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The inhibitory effect of difluoromethane on  $CH_4$  oxidation in reconstructed peat columns and side-effects on  $CO_2$  and  $N_2O$  emissions at two water levels

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1 **Research paper:**

2 **The inhibitory effect of difluoromethane on CH<sub>4</sub> oxidation in**  
3 **reconstructed peat columns and side-effects on CO<sub>2</sub> and N<sub>2</sub>O emissions**  
4 **at two water levels**

5 S. Vicca<sup>1\*</sup>, H. Flessa<sup>2</sup>, N. Loftfield<sup>2</sup>, I.A. Janssens<sup>1</sup>

6

7 <sup>1</sup> Research Group of Plant and Vegetation Ecology, Department of Biology, University of  
8 Antwerp, Universiteitsplein 1, B-2610 Wilrijk, Belgium

9 <sup>2</sup> Institute of Soil Science and Forest Nutrition, University of Göttingen, Büsingenweg 2, D-37073  
10 Göttingen, Germany

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15 Corresponding author: Sara Vicca

16 Email: [Sara.Vicca@ua.ac.be](mailto:Sara.Vicca@ua.ac.be)

17 Telephone: +32 3 820 22 82

18 Fax: +32 3 820 22 72

19

20 **Abstract**

21 Methane emissions from soils are the net result of two processes: methane ( $\text{CH}_4$ )  
22 production and  $\text{CH}_4$  oxidation. In order to understand how both processes respond to  
23 environmental changes, it is necessary to distinguish between  $\text{CH}_4$  production and  
24 oxidation. In bacterial cultures and small soil samples, difluoromethane ( $\text{CH}_2\text{F}_2$ ) was  
25 found to inhibit  $\text{CH}_4$  oxidation reversibly, without affecting  $\text{CH}_4$  production. Hence,  
26  $\text{CH}_2\text{F}_2$  allows the study of  $\text{CH}_4$  production directly and of  $\text{CH}_4$  oxidation indirectly. To  
27 our knowledge, however, the inhibitory effect of  $\text{CH}_2\text{F}_2$  within soil columns has not yet  
28 been evaluated. We therefore tested which  $\text{CH}_2\text{F}_2$  concentration is needed for complete  
29 inhibition of  $\text{CH}_4$  oxidation in reconstructed 28 cm high peat soil columns under different  
30 water levels (WL). We found that soil columns require considerably higher headspace  
31  $\text{CH}_2\text{F}_2$  concentrations for complete inhibition of  $\text{CH}_4$  oxidation than small soil samples.  
32 Inhibition remained complete until ca. 24 h after  $\text{CH}_2\text{F}_2$  exposure. Then, the inhibitory  
33 effect diminished. The time needed for the inhibitory effect to disappear depended on  
34 WL; at a low WL of -15 cm, the inhibitory effect declined slowly and oxidation rates  
35 recovered by 90% only after 12 days. At WL = -5 cm,  $\text{CH}_4$  oxidation recovered much  
36 faster (90% recovery after ca. three days). Last,  $\text{CH}_2\text{F}_2$  addition significantly decreased  
37 the  $\text{N}_2\text{O}$  emissions, whereas  $\text{CO}_2$  emissions remained unaltered.

38

39 *Keywords:* Difluoromethane; Inhibitor; Methane; Carbon Dioxide; Nitrous oxide

## 40 **1. Introduction**

41

42 Methane (CH<sub>4</sub>), the second most important greenhouse gas (IPCC, 2001), is  
43 produced in soils when organic matter is degraded anaerobically by methanogenic  
44 bacteria (Oremland, 1988; Conrad, 1989). In aerobic soils or aerobic surface layers on top  
45 of anaerobic soils, methanotrophic bacteria can oxidize CH<sub>4</sub> to CO<sub>2</sub> (Cicerone and  
46 Oremland, 1988). The net CH<sub>4</sub> flux between soil and atmosphere thus results from two  
47 opposite processes. These processes often respond differently to environmental changes.  
48 Water level drawdown, for example, decreases the production of CH<sub>4</sub>, whereas it can  
49 enhance CH<sub>4</sub> oxidation if CH<sub>4</sub> remains readily available (Jungkunst and Fiedler, 2007). In  
50 order to understand how climate and other environmental factors control methane  
51 emissions, and to accurately incorporate these responses into models, it is essential to  
52 study the separate responses of both production and oxidation of CH<sub>4</sub>.

53 Methane production and oxidation can be differentiated by using selective  
54 inhibitors. Several inhibitors have been used to inhibit CH<sub>4</sub> oxidation, e.g., acetylene and  
55 methyl fluoride, but most of them resulted in an irreversible inhibition or affected not  
56 only CH<sub>4</sub> oxidation, but also CH<sub>4</sub> production (Frenzel and Bosse, 1996; Janssen and  
57 Frenzel, 1997; Matheson et al., 1997). Matheson et al. (1997) and Miller et al. (1998),  
58 however, discovered that difluoromethane (CH<sub>2</sub>F<sub>2</sub>) can inhibit CH<sub>4</sub> oxidation without  
59 affecting CH<sub>4</sub> production. Moreover, they reported that the inhibition is completely  
60 reversible and that CH<sub>2</sub>F<sub>2</sub> can be quickly removed from the system as it is very soluble in  
61 water. Difluoromethane was used in several experiments on small soil samples (e.g., Teh  
62 et al., 2005; 2006), but to our knowledge, a thorough evaluation of the inhibitory effect of

63 CH<sub>2</sub>F<sub>2</sub> in intact soil columns has not been reported before (although CH<sub>2</sub>F<sub>2</sub> was applied  
64 in a few in situ studies; Krüger et al., 2001, 2002 and Shrestha et al., 2008).

65 Miller et al. (1998) used bacterial cultures and 5 g dry soil in serum bottles for  
66 their experiments, and concluded that a headspace concentration of about 0.03% CH<sub>2</sub>F<sub>2</sub>  
67 was sufficient to inhibit the CH<sub>4</sub> oxidation completely. However, we expected that higher  
68 concentrations would be needed when complete soil columns are investigated, because  
69 the inhibitor has to diffuse into the soil to reach the sites of CH<sub>4</sub> oxidation. Furthermore,  
70 the depth of WL is expected to affect the inhibitory effect, because dilution of the CH<sub>2</sub>F<sub>2</sub>  
71 concentration in the aerobic soil is expected to increase with increasing WL depth. In an  
72 incubation experiment, we therefore exposed peat soil cores to two different water levels  
73 and treated these soil cores with different CH<sub>2</sub>F<sub>2</sub> concentrations. Our objectives were (1)  
74 to verify which CH<sub>2</sub>F<sub>2</sub> concentration is needed to inhibit the CH<sub>4</sub> oxidation completely at  
75 the two water levels and (2) to assess whether CO<sub>2</sub> and N<sub>2</sub>O emissions were affected by  
76 CH<sub>2</sub>F<sub>2</sub> concentrations needed to inhibit CH<sub>4</sub> oxidation.

