

WSSI Consortium Projects Progress Report

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WSSI Grant #	25000.01E7	Grant Years:	2010-2013
Progress for the period:		Year: 2010-13 (Final Report)	
University	Duke University		
Project Title	A comparison of wetland functions and services on restored wetlands of the Piedmont: carbon storage and GHG release estimates.		
Prepared by:	Date: 10/31/2014		

NARRATIVE: Summarize activities accomplished during this reporting period using only space provided below (10 pt. min).

Restoration sites in Virginia were studied to determine changes in soil carbon flux in response to the experimental carbon additions. Analysis of bulk density, total carbon and total nitrogen were consistent with earlier studies at the site, although lower amounts of soil organic matter (OM) were found. However, plots that received greater OM amendment loads have higher total carbon and nitrogen and lower bulk density. There is concern that widespread restoration and/or creation of wetlands may present a radiative forcing hazard because of the potential for high rates of methane (CH₄) emissions. Yet data on greenhouse gas (GHG) emissions from restored wetlands remains relatively sparse and there has been little investigation into the GHG effects of amending wetlands with soil organic matter (OM), a practice used to improve function in mitigation wetlands in the Eastern United States. In this study we evaluate the effect of added OM on GHG across an organic matter gradient at the Charles City Wetland (CCW) in Charles City County, Virginia, ten years post original OM additions. Our data suggest that soils heavily loaded with OM are emitting significantly more CO₂ than those that have received little or no OM amendment. Emissions of CH₄ are low compared to those of other forested wetlands in the region and show no relationship with the loading rate of added OM or total soil carbon. We conclude that adding moderate amounts of OM to the CCW does not greatly increase GHG emissions, while the addition of high OM loading rates produces additional CO₂, but not CH₄.

COMMENTS: Note any delays, problems, or special circumstances affecting progress and how you intend to address them.

See the attached final report for a complete analysis of the study. A new method for reducing variability and errors in measuring GHG fluxes from static chambers was developed for this study and is presented in the report as well as annual GHG fluxes

Project Benchmarks and Deliverables	% Completion	Anticip. Completion Date
Collars installed for gas sampling	100	Complete
Wells and soil moisture probes installed and tested	100	March, 2012
A full year of monthly greenhouse gas flux data collected	100	April 2013
Manuscript submitted for publication	1	November 2014

1 WSSI Project Final Report
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5
6 October 29, 2014

7 **Title**

8 The effects of organic matter amendment on greenhouse gas emissions from a mitigation
9 wetland in Virginia's coastal plain

10 **Abstract**

11 There is concern that widespread restoration and/or creation of wetlands may present a
12 radiative forcing hazard because of the potential for high rates of methane (CH₄) emissions. Yet
13 data on greenhouse gas (GHG) emissions from restored wetlands remains relatively sparse and
14 there has been little investigation into the GHG effects of amending wetlands with soil organic
15 matter (OM), a practice used to improve function in mitigation wetlands in the Eastern United
16 States. In this study we evaluate the effect of added OM on GHG across an organic matter
17 gradient at the Charles City Wetland (CCW) in Charles City County, Virginia. Our data suggest
18 that soils heavily loaded with OM are emitting significantly more CO₂ than those that have
19 received little or no OM amendment. Emissions of CH₄ are low compared to those of other
20 forested wetlands in the region and show no relationship with the loading rate of added OM or
21 total soil carbon. We conclude that adding moderate amounts of OM to the CCW does not
22 greatly increase GHG emissions, while the addition of high OM loading rates produces
23 additional CO₂, but not CH₄.

24 **Introduction**

25 Despite making up only five to eight percent of world land cover (Mitsch and Gosselink
26 2007), wetland ecosystems play an important role in regulating the Earth's climate. Wetland
27 soils contain 16 to 33 percent of the earth's soil Carbon (C) pool of 2,500 Pg (Lal 2005;
28 Bridgham et al 2006) and emit 20 to 40 percent of methane (CH₄) (Bloom et al., 2010), an
29 important greenhouse gas (GHG)(Myhre et al 2013).

30 A review of North American wetland C exchange found that because of CH₄ emissions,
31 most wetlands are net emitters of GHG on century timescales and therefore: "...large CH₄
32 emissions from conterminous US wetlands suggest that creating and restoring wetlands may
33 increase net radiative forcing..."(Bridgham et al 2006). Others have claimed that because
34 wetlands are sustainable ecosystems and persistent as C sinks, the widely-used 100-year time
35 horizon is too short, and that: "...wetlands can be created and restored to provide C sequestration
36 and other ecosystem services without great concern of creating net radiative sources on the
37 climate due to methane emissions" (Mitsch et al 2013). But errors in both the math and
38 reasoning underpinning this latter view have been exposed, which reaffirms the potential
39 century-scale impact of restored and created wetland CH₄ emissions on regional climate budgets
40 (Neubauer 2014; Bridgham et al 2014).

41 While this controversy over the C balance of wetland restoration and creation is partly a
42 disagreement about the appropriate use and calculation of global warming potential, versus
43 sustained flux models, which account for annual pulses of GHGs (i.e. Frohling et al., 2006;
44 Neubauer, 2014), it also reflects the great uncertainty (100%) around wetland GHG flux
45 estimates (Bridgham et al 2006). It thus may be particularly difficult to make long-term
46 assumptions regarding restored and created wetland GHG fluxes given their complex histories of

47 human disturbance and intervention and that they routinely fail to achieve the same ecological
48 function of reference ecosystems over short timescales (Zedler and Callaway 1999). An
49 important remaining question is whether restored freshwater wetlands with mineral soils are in
50 fact a sink or source of GHG over policy-relevant timescales?

51 In the eastern United States large areas of wetlands are created as part of compensatory
52 mitigation mandated by section 404 of the Clean Water Act, and they commonly suffer from an
53 initial deficiency of soil organic matter (OM) (Stauffer and Brooks 1997; Whittecar and Daniels
54 1999) compared to “natural” wetlands (Bailey et al 2007). Many studies have advocated for the
55 amendment of created wetlands with OM in the form of salvaged topsoil or mulch to help them
56 achieve reference functionality (Stauffer and Brooks 1997; Whittecar and Daniels 1999; Bruland
57 and Richardson 2004). Indeed, studies have found that moderate loading of OM into a created
58 wetland increase woody plant development (Bailey et al 2007) and soil functions, such as
59 microbial biomass and denitrification enzyme activity (Bruland and Richardson 2009; Sutton-
60 Grier et al 2009).

