub>2</sub> to N<sub>2</sub>, a process called nitrifier denitrification (Megonigal et al. 2004). 56 Alternative processes of consumption have also been suggested, including aerobic 57 denitrification and assimilatory reduction to ammonia (NH<sub>3</sub>) (Chapuis-Lardy et al. 2007, 58 Vieten et al. 2008).

59 There is a lack of knowledge of the potential for and the controllers of N<sub>2</sub>O 60 consumption in forest soils, though mechanisms have been identified. Few studies have 61 investigated N<sub>2</sub>O consumption directly, yet many focusing on emissions have 62 nevertheless cited negative fluxes (from the atmosphere to the soil), which have often 63 gone unexplained (Chapuis-Lardy et al. 2007). Additionally, even where net emission is 64 observed, consumption processes can exert a significant effect on its magnitude (Arah et 65 al. 1991). Much of the uncertainty leading to the large range in estimated forest soil 66 emissions is related to the possibility of an underestimation of the potential for N<sub>2</sub>O 67 consumption, which could depress estimated emissions (Ullah et al. 2008).

68 In other work complementary to the present study, at two deciduous forest sites in 69 southern Quebec, Mont St. Hilaire (MSH) and Morgan Arboretum (MA), we measured 70 soil N<sub>2</sub>O fluxes along hill slope catenas. While overall N<sub>2</sub>O emission has been observed 71 from the forest soils over the growing season, net consumption of N<sub>2</sub>O was observed 72 from well-drained soils at both sites during several summer sampling dates in 2006, 73 particularly in June and July, though the rates remained small (Unpublished data).  $N_2O$ consumption rates ranged from  $3.1 \pm 1$  to  $6.0 \pm 0.5 \ \mu g \ N_2 O-N \ m^{-2} \ h^{-1}$  in well-drained 74 soils under American beech (Fagus grandifolia) and sugar maple (Acer saccharum) at the 75 MSH and MA sites, ranging from 0.1 to 22  $\mu$ g N<sub>2</sub>O-N m<sup>-2</sup> h<sup>-1</sup> in these soils. Overall, there 76 77 was a net emission of N<sub>2</sub>O to the atmosphere from well-drained soils under American beech and sugar maple, averaging  $3.0 \pm 0.7$  and  $5.4 \pm 0.3 \ \mu g \ N_2 O-N \ m^{-2} \ h^{-1}$ , respectively, 78 79 when N<sub>2</sub>O consumption rates were included in the calculation. When consumption rates 80 were excluded from the calculation, net emissions rates averaged  $5.5 \pm 0.5$  and  $6.5 \pm 0.6$  $\mu$ g N<sub>2</sub>O-N m<sup>-2</sup> h<sup>-1</sup>, respectively (Unpublished data). Chapuis-Lardy *et al.* (2007) cited two 81 82 studies in which N<sub>2</sub>O consumption was observed in temperate deciduous forests, with rates ranging from 0.6 to 66  $\mu$ g N<sub>2</sub>O-N m<sup>-2</sup> h<sup>-1</sup> (Dong *et al.* 1998, Goossens *et al.* 2001). 83 84 This suggests that N<sub>2</sub>O consumption through denitrification is occurring within soils and 85 can have significant impacts on net and average fluxes of N<sub>2</sub>O from the soils. 86 The range of environmental factors influencing N<sub>2</sub>O fluxes from these soils is broad, including both biogeochemical factors, such as NO<sub>3</sub>, NH<sub>4</sub>, and organic C 87 88 availability, as well as physical factors such as soil texture, porosity, moisture, and 89 temperature (Megonigal et al. 2004, Chapuis-Lardy et al. 2007, Ullah et al. 2008). These 90 factors may affect the soil microbial populations, favoring certain functional groups and 91 processes over others (Megonigal et al. 2004), as well as differences among topographic 92 position and forest type (Ullah et al. 2008). Knowledge of the ways in which these 93 factors control  $N_2O$  fluxes in different soils is incomplete, particularly as related to 94 consumption processes (Chapuis-Lardy et al. 2007); in general, factors that limit N<sub>2</sub>O 95 diffusion appear to encourage consumption, including low mineral N levels and high 96 moisture contents, suggesting denitrification as a principal mechanism for this 97 consumption (Bandibas et al. 1994, Megonigal et al. 2004). In addition, the enzyme 98 responsible for N<sub>2</sub>O reduction in denitrifiers (Nos) is known to be particularly sensitive to

99 pH and oxygen (O<sub>2</sub>) (Chapuis-Lardy *et al.* 2007).