77

78 **2. Materials and methods**

79

80 2.1. *Soil collection*

81 In September 2006, we collected peat soil from a fen in nature reserve 'Het Wik'  
82 (Genk, Belgium; 50° 57' N, 5° 25' E). The vegetation on this soil was dominated by *Erica*  
83 *cinerea*, *Pieris* sp. and *Sphagnum* sp. Other species occurring were *Eriophorum*  
84 *angustifolium*, *Drosera* sp. and *Typha latifolia*. After peat collection, all vegetation was  
85 removed from the soil and, in order to keep the variation among soil samples as small as  
86 possible, we homogenized the peat by hand and filled it in PVC columns (inner diameter  
87 10 cm) to a height of 28 cm. In 12 columns with homogenized peat the water level (WL)  
88 was set at 5 cm below the soil surface (high water level; WL = -5 cm). In 12 other soil  
89 columns, we maintained a WL of 15 cm below the soil surface (low water level; WL = -  
90 15 cm). In order to control the water level in the soil columns, each column (of which the  
91 lower 5 cm was perforated) was placed in a water bath to obtain communicating water  
92 levels between soil column and water bath. Water levels in the soil columns were  
93 controlled via the water level in the water bath.

94

95 2.2. *Gas flux measurements*

96 The columns were closed by PVC lids (diam. 10 cm; height 10 cm) and placed in  
97 automated incubators, set at 17 °C. A fresh air supply unit continuously flushed the  
98 headspaces with a controlled fresh air flow of 10.0 ml min<sup>-1</sup>. Net CO<sub>2</sub>, CH<sub>4</sub> and N<sub>2</sub>O  
99 fluxes were measured with a GC that contained a flame ionisation detector for CH<sub>4</sub> and a  
100 <sup>63</sup>Ni electron capture detector for CO<sub>2</sub> and N<sub>2</sub>O analysis (Shimadzu, Duisburg, Germany;

101 for details see Loftfield et al., 1997 and Flessa and Beese, 2000). Every three hours, the  
102 input air and the exhaust air of each soil column were analyzed for CO<sub>2</sub>, CH<sub>4</sub> and N<sub>2</sub>O.  
103 Multi-component calibration gases covering the concentrations of 300 to 3000 µl l<sup>-1</sup> CO<sub>2</sub>,  
104 1 to 200 µl l<sup>-1</sup> CH<sub>4</sub> and 0.35 to 2 µl l<sup>-1</sup> N<sub>2</sub>O were measured at the same regular intervals.  
105 Flux rates were calculated from the airflow rate through the microcosm headspace and  
106 the difference in the CO<sub>2</sub>, CH<sub>4</sub>, and N<sub>2</sub>O concentration between the input air and the  
107 exhaust air.

108

### 109 2.3. *Experimental set-up*

110 The inhibition experiment started after the greenhouse gas emissions reached a  
111 stable flux rate (after three months). In order to verify whether CH<sub>2</sub>F<sub>2</sub> inhibited the CH<sub>4</sub>  
112 oxidation, we injected different amounts of CH<sub>2</sub>F<sub>2</sub> (Spectra Gases, Germany) into the  
113 headspace of the soil columns (via tubes in the headspace). Subsequently, we closed the  
114 headspaces and stopped the airflow for 2 h (no measurements were made during these 2  
115 h). Then, the fresh air supply through the column headspaces was resumed. Due to a  
116 technical problem, we have no results of the third inhibition treatment in the high water  
117 level columns. To remove the dissolved CH<sub>2</sub>F<sub>2</sub> from the system, water baths were  
118 completely refreshed between the sequential additions of different CH<sub>2</sub>F<sub>2</sub> concentrations.  
119 Prior to injecting the next CH<sub>2</sub>F<sub>2</sub> concentration, care was taken that the fluxes had  
120 readjusted to their initial values. The CH<sub>2</sub>F<sub>2</sub> concentration sequence that was applied is  
121 given in Table 1.

122 After these inhibition treatments, we tested whether the inhibition of the CH<sub>4</sub>  
123 oxidation by CH<sub>2</sub>F<sub>2</sub> was complete. To this end, we added <sup>13</sup>CH<sub>4</sub> (99 at% <sup>13</sup>C, Spectra

124 Gases, Germany) to the headspaces and checked whether this was oxidized to  $^{13}\text{CO}_2$ . In  
125 detail, we used the following methodology. We first sampled air from the headspaces  
126 four times within one hour after closing the headspaces. Gas samples were collected in 12  
127 ml vacuum vials and were analyzed for  $\delta^{13}\text{C-CO}_2$  by GC-IRMS (Finnigan MAT, Delta  
128 plus; reference standard: PDB). After shortly opening the headspaces for equilibration  
129 with the atmosphere, we closed the headspaces again and injected 6.4 ml of pure  $\text{CH}_2\text{F}_2$   
130 to achieve a headspace  $\text{CH}_2\text{F}_2$  concentration of 1% [vol/vol] (the lowest concentration  
131 that significantly affected the  $\text{CH}_4$  emissions; see results section) in half of the columns  
132 (six per WL; randomly selected). We also stopped the airflow for 2 h, the time needed to  
133 ensure complete inhibition at 1% [vol/vol]  $\text{CH}_2\text{F}_2$ . Subsequently, for each WL, we  
134 injected 5.0 ml  $^{13}\text{CH}_4$  in three inhibited samples and in three samples without  $\text{CH}_2\text{F}_2$   
135 addition, to obtain a headspace concentration of 0.6% [vol/vol]  $^{13}\text{CH}_4$  and then sampled  
136 air for  $^{13}\text{CO}_2$  analysis (three samples within 1 h after  $^{13}\text{CH}_4$  injection). With the  $\delta^{13}\text{CO}_2$   
137 data, we determined the  $\delta^{13}\text{C}$  of the  $\text{CO}_2$  produced in each soil column as the y axis  
138 intercept of a Keeling plot. We are aware that the number of data points was small for  
139 constructing a Keeling plot, increasing the uncertainty of the  $\delta^{13}\text{CO}_2$  to several per mil.  
140 However, our goal was not to study the  $\delta^{13}\text{CO}_2$  of the decomposition or to determine  
141 natural isotope abundance, but to find an indication for  $\text{CH}_4$  oxidation. If  $\text{CH}_4$  was  
142 oxidized, an increase of  $\delta^{13}\text{CO}_2$  in the order of hundreds of per mil of  $\delta^{13}\text{CO}_2$  would be  
143 observed after  $^{13}\text{CH}_4$  injection. Hence, the increased uncertainty in the Keeling plots was  
144 irrelevant for this study. The substantially higher  $\delta^{13}\text{CO}_2$  in the plots treated with  $^{13}\text{CH}_4$  in  
145 the absence of  $\text{CH}_2\text{F}_2$  (see below) demonstrates that our goal was achieved.