61 Few studies have measured GHG emissions from created or restored wetlands and fewer
62 still have done so at sites amended with OM. It is unclear whether or not the practice of adding
63 OM to created wetlands will have an effect on their radiative impact.

64 Increased C substrate and/or productivity due to nutrient content of added OM could
65 enhance CH₄ flux given the relationship between OM loading rate and primary productivity
66 (Bailey 2006), which across wetland systems has been correlated with CH₄ flux rate (Whiting
67 and Chanton 1993). Alternatively, added OM could reduce CH₄ emissions by altering the
68 physical structure of soil. The addition of OM increases soil elevation (Bailey et al 2007) and
69 reduces bulk density (Bruland and Richardson 2009), which could allow surface soil to remain

70 more oxic, facilitating methane oxidation as well as aerobic, rather than anaerobic methanogenic,
71 respiration.

72 The purpose of this study is to investigate how a gradient of added OM affects GHG
73 emissions from a created mitigation wetland on mineral soils. Included in our analysis is an
74 estimate of how long it would take for our restored wetland to change from a GHG source to a
75 sink, calculated as the radiative forcing switchover time following (Frolking et al 2006).

76 **Methods**

77 **Site description**

78 The study took place within the 20.8-hectare Charles City Wetland Mitigation Site
79 (CCW), which is located in Charles City County, Virginia, USA, and owned by the Virginia
80 Department of Transportation (VDOT) as part of its compensatory mitigation program (Bailey et
81 al., 2007; see Fig. 1AB). Precipitation is the dominant hydrologic input and the CCW may hold
82 up to 0.5 m of standing water during cooler months (Bailey et al 2007). Site history is described
83 in detail by Bergschneider (2005) and Bailey et al. (2007), but briefly summarized here. Prior to
84 restoration the site was covered by upland mixed hardwood forest that had been partially
85 converted to agricultural field. The soil was mapped as a complex of Chickahominy (fine,
86 mixed, semiactive, thermic Type Endoaquults) and Newflat (fine, mixed, subactive, thermic
87 Aeric Endoaquults) (Bergschneider 2005). Mitigation efforts attempted to convert field and
88 remnant forest to wetland status during the winter of 1997-1998 by excavating into the subsoil (E
89 or Btg horizon) to the depth of the presumed seasonal high water table. After revegetation, many
90 parts of the site were found to be covered in facultative or upland plant species with much less
91 hydrophytic cover than desired for mitigation purposes, a result attributed to restoration activities
92 in which topsoil was lost, leaving compacted, low organic matter (OM) subsoil at the surface.

93 The addition of an OM source had been proposed as a method for improving function of
94 mitigation wetlands (Stauffer and Brooks 1997), but no data existed regarding the quantity of
95 added OM required to achieve sufficiently improved wetland function in this setting. With a
96 goal of determining optimal OM amendment loads for the wetland, a research group from
97 Virginia Polytechnic Institute and State University implemented a gradient experiment in 2001
98 with 4 replicate plots of 4 OM loading rates (plus control) in wet and dry experimental blocks
99 (see Fig. 1C). Municipal wood and yard waste compost was rototilled into the topsoil of 4.6 by
100 3.1 m plots at loading rates of 56, 112, 224 and 336 kg m⁻² in July, 2002. Control plots received
101 only rototilling. Each plot was planted with five Pin Oak (*Quercus palustris*) and River Birch
102 (*Betula nigra*) saplings, but otherwise the site was allowed to revegetate naturally from seed
103 bank. In January, 2013 we found a mean count of 3.4 *Q. palustris* and 4.6 *B. nigra* survived in
104 each 14.3 m² plot with some volunteer tree species, such as Red Maple (*Acer rubrum*) and Black
105 Willow (*Salix nigra*), established sporadically.

106 **Site Characterization**

107 We measured the relative elevation of each plot near the gas collars used for measuring
108 GHGs using a Topcon RL-H3A laser level and collected soil cores in each plot in September,
109 2011 using a 10-cm diameter soil-corer. Cores were split into 0 to 5 and 5 to 10 cm depth
110 sections in the field. In the lab each core section was weighed wet and a subsample was weighed,
111 oven-dried and re-weighed to estimate wet:dry ratios and calculate bulk density. Subsamples
112 were analyzed for total carbon (C) and total (N) using a CE Instruments Flash (1112 series)
113 Elemental Analyzer. We sampled soils again in September, 2012 using a punch tube and
114 separated depth sections of 0 to 2 cm, 4 to 6, 9 to 11 and 19 to 21 cm in the field, and then
115 composited corresponding depths from three replicate punches. These soils were analyzed for

116 total C, total N (following the same method as above), digested following a nitric-perchloric acid
117 method followed by colorimetric analysis of total phosphorus (P) using a Beckman DU-64
118 spectrophotometer, Meilich-3-extractable P, KCl-extractable nitrate/nitrite (NO_x) and
119 ammonia/ammonium (NH_x) using a Lachat Quickchem 8000 autoanalyzer. We installed litter
120 fall traps (approximately 30 cm^2) in each plot in September, 2012 and litter was collected during
121 subsequent site visits.

122 **Greenhouse Gas Sampling**

123 In late summer 2011 we imbedded 20 cm diameter PVC collars 10 to 15 cm into the soil
124 in each plot of the wet block for static chamber GHG gas sampling (Livingston and Hutchinson,
125 1995). During chamber setup we placed a PVC cap with a rubber gasket over collars, but after
126 sampling in September and October, 2011 and February, 2012, we found that this chamber
127 design and/or sampling technique was producing CH_4 data that frequently failed to follow a
128 linear pattern of accumulation within chamber headspace. CO_2 concentrations accumulated in a
129 linear fashion within headspace as expected, but extraordinarily high initial CH_4 concentrations
130 (up to 1500 ppm; roughly 1000 time ambient concentration) within the headspace indicated that
131 capping the collar and/or standing near the collar during sampling was purging CH_4 stored within
132 soil pores. To mitigate this problem we redesigned our chambers and collars in spring of 2012 to
133 minimize collar disturbance during chamber setup. We accomplished this by building new
134 permanent collars with gutters that could be filled with water, capped and sampled from a
135 distance of 2 m (see Fig. 2). An internal computer fan powered by a 9-volt battery circulates
136 chamber head space from which air samples are extracted using a 2 m tube, 1 mm inner diameter
137 plastic tube. Chamber caps were also equipped with a thermocouple allowing for internal
138 chamber temperature (T) to be recorded during each sample extraction and we coated them with

139 reflective aluminum foil to minimize solar warming as recommended by the US Department of
140 Agriculture (Parkin and Venterea 2010). We installed these new collars in April, 2012 and
141 sampled for estimation of trace gas flux every two months from May, 2012 until January, 2013.