100 We hypothesized that a) wet soil conditions and anaerobic processes impose  $N_2O$ 101 consumption in these forest soils through denitrification; b) that soils with low available 102 mineral N as substrate for N<sub>2</sub>O production during denitrification under anoxic conditions 103 may force full reduction and uptake of atmospheric  $N_2O$ ; and c) that high organic C 104 contents in the soil may increase microbially available energy stores and encourage 105 atmospheric N<sub>2</sub>O consumption, when soils undergo NO<sub>3</sub> limitation. To test these 106 hypotheses, we measured potential N<sub>2</sub>O consumption rates in sixteen soil samples 107 collected from 4 soil types [two organic and two mineral] in the MSH and MA sites in 108 laboratory incubations under elevated headspace N<sub>2</sub>O concentrations to establish their 109 potentials for N<sub>2</sub>O consumption and to identify key conditions that favor this 110 consumption.

111

#### 112 MATERIALS AND METHODS

#### 113 Study Sites

114 The two sites used in this study are representative of southern Québec mixed 115 deciduous forest with a combination of tree species: American beech, sugar maple, 116 yellow birch (*Betula alleghaniensis*), striped maple (*Acer pensylvanicum*), white ash 117 (Fraxinus americana) and eastern hemlock (Tsuga canadensis). The Mont St. Hilaire site 118 is located within an old growth forest occurring on one of the Monteregian hills, 119 approximately 30 km east of Montreal. Two types of well-drained soils were sampled at 120 this site: a sandy loam Brunisol underlying beech-dominated stands and a Podzol 121 underlying hemlock-dominated stands, referred to as 'beech' and 'hemlock', respectively. 122 The Morgan Arboretum site occurs in semi-managed forest located on the western tip of 123 the island of Montreal. Sugar maple is the dominant tree species at this site, overlying 124 well-drained, sandy loam soils referred to as 'maple'.

125

# 126 Soil collection and preparation

Four plots (1 m<sup>2</sup> each) were randomly selected for each of the three soils. In November 2007, soil cores (10 cm diameter) were taken randomly from each of four plots to a depth of 10 cm and bulked, transferred to the laboratory and refrigerated until

130 further analysis. While the upper 10 cm of the maple (mineral) and hemlock (organic) 131 soils showed no soil horizon change, the beech soils contained an organic horizon (O-132 horizon) overlying the mineral soil horizon (A-horizon). For these beech cores, the soil 133 was separated into the organic (0 to 5 cm) and mineral (5 to 10 cm) horizons for separate 134 analysis. The four resulting soils used in this study are: beech mineral, beech organic, 135 hemlock organic and maple mineral. Each of the 16 soil samples (4 plots each for 4 soil 136 types) was homogenized manually and sieved (< 2 mm), then analyzed for soil moisture 137 and pH in water and extracted and analyzed for total dissolved N (TDN), dissolved 138 organic C (DOC), NO<sub>3</sub> and NH<sub>4</sub> contents, as described in Ullah *et al.* (2008). Samples 139 collected from 4 sampling points in each plot for N<sub>2</sub>O consumption incubation and soil 140 analysis incorporated the spatial variability within each plot.

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#### 142 Soil Leaching and Pre-incubations

143 Soil leaching and pre-incubations were performed in January 2008 on field moist 144 soil samples collected from the four locations in the two watersheds to create conditions 145 that favor N<sub>2</sub>O consumption hypothesized above, including NO<sub>3</sub> limitation and anoxic 146 conditions, before testing individual hypotheses under 8 treatments. To leach NO<sub>3</sub>, 30 g 147 of a soil at a time was leached by gravity in 60 ml syringes with the plungers removed 148 and a Whatman GF/D filter placed at the tip. Three sequential washes of 15 ml de-ionized 149 water (DI) were passed through the soils before returning them to field moisture 150 conditions by applying pressure with the syringe plungers. Leachates were collected to 151 ensure that soils were returned to field moisture conditions; these leachates were 152 subsequently acidified and analyzed for DOC and TDN contents. After leaching, 15 g of 153 soils was weighed into 150 ml serum bottles and 20 ml of DI was added. The bottles were 154 capped tight, flushed with oxygen-free  $N_2$  gas for 1 hour to induce anoxic conditions and 155 incubated at room temperature for 5 days to further exhaust soil NO<sub>3</sub> through 156 denitrification. Gas samples for N<sub>2</sub>O concentration determination were collected from the 157 headspace of the pre-incubated leached soil samples at 0, 24, 72 and 120 hours for one set 158 of incubations, to test for the expected NO<sub>3</sub>-depletion process occurring in the soils. 159 After leaching and pre-incubation, soils were incubated under 8 treatments, 160 including 6 treatments applied to the leached and pre-incubated soil samples, 1 treatment

to unleached but pre-incubated samples and 1 treatment to unleached and not pre-

162 incubated soil. A brief description and rationale of each treatment follows, with the