146

147 2.4. *Data analysis*

148 In order to verify whether CH<sub>2</sub>F<sub>2</sub> inhibited CH<sub>4</sub> oxidation, we measured CH<sub>4</sub>  
149 emissions before and after CH<sub>2</sub>F<sub>2</sub> injection. For quantification of the inhibitory effect on  
150 the CH<sub>4</sub> emissions, we opted to exclude the CH<sub>4</sub> release via ebullition. Methane emitted  
151 via ebullition is only partially affected by CH<sub>4</sub> oxidation, because most of the CH<sub>4</sub>  
152 transported via bubbles quickly bypasses the sites of CH<sub>4</sub> oxidation. To exclude CH<sub>4</sub>  
153 emissions via ebullition, we considered only the basal CH<sub>4</sub> flux. To this end, we  
154 computed the median flux (unlike the average, the median is less affected by CH<sub>4</sub>  
155 emissions via ebullition; see Fig. 1 for more details on this issue) over the last five days  
156 before the inhibition and the median flux of 24 h following the injection of CH<sub>2</sub>F<sub>2</sub> (after  
157 24 h, the inhibitory effect on the CH<sub>4</sub> emissions started to decline). The inhibitory effect  
158 was calculated as the difference between the median flux rate before and after inhibition.  
159 The average inhibitory effect presented in Fig. 2, 3 and 4 was computed over all dates on  
160 which an inhibition treatment was applied (e.g., the inhibitory effect at 1% [vol/vol]  
161 CH<sub>2</sub>F<sub>2</sub> represents the average over the T3 replicates inhibited on day 150 and the T2  
162 replicates inhibited on days 194 and 289; see Table 1).

163 In order to ensure that the use of the median flux rate did not confound our results,  
164 we determined the median absolute deviation (MAD) before and after each inhibition. As  
165 expected, MAD was somewhat higher for CH<sub>4</sub> emissions at high WL than at low WL  
166 (one-way ANOVA for CH<sub>4</sub> emissions over the five days before the first inhibition: P =  
167 0.08; Table 1), which reflects the higher abundance of ebullition events at WL = -5 cm.  
168 More importantly, however, we found similar MADs before and after the inhibitions  
169 (paired t-test: P = 0.19). Hence, CH<sub>2</sub>F<sub>2</sub> addition did not enhance bubble ebullition (which

170 would increase MAD) and therefore, we are confident that the median flux was an  
171 appropriate measure to determine the inhibitory effect on CH<sub>4</sub> oxidation.

172 For determining the recovery of the CH<sub>4</sub> oxidation after inhibition, we calculated  
173 for each soil column a 'running median' flux over clusters of ten consecutive data points  
174 (i.e., a first median flux calculated from data point one to data point ten, a second median  
175 flux calculated from data point two to data point 11, etc.). In this way, we were able to  
176 reduce the influence of CH<sub>4</sub> release via ebullition. Subsequently, we determined the time  
177 span between CH<sub>2</sub>F<sub>2</sub> addition and the time at which the CH<sub>4</sub> fluxes showed 75% and 90%  
178 recovery. Note that CH<sub>4</sub> fluxes were similar at the beginning and end of the experiment  
179 (May 2007 and November 2007). Emissions of CO<sub>2</sub> were stable throughout the  
180 experiment as well, whereas N<sub>2</sub>O emissions, in particular at the low WL, increased over  
181 the course of the experiment (data not shown).

182 The second objective was to verify whether CH<sub>2</sub>F<sub>2</sub> affected CO<sub>2</sub> and N<sub>2</sub>O  
183 emissions. Similar to the CH<sub>4</sub> emissions, we calculated the average difference between  
184 the median CO<sub>2</sub> and N<sub>2</sub>O flux before and after the inhibition. For the CO<sub>2</sub> emissions, we  
185 used the same time span as for the CH<sub>4</sub> fluxes, i.e., five days before the inhibition and one  
186 day after the inhibition. For the N<sub>2</sub>O emissions, we opted for a longer time span after the  
187 inhibition, because for high CH<sub>2</sub>F<sub>2</sub> concentrations, we observed an inhibitory effect that  
188 lasted one week after addition of the inhibitor. Therefore, we computed the average N<sub>2</sub>O  
189 flux after CH<sub>2</sub>F<sub>2</sub> injection over the five days following the inhibition.

190 We performed all analyses in Matlab (7.2.0.232, The Mathworks, US). Effects of  
191 WL and CH<sub>2</sub>F<sub>2</sub> addition on CH<sub>4</sub>, CO<sub>2</sub> and N<sub>2</sub>O emissions were tested using ANOVA.  
192 Differences are reported significant at P < 0.10.

193 **3. Results**

194

195 *3.1. Emissions and inhibitory effects*

196 As expected, CH<sub>4</sub> emissions were higher at WL = -5 cm than at WL = -15 cm  
197 (Table 2). Note that Table 2 presents the median fluxes. The CH<sub>4</sub> emissions in this table  
198 thus approximate the CH<sub>4</sub> emission via diffusion. Average CH<sub>4</sub> emissions (including CH<sub>4</sub>  
199 release via ebullition) were 0.08 (± 0.01) and 0.03 (± 0.01) μmol CH<sub>4</sub> m<sup>-2</sup> s<sup>-1</sup>, for WL = -5  
200 and WL = -15 cm, respectively. These emissions were well within the range observed for  
201 undrained fens in temperate regions (Jungkunst and Fiedler, 2007). Average CO<sub>2</sub> and  
202 N<sub>2</sub>O emissions were nearly identical to the median fluxes (data not shown) and both  
203 gases exhibited the highest emissions at WL = -15 cm (Table 2).