142 On each sampling date we collected headspace gas four times over the course of a half-
143 hour incubation from collars in each of the 20 plots. Following placement of the chamber top on
144 the collar we immediately extracted a 50-ml headspace sample via a plastic syringe and
145 deposited it into a mylar gas-tight sample bag. We recorded ambient air T, internal chamber T,
146 soil T at 5 cm depth for initial and subsequent samples taken approximately 5, 15 and 30 minutes
147 following chamber setup. Gas bags were transported to the Duke University Wetland Center
148 laboratory and analyzed within one week of sampling on a Varian 450 Gas Chromatograph (GC)
149 equipped with a flame ionization detector, methanizer, and electron capture detector to analyze
150 CH₄, CO₂ and nitrous oxide (N₂O) concentrations synchronously. All samples were run in
151 duplicate and when duplicate values differed by <10% the mean was used for gas flux
152 calculations. Flux rate was estimated by linear regression of sample concentrations as a function
153 of time elapsed. If a threshold r-squared value of 0.90 was not met, one outlying point was
154 occasionally (approximately 5% of incubations) removed to improve fit.

155 **Supplementary Data**

156 On each sampling date after finishing headspace incubations we measured soil moisture
157 in the top 5 cm using a Fieldscout 100 time domain reflectometry probe (Spectrum
158 Technologies). We recorded the mean of five measurements taken adjacent to each chamber
159 collar.

160 In September 2012 we installed pore water wells in each plot and starting in November,
161 2012 began collecting pore water samples for subsequent analysis of total dissolved C and

162 dissolved organic C using a Shimadzu TOC-5000 A, total phosphorus (P) following persulfate I
163 digestion method (Wetzel and Likens 1979), nitrate/nitrite (NO_x) and ammonia/ammonium
164 (NH_x) using a Lachat Quickchem 8000 autoanalyzer (EPA method 350.1).

165 **Statistical analyses**

166 We used ANOVA to test for differences in gas flux between groups of plots with
167 different OM treatments and linear regression to look for trends in gas flux across the OM
168 gradient. We evaluated all data for normality by generating box-and-whisker, histogram and
169 quantile-quantile plots and log-transformed data if necessary. We explored relationships between
170 gas flux and potential explanatory variables using the Ecodist package (Goslee and Urban, 2007)
171 and by building generalized linear models (GLM). We used JMP Pro 11 (SAS Institute Inc.) to
172 plot GLM outputs. All other statistics were computed using the R programming language (R
173 Core Team 2013) and in Microsoft Excel 2010. We estimated annual emissions of CO_2 and CH_4
174 by extrapolating hourly flux from each sampling day across the nearest adjacent unsampled days.

175 **Carbon Balance**

176 We compare the relative radiative impacts of soil CH_4 and CO_2 fluxes by multiplying
177 CH_4 by its 100-year sustained global warming potential of 38 (Neubauer 2014) and estimate
178 radiative forcing switchover time (Frolking et al 2006) for the CCW using net ecosystem
179 exchange (NEE) data (Bailey 2006) and CH_4 flux data generated in this study. Bailey (2006)
180 found NEE to be negative for most of the CCW plots because of rapid oxidation of added OM,
181 therefore we used his positive mean NEE values from the lowest loading rates (141.1 and 29.9 g
182 $\text{CO}_2\text{-C m}^{-2} \text{ yr}^{-1}$) to generate a range of radiative potential radiative forcing switchover times.

183 **Results**

184 **Hydrology and Soil Elevations**

185 Water level data suggest that the hydrology of CCW is controlled by precipitation inputs
186 with storm events and dry spells driving periodic fluctuations of more than 1 m in the water table
187 (see Fig. 3). Pondered water was present at the site 59 percent of the time from 22 February, 2012
188 to 21 January, 2013 and reached a maximum depth of 14 cm above the mean elevation of
189 unamended plots. The distribution of plot elevations is approximately normally distributed
190 with a standard deviation of 4 cm and two outliers: a 12 cm “hummock” and a -9 cm “hollow.”
191 Pairwise comparison (ANOVA) of plots grouped by OM loading rate shows no significant
192 differences in mean elevation, though there is a weak ($r^2 = 0.18$), but significant ($p < .05$)
193 positive linear trend in elevation across the OM gradient.

194 **Soil Nutrients**

195 Total soil C data shows that while some of the added OM may have been lost since 2005
196 (Bailey et al 2007), particularly from plots loaded with 112 and 224 mg ha⁻¹ OM, the gradient, as
197 originally established, persists (see Fig. 4), with total C in the top 10 cm of soil ranging from
198 approximately 2 to 13 percent. Mean litter fall across the plots during the fall of 2012 was 0.37
199 ± 0.045 kg m⁻², which assuming litter is 50% C by weight (Bocock 1964), represents an input of
200 0.19 ± 0.023 kg C m⁻² yr⁻¹ to surface soils.

201 Total soil C, N and P are generally higher in plots that received higher loading rates of
202 OM, but decreases with depth such that differences between loading rates are negligible at 10
203 and 20 cm depth (see Fig. 5A-5C). KCl-extractable NH_x and NO_x and Mehlich-3-extractable P
204 follow roughly similar patterns, with some exceptions (see Fig. 5D-5F). KCl-extractable NH_x
205 shows no clear pattern related to loading rate and KCl-extractable NO_x is uniformly low with as
206 much variability with depths and loading rates as across them.