163 procedures used:

164 1 Baseline

165 This treatment involved incubation of leached and pre-incubated soils under 166 elevated headspace N<sub>2</sub>O concentrations to investigate if NO<sub>3</sub> limitation and anoxic 167 conditions would result in larger potential N<sub>2</sub>O consumption rates. 15 g of leached and 168 pre-incubated soil was weighed into a 150 ml serum bottle, followed by the addition of 169 20 ml of DI water. The bottles were capped with a gas-tight septa, flushed with oxygen-170 free  $N_2$  gas for 1 hour to induce anoxic condition. After flushing, the headspace  $N_2O$ 171 concentration in the bottles was raised to 2 ppm to ensure unlimited supply of  $N_2O$  and be 172 able to quantify potential  $N_2O$  consumption rates. The incubation was performed on soil 173 slurries on a rotary shaker at 75 rpm to encourage equilibrium solubility of the headspace gases. The incubation period lasted 24 hours, with 5  $cm^3$  gas samples taken at 0, 6, 12 174 175 and 24 hours, stored in pre-evacuated glass vials until analysis for  $N_2O$  concentration on 176 a Shimadzu 14-A gas chromatograph equipped with a 6 m long porapack Q column and 177 an electron capture detector. The end of the the GC column had a 4 valve electronic Valco Valve attached to it, which was timed to vent off separated O2 coming out of 178 179 column to avoid loss in the dector sensitivity for N<sub>2</sub>O detection. Once the O<sub>2</sub> was vented 180 off, the valve swiched back and directed the subsequently separated N<sub>2</sub>O in the column into the detector. The column temperature was adjusted to 60 °C and that of the detector 181 182 to 310 °C. Rate of N<sub>2</sub>O consumption or emission was calculated from the change in 183 concentration over the sampling duration using a linear equation obtained through a 184 calibration curve of known N<sub>2</sub>O standards.

185 2 Glucose-amended

186 This treatment repeats the baseline conditions, with an additional amendment of 187 glucose (approx. 0.8 mg C/g dry soil) to identify the effects of increased C availability, 188 delivered through a needle-fitted syringe to distribute the solution evenly throughout the 189 soil slurry.

190 *3 NO<sub>3</sub>-amended* 

191 This treatment repeats the baseline conditions, with an additional amendment of KNO<sub>3</sub>

- 192 solution (approx. 15 µg NO<sub>3</sub>-N /g dry soil), delivered through a needle-fitted syringe.
- 193 This treatment is a test of whether N<sub>2</sub>O consumption is reduced when inorganic N is
- 194 readily available.
- 195 *4 Unleached*
- 196 This treatment repeats the baseline conditions on soils, which were unleached, but pre-
- 197 incubated to isolate the effects of this method of N-limitation.
- 198 5 No N<sub>2</sub>O amendment
- 199 This treatment repeats the baseline conditions, excluding amendment of bottle
- 200 headspaces with  $N_2O$ , with headspace composed entirely of  $N_2$  gas.
- 201 6 Field moisture, aerobic
- 202 This treatment is designed to remove the gas diffusion limitation imposed in the baseline
- 203 incubation. This was done by incubating the soils at field moisture instead of under slurry
- 204 conditions and under aerobic headspace conditions (but still amended with N<sub>2</sub>O). Soils in
- 205 this treatment were unleached and not pre-incubated.
- 206 7 Unleached, field moisture, no N<sub>2</sub>O amendment
- 207 This treatment represents field conditions, with unleached soils incubated at field
- 208 moisture under aerobic and unamended conditions with ambient air headspace.
- 209 8 Acetylene-amended
- 210 This treatment repeats the baseline conditions but with the additional amendment of 10%
- 211 acetylene in the headspace to inhibit the reduction of N<sub>2</sub>O to N<sub>2</sub> and block nitrification,
- thereby isolating the role of denitrification and confirming that N<sub>2</sub>O consumption
- 213 occurred through denitrification and not nitrification.
- 214

### 215 Statistical analysis

- The gas flux from each bottle was calculated as the slope of the linear regression line best fitting the sample points over 24 hours. Average consumption or emission rates
- and standard errors were calculated based on the four replicates for each soil type and
- treatment. A small constant (the value of the largest emission) was added to each one to
- 220 render all values positive; values were then log-transformed to meet the assumption of
- 221 normality. These values were compared by ANOVA and Fisher LSD using Statistica 6,
- both for differences in fluxes within soil type, by treatment, and among soil types, for

223 each treatment. Pearson correlation coefficients were calculated in SAS 9.1.