204 Addition of 0.2% [vol/vol] CH<sub>2</sub>F<sub>2</sub> enhanced the CH<sub>4</sub> emissions, suggesting that  
205 CH<sub>4</sub> oxidation was being inhibited, but the increase was not statistically significant at P <  
206 0.10, for either water level (Fig. 2). A CH<sub>2</sub>F<sub>2</sub> concentration of 1% [vol/vol] or higher  
207 always resulted in a significant inhibitory effect (i.e., an increase of the CH<sub>4</sub> emission),  
208 except for 2% [vol/vol] CH<sub>2</sub>F<sub>2</sub> at WL = -5 cm. For this concentration, however, we could  
209 use only three replicates due to a technical problem with the fourth replicate, resulting in  
210 a loss of statistical power. Even though the inhibitory effect was not constant across  
211 CH<sub>2</sub>F<sub>2</sub> concentrations of 1% [vol/vol] and higher, the differences between these  
212 concentrations were never significant (Fig. 2). Note that the inhibitory effect (and thus the  
213 CH<sub>4</sub> oxidation) at 1% [vol/vol] represented the equivalent of 75% and 64% of the basal  
214 CH<sub>4</sub> flux rate, for WL = -5 cm and WL = -15 cm, respectively. Hence, in this study, CH<sub>4</sub>  
215 oxidation consumed the majority of the CH<sub>4</sub> released via diffusion. When also taking into

216 account the CH<sub>4</sub> release via ebullition, CH<sub>4</sub> oxidation removed only 58% and 43% of the  
217 total CH<sub>4</sub> emitted, for WL = -5 cm and WL = -15 cm, respectively.

218

### 219 3.2. *Completeness of inhibition and recovery of CH<sub>4</sub> emissions*

220 In order to verify whether CH<sub>4</sub> oxidation was inhibited completely, we sampled  
221 air for δ<sup>13</sup>CO<sub>2</sub> analysis before and after injection of CH<sub>2</sub>F<sub>2</sub> and/or <sup>13</sup>CH<sub>4</sub>. Prior to the  
222 additions, the δ<sup>13</sup>CO<sub>2</sub> values of the produced CO<sub>2</sub> were similar for all treatments, ranging  
223 from -25.63 to -27.12‰. Addition of <sup>13</sup>CH<sub>4</sub> in the absence of CH<sub>2</sub>F<sub>2</sub> resulted in a two-  
224 orders-of-magnitude increase of the δ<sup>13</sup>CO<sub>2</sub> (Table 3). The δ<sup>13</sup>CO<sub>2</sub> produced in the soil  
225 columns treated with CH<sub>2</sub>F<sub>2</sub> and <sup>13</sup>CH<sub>4</sub> differed neither from the controls, nor from the  
226 soil columns treated only with CH<sub>2</sub>F<sub>2</sub>. On the other hand, we did observe some  
227 differences between the δ<sup>13</sup>CO<sub>2</sub> before and after addition of CH<sub>2</sub>F<sub>2</sub> and <sup>13</sup>CH<sub>4</sub> (for all  
228 treatments). These differences were expected given the small number of samples used to  
229 construct the Keeling plots (n = 3) and the associated low precision. Because these  
230 differences were inconsistent and always two orders of magnitude smaller compared to  
231 those in the soil columns treated only with <sup>13</sup>CH<sub>4</sub>, we considered these differences  
232 irrelevant.

233 Inhibitions remained complete from at least 2 h after headspace injection until ca.  
234 24 h after the injection and then started to decline. We determined the time span (at a  
235 concentration of 1% [vol/vol] CH<sub>2</sub>F<sub>2</sub>) required for 75% and 90% recovery of the CH<sub>4</sub>  
236 emissions. At WL = -5 cm, roughly two days were needed for 75% recovery and about  
237 three days for 90% recovery (Table 4). At WL = -15 cm, the CH<sub>4</sub> oxidation showed 75%

238 recovery only after eight days and 90% recovery after 12 days (Table 4), which was  
239 significantly longer than for the high WL ( $P < 0.001$  for both 75% and 90% recovery).

240

### 241 3.3. *Inhibitory effects on CO<sub>2</sub> and N<sub>2</sub>O emissions*

242 We further studied whether CH<sub>2</sub>F<sub>2</sub> affected CO<sub>2</sub> and N<sub>2</sub>O emissions. The CO<sub>2</sub>  
243 emissions exhibited no significant response to CH<sub>2</sub>F<sub>2</sub> addition (Fig. 3). Moreover, we  
244 observed no trend in the inhibitory effect on the CO<sub>2</sub> emissions over the different CH<sub>2</sub>F<sub>2</sub>  
245 concentrations. The N<sub>2</sub>O emissions, in contrast, did respond to CH<sub>2</sub>F<sub>2</sub> addition (Fig. 4).  
246 For WL = -5 cm, net N<sub>2</sub>O emissions were too low (near detection limit) for a reliable  
247 evaluation of a possible inhibitory effect. Hence, we determined the response of the N<sub>2</sub>O  
248 emissions to CH<sub>2</sub>F<sub>2</sub> addition only for WL = -15 cm. For this WL, CH<sub>2</sub>F<sub>2</sub> concentrations  
249 equal to or higher than 1% [vol/vol] significantly decreased the net N<sub>2</sub>O flux.  
250 Difluoromethane concentrations of 1% [vol/vol] reduced the N<sub>2</sub>O emissions by about 6%  
251 and concentrations of 3 and 5% [vol/vol] CH<sub>2</sub>F<sub>2</sub> roughly halved N<sub>2</sub>O emissions (Fig. 4).  
252 We remark that the N<sub>2</sub>O emissions increased over the course of the experiment (see also  
253 the positive CH<sub>2</sub>F<sub>2</sub> effect for the control columns in Fig. 4), which implies that the values  
254 in Fig. 4 and Table 2 should not be compared.

255 Last, the recovery of the N<sub>2</sub>O production at 1% [vol/vol] CH<sub>2</sub>F<sub>2</sub> was remarkably  
256 faster than recovery of CH<sub>4</sub> oxidation, with complete recovery of the N<sub>2</sub>O emissions  
257 after, on average, two days (SD = 0.9). Because of the fast recovery and the increase of  
258 the N<sub>2</sub>O emissions over time, we did not determine the time needed for 75% and 90%  
259 recovery.

260 **4. Discussion**

261

262 *4.1. Emissions and inhibitory effects*

263 In accordance with many other studies (see review by Jungkunst and Fiedler,  
264 2007), we observed higher CH<sub>4</sub> emissions at high WL than at low WL, whereas CO<sub>2</sub>  
265 emissions were higher at WL = -15 cm as compared to WL = -5 cm. Also for N<sub>2</sub>O, we  
266 observed higher net flux rates at WL = -15 cm than at WL = -5 cm. This is in agreement  
267 with the conceptual model of Davidson et al. (2000), predicting highest N<sub>2</sub>O production  
268 at intermediately high water-filled-pore-space, where nitrification and denitrification can  
269 proceed simultaneously at aerobic and anaerobic microsites interspersed close to each  
270 other in the soil matrix.