207 We observe a strong linear correlation ($r^2=.96$) between total soil C and N had throughout
208 the top 20 cm of soil (see Fig. 6A). The relationship between total N and KCl-extractable NH_x is
209 much weaker ($r^2=.56$; see Fig. 6C). Total soil C and P show a logistic correlation ($r^2=.67$; Fig.
210 6B), and total P and Mehlich-3-extractable P show a relatively weaker but significant ($r^2=.58$)
211 quadratic correlation (see Fig. 6D).

212 **GHG Fluxes**

213 We analyze the three most-important GHGs (CO_2 , CH_4 and N_2O). Because of the high
214 spatial and temporal heterogeneity in N_2O flux (Firestone 1982; Groffman et al 2009), and the
215 fact we that found N_2O flux to be below minimum detection thresholds for approximately 90
216 percent of incubations we focused our results and discussion on CH_4 and CO_2 flux.

217 **CO_2 Flux**

218 The highest CO_2 fluxes ($>400 \text{ mg m}^{-2} \text{ hr}^{-1}$) were observed during warmer, drier months
219 and contrast with fluxes approaching minimum analytical detection limits during cold, wet
220 months (Fig. 7). CO_2 emissions from soil directly responded to increases in soil T (Fig.8) and in
221 general, the higher CO_2 emissions are associated with higher OM loading rates; linear regression
222 of log-transformed CO_2 flux as a function of OM treatment shows significant positive
223 relationships across all sampling months except September (Table 1). The relationship between
224 OM and CO_2 emission is strongest during peak flux in July which is one of only two months (the
225 other being January) where significant differences in CO_2 flux between OM treatments occur.
226 From summed monthly data we estimate an annual soil CO_2 flux ranging from 0.33 ± 0.019 to
227 $0.71 \pm .11 \text{ kg CO}_2\text{-C}$ from the respective low to high end of the OM gradient.

228 A GLM with three parameters: soil T, soil volumetric water content (SVWC), and soil
229 total C (top 5 cm), explains much of the variability ($r^2 = 0.75$) in CO_2 flux across all sampling

230 dates (Fig. 9A). During any given sampling date soil T and soil moisture are essentially constant
231 across plots (relative to seasonal changes) and cannot explain differences in soil respiration. For
232 example, variability in July CO₂ flux could only be partially explained ($r^2 = 0.61$) by a GLM
233 incorporating soil total C (top 5 cm) and soil total N (20 cm depth; Fig. 9B).

234 **CH₄ Flux**

235 We find CH₄ flux rates above minimum analytical detection thresholds only when soil T
236 was at least 18 °C and some ponded water was present at the CCW (see Table 2). We identify a
237 threshold of 50 percent SVWC, below which CH₄ was never greater than 0.13 mg CH₄ m⁻² hr⁻¹
238 (Fig. 10). When conditions at the CCW are favorable for methanogenesis (soil T > 15 °C and
239 ponded water), flux rates are highly variable across plots. Maximum observed CH₄ flux rates are
240 approximately 3 to 5 mg m⁻² hr⁻¹. We estimate an annual efflux of 40.5 kg CH₄-C ha⁻¹ yr⁻¹ from
241 our bi-monthly measurements (see Table 4). We are unable to detect any statistically significant
242 patterns in CH₄ flux related to soil C or OM loading rate.

243 **Carbon Balance**

244 During the sampling dates when CH₄ flux was large enough to be detectable, its
245 contribution to radiative forcing was relatively minor compared to soil CO₂ flux based on a 100-
246 year sustained global warming potential of 38 for CH₄ (Neubauer 2014) (Fig. 11).

247 **Discussion**

248 **Hydrology**

249 Wetland GHG flux is moderated by hydrologic dynamics because saturation inhibits
250 decomposition and creates conditions favorable for CH₄ emission (Whalen 2005). Thus it is
251 important to consider site hydrology as we discuss gas flux. Our hydrologic data are consistent
252 with previous work indicating that the CCW is a groundwater recharge system with hydrologic

253 inputs dominated by precipitation (Despres 2004). The CCW was relatively wet during the 2012
254 growing season when it received 82 cm of rain (7.5 percent above mean; National Climatic Data
255 Center; Lawrimore et al 2011) and held ponded water 52 percent of the time. This contrasts with
256 conditions during the 2005 growing season when the CCW received 10 percent less rainfall than
257 average (National Climatic Data Center; Lawrimore et al 2011) and water was ponded just 25%
258 of the time (Bailey et al 2007).

259 **Elevation and OM incorporation**

260 During OM addition to the CCW in 2001 there was difficulty in completely incorporating
261 the highest OM loading rates into plots, which led to mounding (Daniels et al 2005). We found
262 micro-elevational differences between plots to be less pronounced in 2012 compared to
263 conditions in 2005 reported by Bailey et al. (2007). The relationship between OM loading rate
264 and elevation was far weaker in 2012 (see Table 3), which could be the result of settling or
265 subsidence due to more rapid OM oxidation in elevated, high-OM plots. The higher rates of soil
266 respiration that we and Bailey et al. (2007) detected coming from higher OM plots are consistent
267 with an oxidation-subsidence explanation for the loss of elevation, as is the discrepancy in total
268 soil C between 2005 and 2012 we observed (see Fig. 4).

269 **CO₂ flux**

270 Our annual soil respiration budget cannot account for the soil C losses in the 112 and 224
271 Mg ha⁻¹ plots we observe between 2005 and 2012, which were approximately 1 and 5 percent,
272 respectively, corresponding to respective losses of 1.5 and 5.5 kg C m⁻² yr⁻¹ over seven years.
273 This rate is an order of magnitude greater than our estimated annual soil respiration loss from
274 these plots: 0.42 and 0.49 ± 0.032 kg CO₂-C m⁻² yr⁻¹ respectively. Therefore we suspect that
275 some of the C loss may be due to leaching of dissolved OM and/or transport of particulate OM

276 during floods. In calculating the annual budget we assume that CO₂ flux will be similar on
277 average across a 2-month window to what we measure during our relatively short period of
278 observations, which means that our calculations are susceptible to bias from idiosyncrasies of
279 weather preceding each sampling date. Such effects could be especially pronounced during the
280 fall and spring, which experience extreme within-season and inter-annual climate variability.