224

#### 225 **RESULTS**

The hemlock organic soil sample had the highest DOC content (321  $\mu$ g C g<sup>-1</sup> dry 226 soil), followed by beech organic (180  $\mu$ g C g<sup>-1</sup> dry soil), with beech and maple mineral 227 samples having the lowest DOC content (49 and 41  $\mu$ g C g<sup>-1</sup> dry soil, respectively; Table 228 229 1). The same trend is observed for TDN, with the hemlock organic soil having the highest content (17.8  $\mu$ g N g<sup>-1</sup> dry soil) and maple the lowest (5.7  $\mu$ g N g<sup>-1</sup> dry soil). In terms of 230 NO<sub>3</sub> and NH<sub>4</sub> contents, however, the hemlock organic soil had the lowest values (0.9 and 231 3.1  $\mu$ g N g<sup>-1</sup> dry soil, respectively), with beech mineral (3.7 and 5.9  $\mu$ g N g<sup>-1</sup> dry soil) and 232 organic (3.4 and 5.6 µg N g<sup>-1</sup> dry soil) exhibiting the highest contents of both ions, trailed 233 by maple mineral, which showed slightly lower contents (2.7 and 5.0  $\mu$ g N g<sup>-1</sup> dry soil). 234 Leaching slightly reduced DOC and TDN contents for all soils, though the hemlock 235 organic soil showed a large reduction in DOC content (80  $\mu$ g C g<sup>-1</sup> dry soil) with leaching 236 237 (Table 1).

238  $N_2O$  exchange from pre-incubated soils was low, ranging from a net emission of 0.22 ng N<sub>2</sub>O-N  $g^{-1} d^{-1}$  from the hemlock soil to a net consumption of 1.8 ng N<sub>2</sub>O-N  $g^{-1} d^{-1}$ 239 240 from the beech mineral soil. Pre-incubated leached soils showed significantly smaller 241  $N_2O$  fluxes than those incubated under elevated headspace  $N_2O$  (Fig. 1). Substantial net 242  $N_2O$  consumption was observed in the soils under the conditions hypothesized to 243 facilitate N<sub>2</sub>O reduction: treatment 1 with anoxic conditions and with 2 ppm N<sub>2</sub>O-244 amended headspace (Fig. 1). The greatest potential for N<sub>2</sub>O consumption was found in the hemlock organic soil, with a consumption of 111 ng N<sub>2</sub>O-N  $g^{-1} d^{-1}$  (Fig. 1c), 245 significantly larger (p < 0.05) than the beech organic soil with 74.5 ng N<sub>2</sub>O-N g<sup>-1</sup> d<sup>-1</sup> (Fig. 246 247 1 b). The N<sub>2</sub>O consumption rate was significantly slower in the two mineral soils: beech  $(45.2 \text{ ng } N_2\text{O-N } \text{g}^{-1} \text{d}^{-1})$  and maple  $(40.7 \text{ ng } N_2\text{O-N } \text{g}^{-1} \text{d}^{-1})$ , Fig. 1a, d). 248 When soils were incubated at field moisture or under aerobic conditions, the N<sub>2</sub>O 249 250 consumption rate decreased greatly in all soils, even in the presence of high 251 concentrations of  $N_2O$  in the headspace (treatments 6 and 7, Fig. 1). When the headspace

252 remained unamended with  $N_2O$ , most soils switched to slow rates of  $N_2O$  emission,

253 despite saturated and anoxic conditions (Treatment 5). For the incubation at field