271 Matheson et al. (1997) reported that CH<sub>2</sub>F<sub>2</sub> inhibits methane monooxygenase  
272 (MMO), the enzyme produced by methanotrophic bacteria to oxidize CH<sub>4</sub>. Miller et al.  
273 (1998) found that headspace concentrations as low as 0.03% CH<sub>2</sub>F<sub>2</sub> were sufficient to  
274 completely inhibit CH<sub>4</sub> oxidation in 5 g of dry soil. In contrast, we observed that in our  
275 reconstructed soil columns a headspace CH<sub>2</sub>F<sub>2</sub> concentration of 0.2% [vol/vol] was  
276 insufficient for complete inhibition of CH<sub>4</sub> oxidation. Concentrations equal to or above  
277 1% [vol/vol] CH<sub>2</sub>F<sub>2</sub>, on the other hand, did enhance the CH<sub>4</sub> emissions significantly and  
278 thus inhibited the CH<sub>4</sub> oxidation. Moreover, headspace injection of <sup>13</sup>CH<sub>4</sub> followed by  
279 δ<sup>13</sup>CO<sub>2</sub> analyses verified that this CH<sub>2</sub>F<sub>2</sub> concentration of 1% [vol/vol] inhibited the CH<sub>4</sub>  
280 oxidation completely. In contrast to Miller et al. (1998), we approached natural  
281 conditions more closely, using 28 cm high soil columns in which the inhibitor had to  
282 diffuse into the soil, to the sites where CH<sub>4</sub> oxidation occurs. Also Krüger et al. (2001,

283 2002) and Shrestha et al. (2008) applied 1% [vol/vol] CH<sub>2</sub>F<sub>2</sub> to inhibit CH<sub>4</sub> oxidation in  
284 situ, but no evaluation of the inhibitory effect was reported in these studies.

285 Besides effects of CH<sub>2</sub>F<sub>2</sub> on CH<sub>4</sub> oxidation, Miller et al. (1998) found that, in  
286 some cases, CH<sub>2</sub>F<sub>2</sub> also (partially) inhibited CH<sub>4</sub> production via acetate fermentation,  
287 albeit at concentrations considerably higher than those needed to inhibit CH<sub>4</sub> oxidation.  
288 In our experiment, we could not prove that CH<sub>4</sub> production was unaffected by CH<sub>2</sub>F<sub>2</sub>.  
289 Nonetheless, we believe that inhibitory effects on methanogenesis were very small or  
290 absent, because we observed no significant decrease of the CH<sub>4</sub> emission with increasing  
291 CH<sub>2</sub>F<sub>2</sub> concentrations. At WL = -15 cm, net CH<sub>4</sub> fluxes even showed a slight (but not  
292 significant) increase with increasing CH<sub>2</sub>F<sub>2</sub> concentration.

293

#### 294 4.2. *Recovery of the CH<sub>4</sub> emissions*

295 In accordance with Miller et al. (1998), who reported that the inhibition of CH<sub>4</sub>  
296 oxidation by CH<sub>2</sub>F<sub>2</sub> was reversible, we observed a decrease of the inhibitory effect  
297 (starting about 24 h after CH<sub>2</sub>F<sub>2</sub> addition). We found that the inhibitory effect diminished  
298 significantly more slowly at low WL than at high WL. Possibly, CH<sub>2</sub>F<sub>2</sub> was more easily  
299 removed from the few air-filled pores at WL = -5 cm than from the numerous air-filled  
300 pores (at greater depth) at WL = -15 cm. Moreover, CH<sub>2</sub>F<sub>2</sub> is very soluble in water  
301 (Horvath, 1982) and was thus more quickly diluted at WL = -5 cm than at WL = -15 cm,  
302 potentially clarifying the differences in recovery time. Further, the time needed for 75%  
303 recovery of the CH<sub>4</sub> emissions at WL = -5 cm agrees with observations of Eller and  
304 Frenzel (2001), who reported 70% recovery of the CH<sub>4</sub> emissions one day after CH<sub>2</sub>F<sub>2</sub>  
305 addition of a headspace concentration of 1% [vol/vol] to flooded rice soil samples.

306

307 *4.3. Inhibitory effects on CO<sub>2</sub> and N<sub>2</sub>O emissions*

308 In addition to CH<sub>2</sub>F<sub>2</sub> effects on CH<sub>4</sub> emissions, we also determined the response  
309 of the CO<sub>2</sub> and N<sub>2</sub>O emissions to CH<sub>2</sub>F<sub>2</sub>. Concentrations up to 5% [vol/vol] CH<sub>2</sub>F<sub>2</sub> did  
310 not affect CO<sub>2</sub> emissions. Apparently, the contribution of CH<sub>4</sub> oxidation to the CO<sub>2</sub>  
311 emissions (less than 10%, on average; see Fig. 2 and Table 2) was less than the inherent  
312 variation of the CO<sub>2</sub> emissions.

313 Net N<sub>2</sub>O emissions (at WL = -15 cm), on the other hand, decreased at CH<sub>2</sub>F<sub>2</sub>  
314 concentrations equal to or above 1% [vol/vol], to reach a minimum after approximately  
315 24 h. Subsequently, N<sub>2</sub>O emissions increased again and reached their initial flux rate  
316 several days after the CH<sub>2</sub>F<sub>2</sub> injection (data not shown). At CH<sub>2</sub>F<sub>2</sub> concentrations higher  
317 than 1% [vol/vol], the inhibitory effect on the N<sub>2</sub>O emissions became larger (Fig. 4) and  
318 also the recovery time increased (data not shown). In accordance with our results, Miller  
319 et al. (1998) reported inhibition of NH<sub>4</sub><sup>+</sup> oxidation and of NO<sub>3</sub><sup>-</sup> and NO<sub>2</sub><sup>-</sup> production  
320 after CH<sub>2</sub>F<sub>2</sub> addition. This was not surprising, as ammonium and methane  
321 monoxygenases are similar in function (Bédard and Knowles, 1989). Hence, CH<sub>2</sub>F<sub>2</sub> can  
322 reduce N<sub>2</sub>O production by affecting both N<sub>2</sub>O producing processes: nitrification and  
323 denitrification. Nitrification is inhibited by the effects on ammonium monoxygenase,  
324 while denitrification is reduced because nitrate production via nitrification decreases.