281 However, the overall seasonal pattern in soil CO₂ flux we observe is similar to what
282 Bailey (2006) reported from the CCW for 2005/2006 with peak respiration of greater than 400
283 mg m⁻² hr⁻¹ during summer dry spells and low CO₂ flux of less than 100 mg m⁻² hr⁻¹ during wet
284 winter months. The positive relationship between CO₂ flux and soil OM loading rate is also
285 consistent with Bailey's (2006) results.

286 Soil respiration rate is typically limited by T and oxygen availability, so it is not
287 surprising that soil T and SVWC are the two most important terms in our generalized linear
288 model explaining log-transformed CO₂ flux variability across seasons, with r-squared values of
289 .50 and .49 respectively. Soil T and SVWC are slightly correlated with each other (r-squared of -
290 .40), but this relationship is driven by one sampling date in July when the site was both very
291 warm and very dry. Including both soil T and SVWC improves model r-squared to .71. The
292 third model parameter, total surficial soil C simply reflects the amount of OM available to be
293 decomposed. The effects of OM on CO₂ flux become obvious when the site is sufficiently dry
294 (i.e. July), but during wetter periods the importance of surface soil C is obscured. So while soil
295 C is very weakly correlated with log-transformed CO₂ flux across all sampling dates (r-squared
296 of .05), including it in the GLM helps improve fit (r-squared of .75) and reduces the Akaike
297 information criterion (AIC).

298 With T and soil moisture held relatively constant across the site during a given sampling
299 date, we found surface soil C to be the most important parameter explaining CO₂ flux in July (r-
300 squared of .52). The inclusion of total soil N at 20 cm depth improved our model r-squared to
301 .63 and it was not highly correlated with surface soil C (r-squared of .24). We assume that soil N
302 at depth correlates with CO₂ flux because a greater N pool in the rooting zone should stimulate
303 higher rates of autotrophic and heterotrophic respiration related to N mineralization (Schlesinger
304 1997).

305 **CH₄ flux**

306 Hydrology and T both control rates of methane production by dictating oxygen
307 availability and demand (Whalen 2005), which explains why we found CH₄ flux to be very low
308 during cold and/or dry periods. CH₄ flux variability is consistent with results from other forested
309 wetlands of the Southeastern US but our annual CH₄ flux estimate was on the low end of the
310 range of published estimates for analogous systems (See table 4).

311 CH₄ flux shows no significant relationship with OM loading rate, suggesting that if
312 excess nutrients and enhanced primary productivity are increasing methane production, then the
313 increase is being cancelled out by increased oxidation. This result contrasts with that of
314 Ballantine et al. (in press) which shows that addition of OM led to higher rates of potential net
315 methane emissions from intact soil cores compared to controls (Ballantine et al., in press).
316 Higher soil moisture in amended plots correlate with Ballantine et al.'s observed differences in
317 CH₄ production and they suggest that OM amendments increased water retention creating
318 conditions more favorable for methanogenesis. At CCW, OM additions appeared to have the
319 opposite effect on soil moisture because of the slight mounding effect described above. Our data
320 from the relatively drier months of May and July show weak (r-squared of 0.16 and .014,

321 respectively), marginally significant ($p < 0.09$ and $p < 0.11$, respectively) relationship between
322 SVWC and OM loading rate.

323 The higher CH_4 production in response to added OM found by Ballantine et al. (in press)
324 appears to be an indirect effect caused by increased soil moisture as there was no relationship
325 between C quality and CH_4 flux among different types of added OM. Therefore increasing soil C
326 by adding OM does not necessarily provide additional C substrate for methanogens, but it may
327 alter methane production and/or oxidation because of indirect hydrologic effects. Heavy OM
328 addition may elevate the soil surface allowing for more oxic conditions, or conversely, increased
329 OM may enhance water holding capacity facilitating anoxia (Ballantine et al., in press).

330

331 **Carbon Balance**

332 The radiative forcing switchover time (Frolking et al 2006) for CCW is highly uncertain
333 because of high variability in NEE (Bailey 2006) and CH_4 flux data (this study). Furthermore in
334 this analysis we must assume that CH_4 emissions and NEE will remain constant over many
335 decades. In reality NEE is likely to be dynamic over at least several decades of succession
336 (Odum 1969). Therefore it would take a long-term monitoring approach to improve certainty of
337 radiative forcing switchover time for the CCW. Despite these shortcomings we may conclude
338 that CCW has a relatively short radiative forcing switchover time due to its low CH_4 flux. The
339 CO_2 sequestration: CH_4 -flux ratio of CCW ranges from 96 to 20, corresponding to a radiative
340 forcing switchover time range of 0 to approximately 200 years following Neubauer's (2014)
341 model. CCW will likely become a net GHG sink more quickly than at least six out of eight
342 wetlands analyzed by Neubauer (2014).

343 **Conclusions**

344 We found little evidence to suggest that added composted yard waste increases CH₄ or
345 N₂O emissions from CCW a decade after restoration. CH₄ emissions are only significant when
346 soils are warm and water levels and soil moisture are high. Even when CH₄ flux is at its greatest
347 magnitude, it still represents a relatively modest contribution to global warming potential
348 compared to soil CO₂ flux.

349 Yet even if CCW were to produce no CH₄, it would still be a net CO₂ source at high OM
350 loading rates because of negative NEE (Bailey 2006), at least until the excess OM is respired.
351 Therefore we recommend that only moderate levels of OM need to be added to created wetlands.
352 Adding more than ~160 Mg ha⁻¹ does not improve soil geochemistry (Bruland and Richardson
353 2009) and excess OM simply decomposes while adding little in the way of tangible productivity
354 increases, not to mention incurring greater material transport and associated construction costs.