- 254 moisture, under aerobic, unamended conditions, the average fluxes were near-zero for all
- soils (Incubation 7). Leaching with  $H_2O$  appeared to slightly increase the potential for
- 256 N<sub>2</sub>O consumption for most soils, but the average flux was significantly different from
- that of the unleached soils only for the beech mineral soil (Treatment 4, Fig. 1a).
- The addition of organic C in the form of glucose (treatment 2) yielded a slightly faster N<sub>2</sub>O consumption rate in the two beech soils, but was statistically indistinguishable from the baseline treatment 1 for all soils (Fig. 1). The addition of NO<sub>3</sub> solution to the soils clearly decreased the N<sub>2</sub>O consumption rate for all soils, though not significantly (treatment 3). The addition of headspace acetylene (treatment 8) resulted in near-zero N<sub>2</sub>O fluxes in all soils, canceling the strong consumption observed in treatment 1 and was statistically significant (p < 0.05) in all cases.
- There were few significant differences in N<sub>2</sub>O fluxes among the four soils for each treatment: significant (p < 0.05) differences occurred only in treatments 1, 2 and 4, with the hemlock and beech organic soils generally with the largest potential consumption rates (Fig. 1).
- A correlation analysis revealed that the potential  $N_2O$  consumption rates under treatment 1 are significantly and negatively correlated with both DOC and original soil moisture, suggesting that soils with higher C contents and field moisture levels have a greater capacity to consume  $N_2O$  (Table 2). In addition, TDN showed a similar negative correlation with  $N_2O$  fluxes, though only significant at the 10% level.
- 274

#### 275 **DISCUSSION**

276 The strong potential  $N_2O$  consumption rates obtained from the baseline incubation 277 reveal that these well-drained forest soils have a significant capacity for N<sub>2</sub>O reduction 278 under conditions of anoxia and N limitation. Though conditions in this laboratory study 279 are incomparable in many ways to field conditions, the occurrence of potential  $N_2O$ 280 consumption rates at such magnitude suggests that these processes could play a 281 significant role in determining net fluxes of N<sub>2</sub>O from the soils, even though these soils 282 are weak net sources of N<sub>2</sub>O throughout most of the growing season. This result suggest a 283 need for in situ N<sub>2</sub>O consumption studies and the inclusion of N<sub>2</sub>O reduction processes in 284 the consideration of N cycling and gas fluxes in these soils, and forest soils in general.

285 The conditions which facilitated N<sub>2</sub>O reduction in this study included: a) the 286 imposition of enhanced  $N_2O$  consumption, through anaerobiosis and soil water 287 saturation; b) low availability of electron acceptors, notably NO<sub>3</sub>, achieved by leaching 288 and pre-incubating soils; and c) high organic C contents to encourage reduction of  $N_2O$  as 289 an alternate electron acceptor. The baseline incubation (treatment 1) showed that the 290 combination of these factors yielded significant consumption in all four soils, both 291 mineral and organic. The near-zero N<sub>2</sub>O fluxes obtained under conditions similar to those 292 in the field (treatment 7) represented a clear contrast to the baseline incubation. The 293 remaining six treatments served to identify the effect of the different variables on 294 potential N<sub>2</sub>O consumption.

295 The glucose addition (treatment 2) increased N<sub>2</sub>O consumption levels only (and 296 then not statistically significant) in the beech soils, both organic and mineral. Soil C:N 297 ratios (Table 1) influenced  $N_2O$  consumption, where beech and hemlock showed faster 298 consumption rates and their C:N ratios were larger than those of soils under sugar maple. 299 Cavigelli and Robertson (2001) also noted that the Nos enzyme is particularly sensitive to 300 a low C:N ratio and stated that the organic C level in a soil is an important factor for soil 301 denitrifier populations. This might explain the low baseline  $N_2O$  consumption of the 302 maple mineral soil, as well as the absence of an increase in consumption upon glucose 303 amendment. The hemlock organic soil may not have shown increased N<sub>2</sub>O consumption 304 upon glucose addition, owing to the high DOC levels of organic C in these soils. Larger 305 nitrification rates in soils under the sugar maple compared to those under American beech 306 and hemlock in these plots may have led to the evolution of denitrifiers with low affinity 307 for  $N_2O$  consumption in soils under sugar maple trees (Ullah and Moore, 2009). 308 Additionally, in situ N<sub>2</sub>O consumption rates in soils under American beech were 2 times 309 larger than under sugar maple in these sites (Ullah and Moore, in prep). We hypothesize 310 that soils with larger soil C content and C:N ratio in deciduous forests consume more 311  $N_2O$  than soils under smaller C:N ratios. We suggest further microbial studies to validate 312 this hypothesis under field conditions. 313 Our strong correlation between N<sub>2</sub>O consumption and extractable DOC

314 concentration, within the limited range of soils, is consistent with reports on the

315 importance of organic C (as well as moisture, oxygen, disturbance and pH levels) in

determining N<sub>2</sub>O consumption potentials, which may also reflect differences in microbial communities among soil types (Parkin 1987, Cavigelli and Robertson 2001, Wallenstein *et al.* 2006). Given the limited, non-significant increases in N<sub>2</sub>O consumption upon glucose amendment in the beech soils, further experiments are needed to determine whether different and larger organic C amendments would have a significant effect, or whether the microbial populations present in the soils limit the response over the 24-hour incubation period.