325

326 *4.4. Conclusions*

327 Our results indicated that CH<sub>2</sub>F<sub>2</sub> is capable of completely inhibiting CH<sub>4</sub>  
328 oxidation in a reversible manner also in soil columns, but that considerably higher CH<sub>2</sub>F<sub>2</sub>

329 concentrations are needed to inhibit CH<sub>4</sub> oxidation in soil columns than previously  
330 reported for small soil samples. Furthermore, we demonstrated that CH<sub>2</sub>F<sub>2</sub> addition can  
331 substantially decrease N<sub>2</sub>O emissions. The CO<sub>2</sub> emissions remained unaffected by CH<sub>2</sub>F<sub>2</sub>,  
332 because, for both water levels, the contribution of CH<sub>4</sub> oxidation to the CO<sub>2</sub> flux was  
333 smaller than the natural variation in the latter flux.

334         With regard to future experiments, we conclude that CH<sub>2</sub>F<sub>2</sub> can be a useful tool  
335 for studying controls on CH<sub>4</sub> production and/or oxidation. However, as highlighted in this  
336 experiment, CH<sub>4</sub> release via ebullition can considerably obscure the inhibitory effect and  
337 hence also the determination of CH<sub>4</sub> production and oxidation rates (especially at high  
338 water levels). This problem can be encountered by making continuous measurements and  
339 using median instead of mean flux rates.

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423 **Tables**

424

425 Table 1: Experimental sequence of inhibition treatments with various headspace CH<sub>2</sub>F<sub>2</sub>  
426 concentrations (% [vol/vol]). Each treatment (T) consisted of four replicates. Due to a  
427 technical problem, we have no data of the inhibitions performed on day of year (doy) 239  
428 for the high water level (WL).

429

	<b>WL = -5 cm</b>			<b>WL= -15 cm</b>		
<b>Doy</b>	<b>T 1</b>	<b>T 2</b>	<b>T 3</b>	<b>T 1</b>	<b>T 2</b>	<b>T 3</b>
150	0	0.2	1	0	0.2	1
194	0	1	3	0	1	3
239	0	/	/	0	3	5
289	0	1	2	0	5	/

430

431

432 Table 2: Median net CO<sub>2</sub>, CH<sub>4</sub> and N<sub>2</sub>O fluxes (μmol m<sup>-2</sup> s<sup>-1</sup>) for the two water levels  
 433 (WLs), calculated from the fluxes over the last five days before the first inhibition (day of  
 434 year = 146 till 150) and the standard deviation (SD) over the 12 replicates for each WL.  
 435 We also present a measure for the dispersion of the median fluxes: the average median  
 436 absolute deviation (MAD) over the 12 replicates.

	<b>WL = -5 cm</b>			<b>WL = -15 cm</b>			
	<b>Median</b>	<b>SD</b>	<b>MAD</b>	<b>Median</b>	<b>SD</b>	<b>MAD</b>	<b>P value</b>
<b>CH<sub>4</sub></b>	0.036	0.028	0.02	0.013	0.013	0.005	0.013
<b>CO<sub>2</sub></b>	1.41	0.28	0.043	3.27	0.47	0.040	< 0.001
<b>N<sub>2</sub>O</b>	1.2 * 10 <sup>-5</sup>	8.6 * 10 <sup>-5</sup>	1.2 * 10 <sup>-5</sup>	3.3 * 10 <sup>-4</sup>	3.5 * 10 <sup>-4</sup>	2.3 * 10 <sup>-5</sup>	0.006

437

438 The P value represents the significance level of the difference between the two water  
 439 levels.

440

441 Table 3: Isotopic signature ( $\delta^{13}\text{CO}_2$  in ‰) of the  $\text{CO}_2$  produced in the soil columns before  
 442 and after addition of  $\text{CH}_2\text{F}_2$  and/or  $^{13}\text{CH}_4$ , calculated via a Keeling plot and averaged over  
 443 the three replicates used for each treatment of the two water levels (WLs). We also  
 444 present the 95% confidence interval (95% CI) of this average.  
 445

	WL = -5 cm				WL = -15 cm			
	Before injection	95% CI	After injection	95% CI	Before injection	95% CI	After injection	95% CI
<b>Control, no <math>^{13}\text{CH}_4</math></b>	-26.39	1.01	-27.61	3.02	-26.89	0.99	-19.56	4.09
<b>Control, <math>^{13}\text{CH}_4</math></b>	-26.38	1.10	825.17	51.47	-27.12	0.65	688.77	26.88
<b>Inhibited, no <math>^{13}\text{CH}_4</math></b>	-26.11	0.98	-19.94	1.81	-25.63	0.83	-29.15	1.53
<b>Inhibited, <math>^{13}\text{CH}_4</math></b>	-26.62	0.80	-21.00	3.78	-25.84	0.87	-24.07	3.21

446

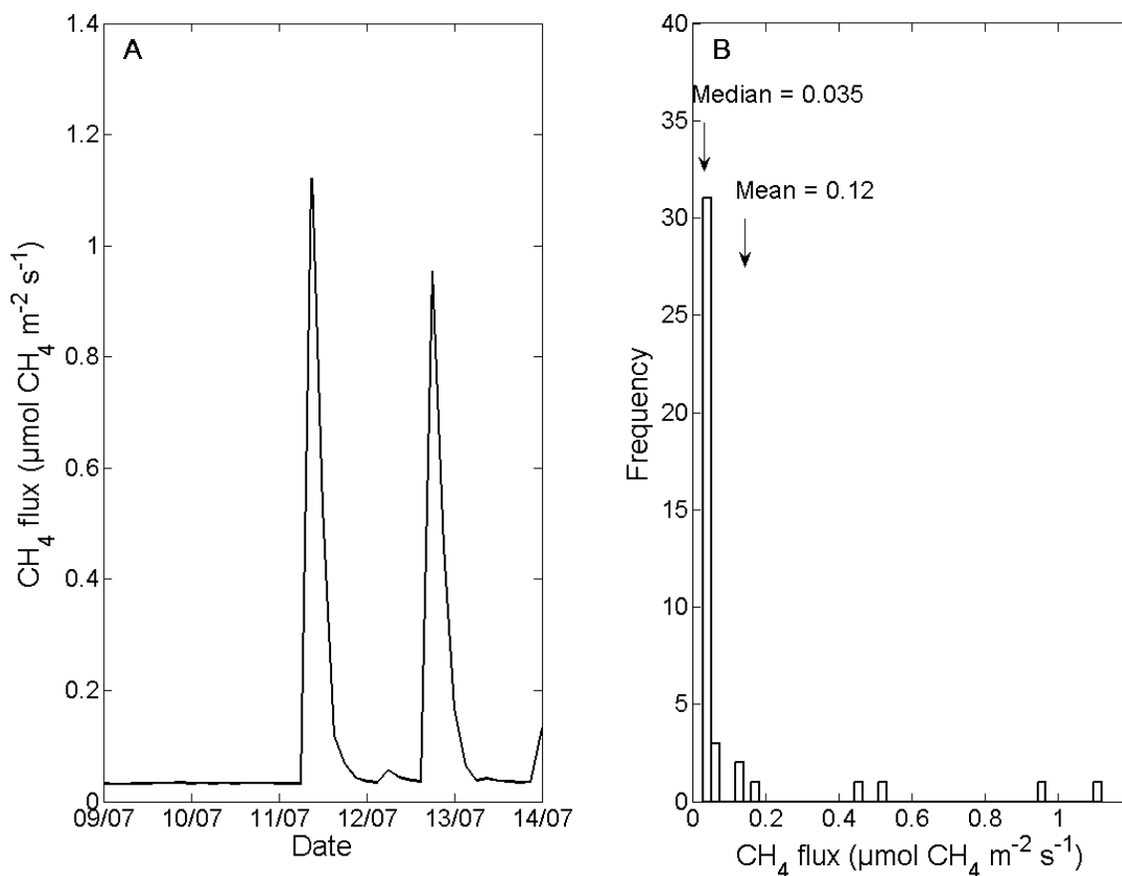
447 Table 4: Mean time (days) after addition of 1% [vol/vol] CH<sub>2</sub>F<sub>2</sub> needed to reach 75% and  
448 90% recovery from the inhibition for the two water levels (WLs), and the standard  
449 deviation (SD) on this mean.