355

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454

455

456

457 **Tables**

458 Table 1. Summary of linear regression and ANOVA tests for differences and trends in log-
459 transformed carbon dioxide (CO₂) emissions between and across gradient of plots treated with

460 different levels of organic matter (OM) at the Charles City Wetland in Charles City County,
461 Virginia. Values that meet $p < 0.05$ are bolded

Month	linear regression		ANOVA
	p-value	r-squared	p-value
May	0.028	0.24	0.165
July	<0.001	0.55	0.018
Sept.	0.133	0.12	0.116
Nov.	0.009	0.40	0.126
Jan.	0.043	0.21	0.003

462

463 Table 2. Summary of monthly averages (\pm SD) soil temperature (at 5 cm depth), hydrology and
464 soil carbon emissions from the Charles City Wetland in Charles City County, Virginia. All data
465 collected in 2012 except for January, 2013

Month	Soil Temp.	Water Level	Soil volumetric	CH ₄	CO ₂
	$^{\circ}$ C	cm	water content %	emissions	emissions
				$\text{mg}\cdot\text{m}^{-2}\cdot\text{hr}^{-1}$	
May	18.1 \pm 0.7	5.3 \pm 2.8	65.6 \pm 7.5	0.82 \pm 0.88	117 \pm 94
July	24.6 \pm 0.3	-39.9 \pm 28.6	29.9 \pm 11.5	0.02 \pm 0.04	595 \pm 191
Sept.	20.0 \pm 1.5	6.0 \pm 2.3	60.2 \pm 6.4	1.29 \pm 1.41	188 \pm 130
Nov.	10.1 \pm 0.4	3.0 \pm 2.3	55.5 \pm 6.7	0.02 \pm 0.04	124 \pm 63
Jan.	5.9 \pm 0.8	9.2 \pm 2.8	60.5 \pm 6.2	0	32 \pm 27

466

467 Table 3. Review of methane (CH₄) emissions rates in $\text{kg CH}_4\text{-C ha}^{-1} \text{ yr}^{-1}$ from natural and
468 restored forested wetlands of the Southeastern United States.

CH ₄ flux	Location	Type	Reference
554	Newport News Swamp, Va.	Natural	(Wilson et al 1989)
427	Newport News Swamp, Va.	Natural	(Wilson et al 1989)
311	Ogeechee River, Ga. (west)	Natural	(Pulliam 1993)
297	Okeefenokee Swamp, Ga.	Natural	(Flebbe 1982)
262	Creeping Swamp, NC	Natural	(Mulholland 1981)
107	Timberlake Restoration Preserve, NC	Restored	(Morse et al 2012)
92	Ogeechee River, Ga. (east)	Natural	(Pulliam 1993)
72	Palmetto Peartree Preserve, NC	Natural	(Morse et al 2012)
41	Charles City Wetland, Va.	Restored	This study
14	Timberlake Restoration Preserve, NC	Restored	(Morse et al 2012)
0.5	Timberlake Restoration Preserve, NC	Restored	(Morse et al 2012)

469

470 Table 4. Comparison of microtopographic and growing season hydrologic conditions at the
 471 Charles City Wetland in Charles City County, Virginia between 2005 (Bailey et al. 2007) and
 472 2012 (this study)

Year	Rainfall (Apr. - Oct.; cm)*		Elevation across OM loading rates		
	total	depart. from normal	range (cm)	lin. reg. r-squared	lin. reg. p-value
2005	69	-7.5	11	0.55	<0.001
2012	82	+5.7	6	0.17	<.05

473 *Source: National Climatic Data Center (Lawrimore et al. 2011)

474

475 **Figure captions**

476 **Fig. 1** Location of Charles City Wetland in Charles City County, Virginia, USA, with A)
 477 showing the geographic location of Charles City County, Virginia, B) showing the siting of the
 478 experimental block within the wetland, and C) indicating the arrangement of plots, treatments
 479 and wells (labeled a through e) within the block, and site dimensions

480 **Fig. 2** Illustrations of the reduced-disturbance static chamber design: A) Photograph of chamber
 481 being deployed in the Charles City Wetland in Charles City County, Virginia; B) schematic of
 482 chamber disassembled to reveal water fillable gutter on rim of collar that creates an air tight seal,
 483 internal fan to mix headspace air and thermocouple to monitor internal chamber temperature; C)
 484 schematic of chamber assembled

485 **Fig. 3** Water level as recorded by five 1.5 meter Odyssey loggers (Dataflow Systems,
 486 Christchurch, New Zealand) placed in water level wells (W1 through W5) at the Charles City
 487 Wetland in Charles City County, Virginia, USA from 22 February, 2012 to 21 January, 2013.
 488 Positive values indicate standing water. Overlaid precipitation data is from a station in nearby
 489 James City County, Virginia (National Climate Data Center)

490 **Fig. 4** Linear regressions of mean (\pm SE) total carbon in top 10 cm of soil across organic matter
 491 amendment plots at the Charles City Wetland in Charles City County, Virginia, USA. 2005 data
 492 from Bailey *et al* (2007.)

493 **Fig. 5** Depth profiles of mean (\pm SE): A) percent soil carbon by mass, B) percent soil nitrogen by
 494 mass, C) total soil phosphorus by mass in $\mu\text{g g}^{-1}$, D) extractable ammonia/ammonium in $\mu\text{g NH}_x\text{-}$
 495 N g^{-1} dry soil, E) extractable nitrate/nitrite in $\mu\text{g NO}_x\text{-N g}^{-1}$ dry soil, F) extractable phosphorus in
 496 $\mu\text{g NO}_x\text{-N g}^{-1}$ dry soil. Different dash patterns represent loading rates of organic matter in Mg
 497 ha^{-1} added to the Charles City Wetland in Charles City County, Virginia, USA

498 **Fig. 6** Relationships between total soil elemental content and extractable nutrients by depth, as
 499 indicated by shapes, at the Charles City Wetland in Charles City County, Virginia, USA. A)

500 shows a linear relationship between percent total soil carbon by mass and percent total soil
501 nitrogen by mass, B) a logarithmic relationship between percent total soil carbon by mass and
502 total soil phosphorus by mass in ppm, C) a linear relationship between percent total nitrogen of
503 soil by mass and extractable ammonia/ammonium in $\mu\text{g NH}_x\text{-N g}^{-1}$ dry soil, D) a quadratic
504 relationship between total soil phosphorus by mass in $\mu\text{g.g}^{-1}$ and extractable phosphorus in $\mu\text{g g}^{-1}$
505 dry soil

506 **Fig. 7** Mean carbon dioxide flux as a function of soil temperature at 5 cm depth from the organic
507 matter experimental plots at the Charles City Wetland in Charles City County, Virginia across
508 eight sampling dates from November, 2011 to January, 2013. Error bars represent ± 1 standard
509 deviation