323 The addition of  $NO_3$  (treatment 3) and the incubation of unleached soils 324 (treatment 4) were performed to contrast  $N_2O$  consumption with those of the baseline 325 incubation (treatment 1), where an effort was made to eliminate as much  $NO_3$  from the 326 soils as possible. This was to test whether denitrifiers will turn to an alternate, though less 327 energetically favorable electron acceptor, the  $N_2O$  provided through headspace 328 amendment (Bandibas et al. 1994), a technique of limiting the availability of electron 329 acceptors successfully employed in other studies (e.g. Firestone et al. 1980, Holtan-330 Hartwig *et al.* 2000). This hypothesis was supported by our results, showing a decreasing 331 trend, although not statistically significant at p < 0.05, in treatments 3 and 4 (Figure 1). 332 The near-zero N<sub>2</sub>O fluxes in pre-incubation conditions support the hypothesis that this 333 treatment lowered NO<sub>3</sub> in the soil, allowing the uptake and reduction of  $N_2O$  as an 334 alternate electron acceptor in the following treatments. Leaching the soils clearly 335 depressed total N levels in all cases (Table 1). The differences in N<sub>2</sub>O consumption 336 between the leached, pre-incubated soils and the same soils with  $NO_3$  added, reflect the 337 control of NO<sub>3</sub>-availability: adding NO<sub>3</sub> after leaching and pre-incubating the soils 338 effectively cancels out the effect of the initial leaching. There was a decrease in  $N_2O$ 339 consumption by about one third upon NO3 addition in the beech soils, whereas the effect 340 was less pronounced in the hemlock and maple soils.

Thus, while NO<sub>3</sub> availability is significant, other conditions such as anaerobiosis,
moisture and N<sub>2</sub>O play key roles in determining whether N<sub>2</sub>O consumption will occur.
The correlation between low NO<sub>3</sub> levels and N<sub>2</sub>O consumption is widely cited, but most
studies go on to describe the conditions of anaerobiosis and saturation as predominantly
important (Blackmer and Bremner 1976, Firestone *et al.* 1980, Bandibas *et al.* 1994,
Holtan-Hartwig *et al.* 2000, Rosenkranz *et al.* 2006, Wallenstein *et al.* 2006). A possible

reason why a major decrease in  $N_2O$  consumption was not observed in treatments 3 and 4 is that the  $NO_3$  was quickly exhausted, leading to a  $NO_3$ -limitation similar to that of the baseline and creating larger  $N_2O$  consumption later during the 24-hour incubation. Firestone *et al.* (1980) suggested that Nos enzyme could be sequentially produced during the incubation, resulting first in an increase and then a decrease in headspace  $N_2O$ concentration. We recommend further studies with <sup>15</sup>N tracers to validate this hypothesis.

353 Excess N<sub>2</sub>O appears to be critical in its consumption as an alternate electron 354 receptor upon  $NO_3$ -limitation (Mei *et al.* 2004). This is reflected in treatment 5, without 355 the  $N_2O$  headspace amendment, where, for all but the hemlock soil, there was a small 356  $N_2O$  production (Fig. 1) indicating that, all other conditions being equal, when  $N_2O$  is not 357 abundantly available in the soil pore spaces, N<sub>2</sub>O consumption will likely be small, or is 358 severely limited. The leaching and pre-incubation is effective at reducing available  $NO_3$ 359 as a substrate for denitrification but the small rates of  $N_2O$  indicate that complete 360 reduction to  $N_2$  (and thus  $N_2O$  consumption) may be limited due to lower soil pore space 361 N<sub>2</sub>O concentrations. Indeed, the lack of substrates for denitrifiers under treatment 5 likely 362 limited their activity in either the production or consumption, resulting in the fluxes. For 363 the hemlock soil, the large standard error makes the interpretation of the small 364 consumption rates for this incubation ambiguous, but perhaps suggests that complete 365 reduction can occur under these circumstances, if enough organic C is available to 366 encourage microbial activity, but this is variable and ephemeral. Fast denitrification rates 367 are often associated with high organic C in soils (Parkin 1987, Wrage et al. 2001) and more work could be done to establish the effect of varying the headspace N2O 368 369 concentrations, from 2 ppm (treatment 1) to ambient.