450

	<b>75% reduction of inhibitory effect</b>		<b>90% reduction of inhibitory effect</b>	
	<b>Mean</b>	<b>SD</b>	<b>Mean</b>	<b>SD</b>
<b>WL = -5 cm</b>	1.9	2.1	2.9	2.0
<b>WL = -15 cm</b>	7.8	3.7	11.6	5.5

451

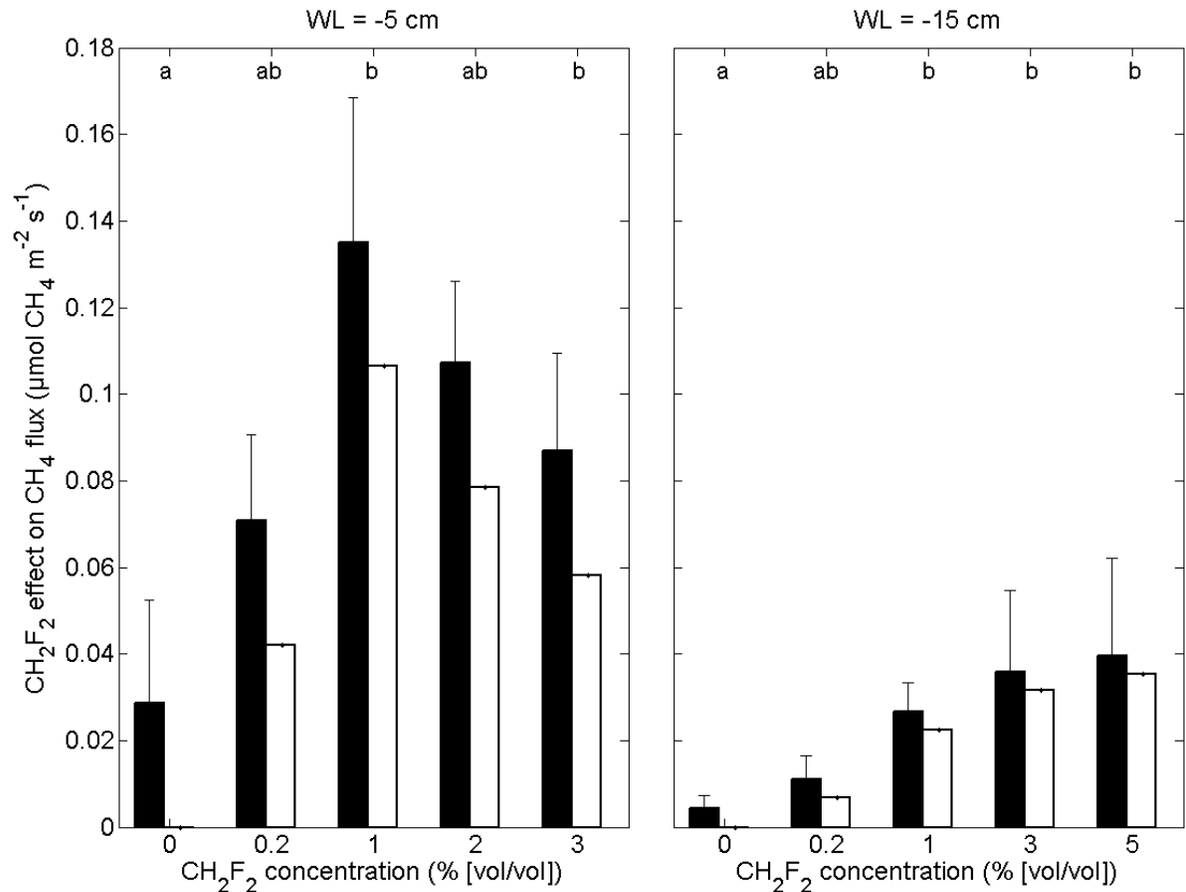
452 **Figures**



453

454 Figure 1: We present an example of the CH<sub>4</sub> fluxes of one replicate over five days before  
 455 an inhibition (A) and the histogram showing the distribution of these CH<sub>4</sub> fluxes (B). We  
 456 also present the median and mean CH<sub>4</sub> flux computed from these data. In this figure, we  
 457 demonstrate that CH<sub>4</sub> release via ebullition can substantially affect the mean CH<sub>4</sub> flux,  
 458 whereas the effect on the median CH<sub>4</sub> flux is small or inexistent (adding or removing a  
 459 few ebullition events will not affect the median flux, but will enhance or reduce the mean  
 460 CH<sub>4</sub> flux substantially).

461



462

463 Figure 2: Inhibitory effect on the net CH<sub>4</sub> fluxes at different CH<sub>2</sub>F<sub>2</sub> concentrations and for

464 the two water levels (WLs). Black bars were calculated as the average difference between

465 the median CH<sub>4</sub> flux before the inhibition (over five days) and after the inhibition (over

466 24 h). White bars represent the CH<sub>2</sub>F<sub>2</sub> effect corrected for changes in the control columns

467 (i.e., we subtracted the effect at 0% [vol/vol] from the effect at each inhibition treatment).