510 **Fig. 8** Mean ($\pm\text{SE}$) carbon dioxide flux from the organic matter experimental plots at the Charles
511 City Wetland in Charles City County, Virginia across nine sampling dates from September, 2011
512 to January, 2013. Different dash patterns represent loading rates of organic matter in Mg ha^{-1}

513 **Fig. 9** Actual carbon dioxide flux compared to linear model predictions at the Charles City
514 County Wetland in Charles City County, Virginia for: A) data across five sampling dates from
515 May 2012 to January 2013 and multiple regression predictions based on soil temperature (5 cm
516 depth), soil volumetric water content, and total soil carbon (top 5 cm); and B) data from 22 July
517 2012 and linear predictions based on total soil carbon (top 5 cm) and total soil nitrogen at 20 cm
518 depth. Dashed curves represent 95 percent confidence intervals for the regression line. Dashed
519 horizontal line indicates mean carbon dioxide flux value

520 **Fig. 10** Methane flux (CH_4) rates as a function of soil volumetric water content measured from
521 the organic matter experimental plots at the Charles City Wetland in Charles City County across
522 five sampling dates from May 2012 to January 2013

523 **Fig. 11** Carbon dioxide (CO_2) and methane (CH_4) flux from soil across five levels of organic
524 matter loading rates estimated from sampling on 7 May and 26 September, 2012 at the Charles
525 City Wetland in Charles City County, Virginia, USA. Note: CH_4 was converted to CO_2 -
526 equivalents by multiplying by 38—its 100-year sustained global warming potential following
527 Neubauer (2014). Error bars represent standard errors of the mean

Figures (color) for Peterson Report

Fig. 1 (174 mm)

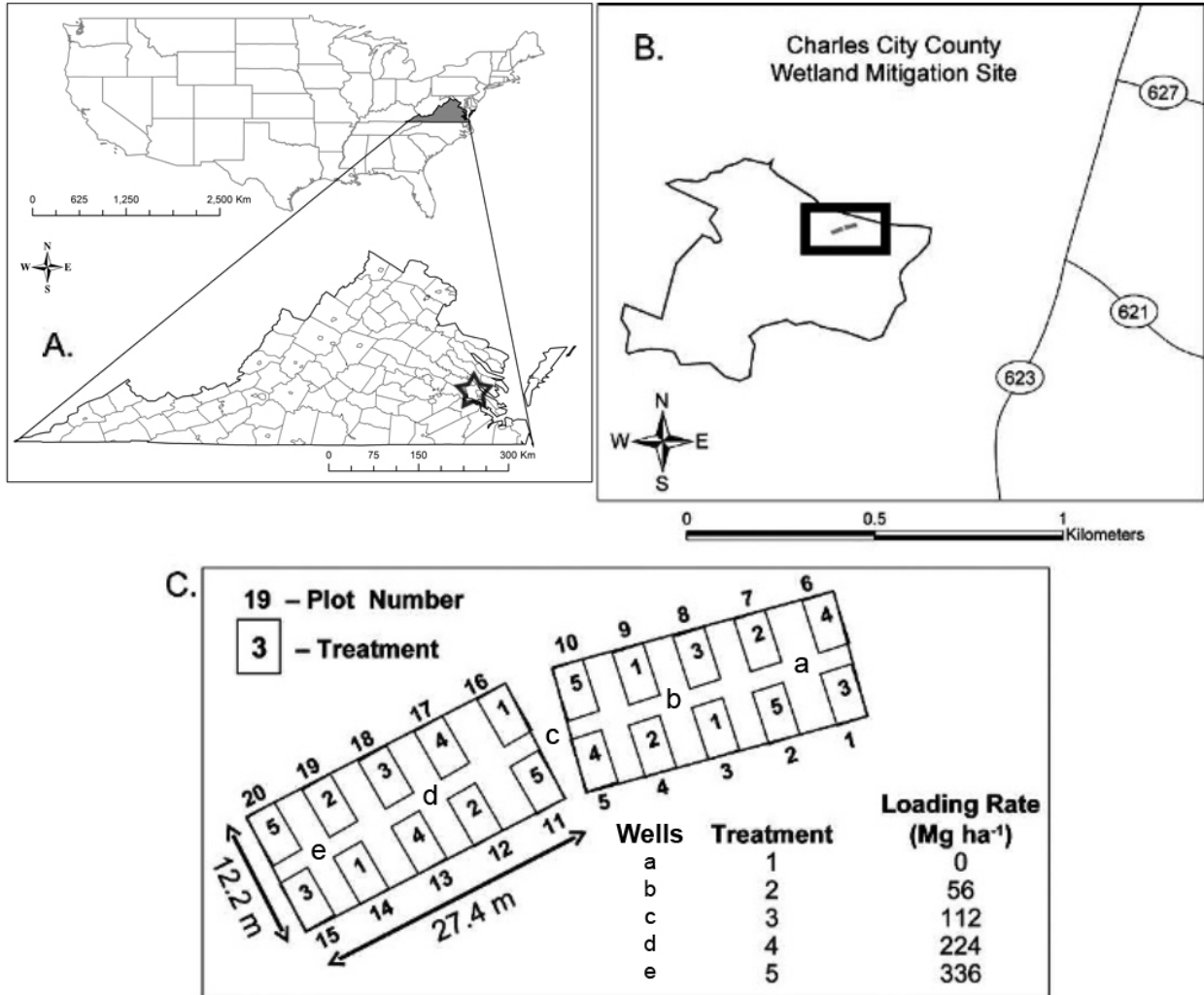


Fig. 2 (129 mm wide)