370 In treatment 6, soils were incubated aerobically and at field moisture to identify 371 the effect of anoxic conditions, resulting in a significant decrease in  $N_2O$  consumption. 372 Anoxic conditions play a critical role in  $N_2O$  consumption in soils, for which there is 373 support in the literature:

a) When oxygen is allowed to diffuse into the soil, denitrifier activity and Nos activity in

375 particular are impeded and restricted to microsites of anoxia (Firestone *et al.* 1979,

376 Bandibas et al. 1994, Cavigelli and Robertson 2001).

b) Low moisture allows greater gas diffusion into and out of the soil profile and any

remaining NO<sub>3</sub> reduced to N<sub>2</sub>O is able to diffuse out of the soil without further reduction to N<sub>2</sub>, whereas higher moisture content facilitates N<sub>2</sub>O entrapment and reduction to N<sub>2</sub> (Clough *et al.* 2005; Ullah et al. 2005).

c)  $N_2O$  present in the headspace diffuses into the soil profile more easily under low moisture conditions, which paradoxically may tend to encourage consumption in the drier soils by increasing the availability of the  $N_2O$  as a substrate in the redox chain (Bandibas *et al.* 2004). This could be the reason why  $N_2O$  fluxes remained negative, though reduced in magnitude, in treatment 6. This effect is evidenced further in treatment 7, where in addition to the removal of the gas diffusion limitation, the  $N_2O$  headspace amendment is removed (as well as leaching), and fluxes diminish to near-zero or slight production.

388 Clough et al. (2005) suggest that this is due to the decreased time in which potential

389 reduction can occur under low moisture conditions because of increased gas diffusion and

390 prevalence of oxic conditions in soil profile.

391 d) Though the presence of anoxic conditions appears to exert a stronger control over  $N_2O$ 

392 consumption than NO<sub>3</sub> availability, the NO<sub>3</sub>-limitation is likely required initially to

393 encourage N<sub>2</sub>O consumption as the predominant denitrifier activity in anoxic, saturated

394 conditions, as suggested in treatments 3 and 4. The two controls are interdependent, and

their importance can both be traced to the availability of the various reactants in the

396 denitrification redox chain, which in all cases is concentration and diffusion-dependent.

397 The addition of acetylene in treatment 8 inhibited the reduction of N<sub>2</sub>O to N<sub>2</sub> and 398 removed the strong  $N_2O$  consumption under treatment 1 (Schuster and Conrad 1992). 399 This eliminates the possibility that other factors, such as simple diffusion into soil water, 400 is causing the decrease in concentration of  $N_2O$  in the headspace over the incubation 401 period, confirming that the N<sub>2</sub>O is in fact being reduced to N<sub>2</sub> through denitrification. 402 Denitrification is clearly implicated as the process by which this reduction occurs under 403 baseline conditions since the acetylene amendment also inhibits nitrification processes. 404  $N_2O$  is not accumulating in the headspace, implying that the  $N_2O$  being reduced in 405 treatment 1 was primarily amended headspace N<sub>2</sub>O through denitrification. 406 These well-drained soils are capable of consuming  $N_2O$ , so the summer field

407 consumption that first motivated this study are likely not anomalous. Although great care 408 needs to be taken in extrapolating these results to field conditions, the potential for  $N_2O$ 

409 consumption in the field likely results from denitrification processes occurring in isolated

410 anoxic, wet microsites within the soil, where locally produced  $N_2O$  can be retained and

411 reduced to N<sub>2</sub>. This N<sub>2</sub>O consumption is usually not detectable in the field as emission

412 rates are slow emissions, though the emission rate may be decreased by  $N_2O$ 

413 consumption (Ullah *et al.* in prep.). Net field N<sub>2</sub>O consumption could be anticipated after

- 414 heavy rainfall, where soils are strongly NO<sub>3</sub>-limited and with large organic C contents
- 415 (Seitzinger *et al.* 2006).