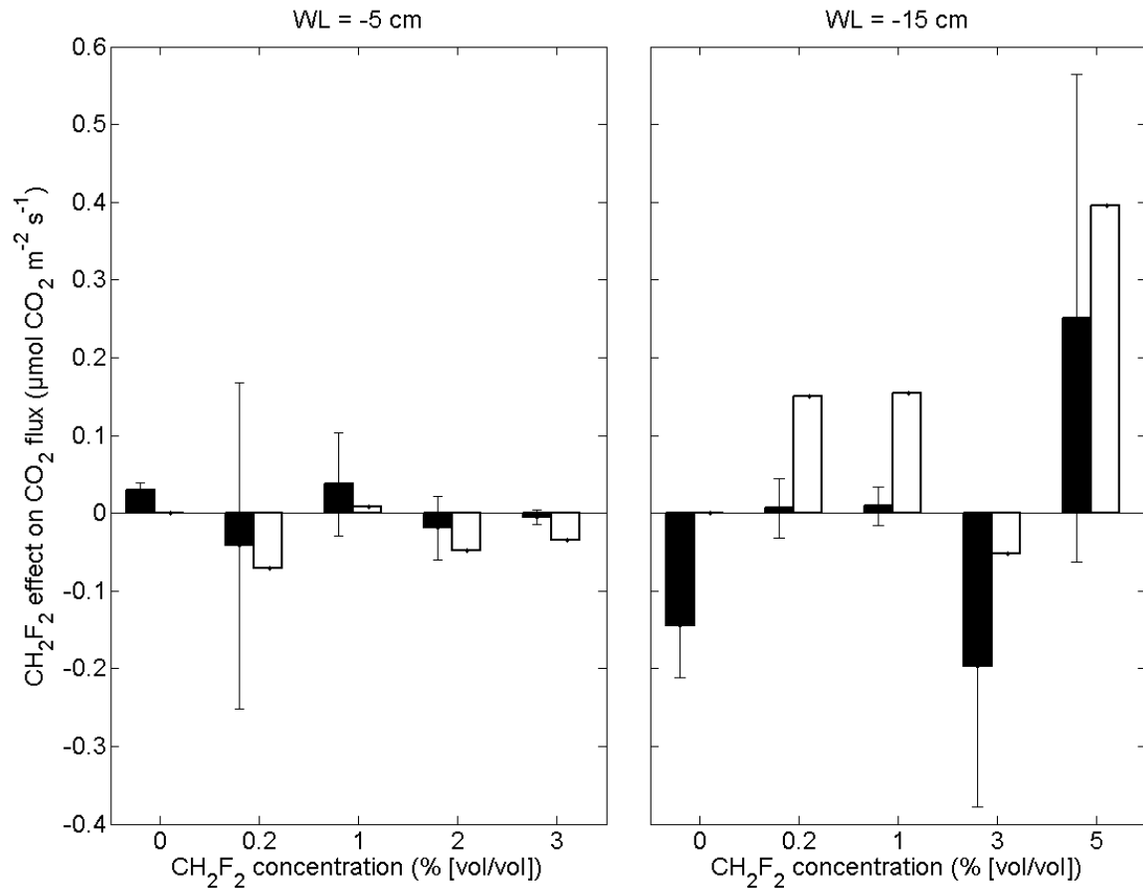
468 Error bars represent the standard error and for each WL, letters indicate the differences

469 significant at P < 0.10. Note that the lack of a significant inhibitory effect at 2% [vol/vol]

470 CH<sub>2</sub>F<sub>2</sub> for WL = -5 cm is very likely due to loss of statistical power, as we could use only

471 three of the four replicates at this CH<sub>2</sub>F<sub>2</sub> concentration.

472



473

474 Figure 3: Difluoromethane effect on the net CO<sub>2</sub> fluxes at different CH<sub>2</sub>F<sub>2</sub> concentrations

475 and for the two water levels (WLs). Black bars were calculated as the average difference

476 between the median CH<sub>4</sub> flux before the inhibition (over five days) and after the

477 inhibition (over 24 h). White bars represent the CH<sub>2</sub>F<sub>2</sub> effect corrected for changes in the

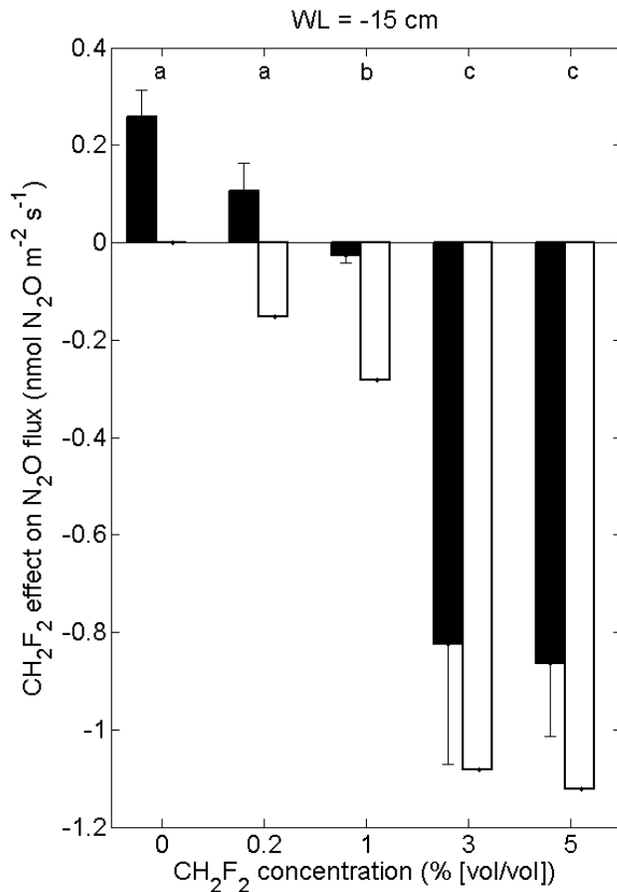
478 control columns (i.e., we subtracted the effect at 0% [vol/vol] from the effect at each

479 inhibition treatment). Error bars represent the standard error. None of the differences are

480 statistically significant at  $P < 0.10$ .

481

482



483

484 Figure 4: Difluoromethane effect on the net N<sub>2</sub>O fluxes at different CH<sub>2</sub>F<sub>2</sub> concentrations  
 485 for WL = -15 cm. Black bars were calculated as the average difference between the  
 486 median CH<sub>4</sub> flux before the inhibition (over five days) and after the inhibition (over 24  
 487 h). White bars represent the CH<sub>2</sub>F<sub>2</sub> effect corrected for changes in the control columns  
 488 (i.e., we subtracted the effect at 0% [vol/vol] from the effect at each inhibition treatment).  
 489 Error bars represent the standard error and for each WL, letters indicate the differences  
 490 significant at P < 0.10.