416

# 417 CONCLUSIONS

418 Well-drained forest soils in southern Québec exhibited potential N<sub>2</sub>O

- 419 consumption through denitrification when incubated under anoxic and saturated moisture
- 420 conditions, and when amended with high levels of atmospheric  $N_2O$ . Mineral N
- 421 limitation within the soils likely stimulated the reduction of N<sub>2</sub>O to N<sub>2</sub>. Organic soils
- 422 showed generally greater N<sub>2</sub>O consumption potentials than mineral soils. Conditions
- 423 favoring N<sub>2</sub>O consumption may occur in wet, anaerobic microsites within the soil profile,
- 424 and such consumption processes could bear significantly on the net flux of  $N_2O$  from
- 425 these soils. Our results also suggest that soil with larger soil C:N ratios exhibiting lower
- 426 nitrification rates may possess higher affinity for N<sub>2</sub>O consumption than soils with
- 427 smaller ratios.

428

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436

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# 516 **TABLES**

Variables	Beech mineral	Beech organic	Hemlock organic	Maple mineral
pН	$4.7\pm0.5$	$4.7\pm0.5$	4.5 ± 0.39	$5.5\pm0.03$
Bulk density (g cm <sup>-3</sup> )	$0.99 \pm 0.21$	$0.56\pm0.10$	$0.48\pm0.1$	$1.19\pm0.04$
DOC ( $\mu$ g C g <sup>-1</sup> dry soil)	$49\pm2$	$180 \pm 18$	321 ± 29	$41 \pm 2$
DOC leached ( $\mu g C g^{-1} dry soil$ )	$45 \pm 2$	$150\pm14$	$241\pm18$	$36\pm2$
TDN (µg N g <sup>-1</sup> dry soil)	$8.5 \pm 2.4$	$15.6\pm2.3$	$17.7\pm0.9$	5.7 ± 2.3
TDN leached ( $\mu g N g^{-1} dry soil$ )	$8 \pm 2.3$	$12.1 \pm 1.1$	$14.4\pm0.7$	$4.8 \pm 1.3$
$NO_3$ (µg N g <sup>-1</sup> dry soil)	3.7	3.4	0.9	2.7
$NH_4$ (µg N g <sup>-1</sup> dry soil)	5.9	5.6	3.1	5
Soil C:N ratio (0-10 cm depth)	$26 \pm 1.3^{*}$		$25 \pm 1.5$	$16 \pm 0.7$
Leaf litter N input (g m <sup>-2</sup> )	$4.6 \pm 1.4^{**}$		$1.9\pm0.3$	$4.9\pm0.1$
Leaf litter fall C:N ratio	61	± 6**	71 ± 3	$52\pm0$

517 Table 1. Chemical properties of the 4 soil types and leaf litter input and litter C:N ratio.

<sup>518</sup> 

 <sup>\*</sup> C:N ratio in soils under beech trees represent an average of both organic and mineral layer as only a 0-10 cm depth sample was
 taken for this purpose. \*\* Represents total litter N input on the soil surface and its C:N ratio.

521 Table 2. Pearson correlation coefficient (*r*) between the properties of the four soils (see

522 Table 1) and the N<sub>2</sub>O consumption rate under treatment 1. Coefficients with *p*-values that

- 523 are significant at the 5% level are listed in **bold** text and those at the 10% level are
- *italicized*.

	N <sub>2</sub> O flux	DOC	TDN	NO <sub>3</sub>	NH <sub>4</sub>	pН	Bulk density
N <sub>2</sub> O flux	-						
DOC	-1.00	-					
TDN	-0.92	0.94	-				
NO <sub>3</sub>	0.82	-0.78	-0.52	-			
$\mathbf{NH}_4$	0.83	-0.80	-0.55	1.00	-		
pН	0.63	-0.67	-0.85	0.10	0.13	-	
Bulk density	0.71	-0.73	-0.83	0.30	0.34	0.94	-
Field soil moisture	-0.97	0.98	0.97	-0.68	-0.70	-0.79	-0.85

# 528 FIGURES

529 Figure 1. Average N<sub>2</sub>O exchange rates (± standard error) for the four soil samples under

- 530 the Pre-incubation and the 8 treatments: 1 Baseline; 2 Glucose-amended; 3 NO<sub>3</sub>-
- 531 amended; 4 Unleached; 5 No N<sub>2</sub>O amendment; 6 Field moisture, aerobic; 7 Unleached,
- 532 field moisture, no N<sub>2</sub>O amendment; 8 Acetylene-amended. Negative values indicate
- 533 consumption by the soil. Statistically significant differences (p < 0.05) among treatments
- 534 for each soil for are indicated by lower case lettering and among the four soils under
- treatments 1, 2, 3 and 4 are indicated by bold upper case lettering.