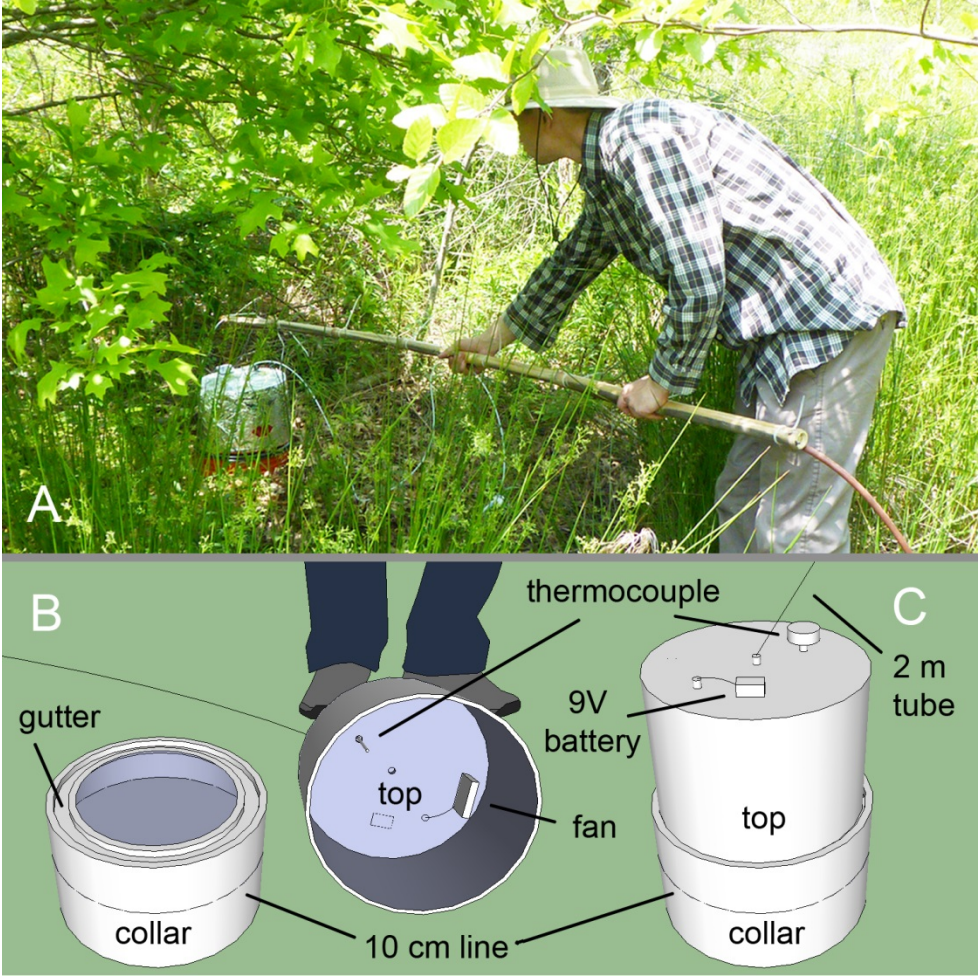


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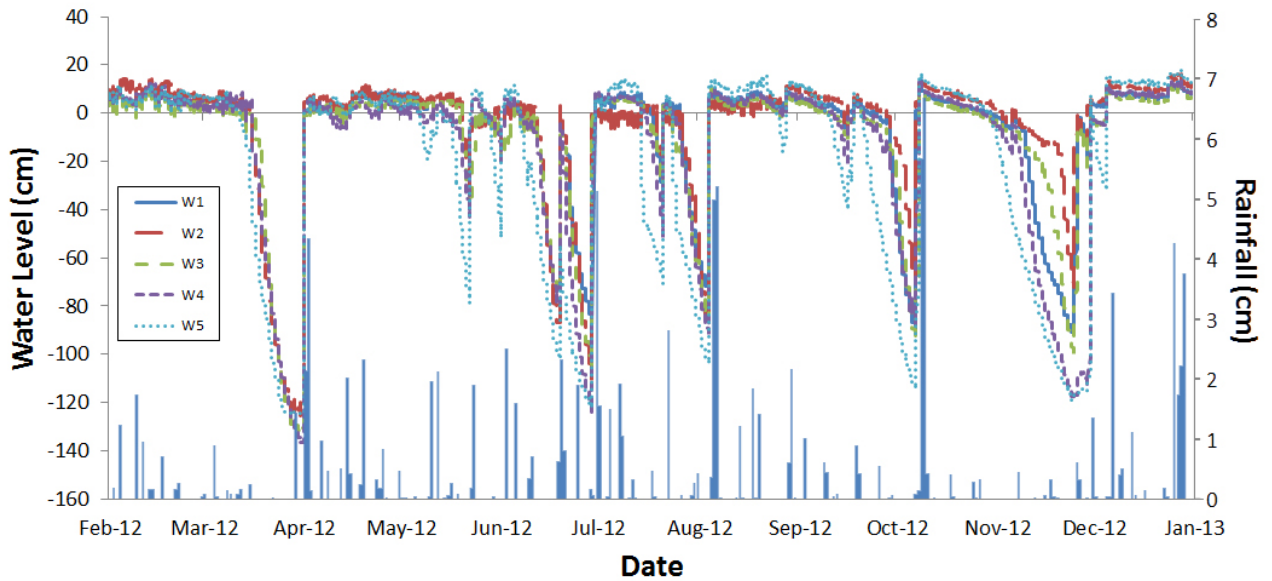


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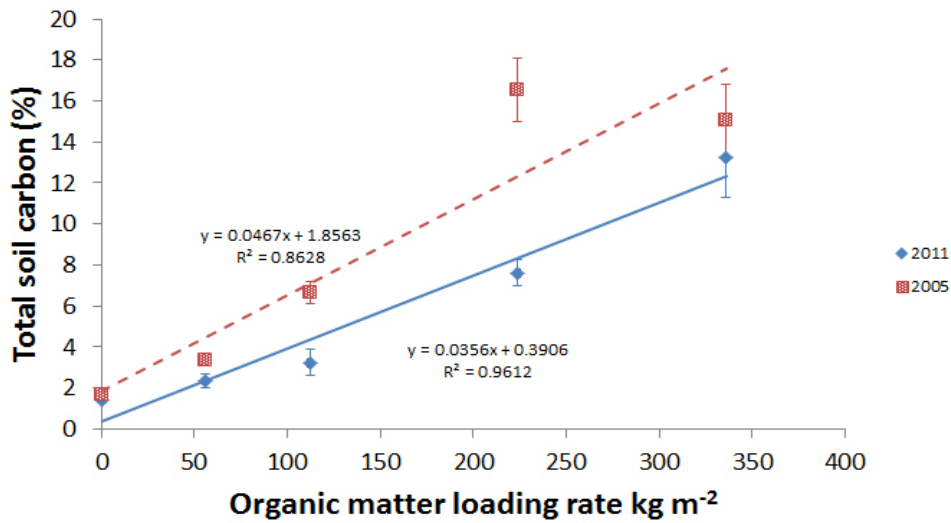


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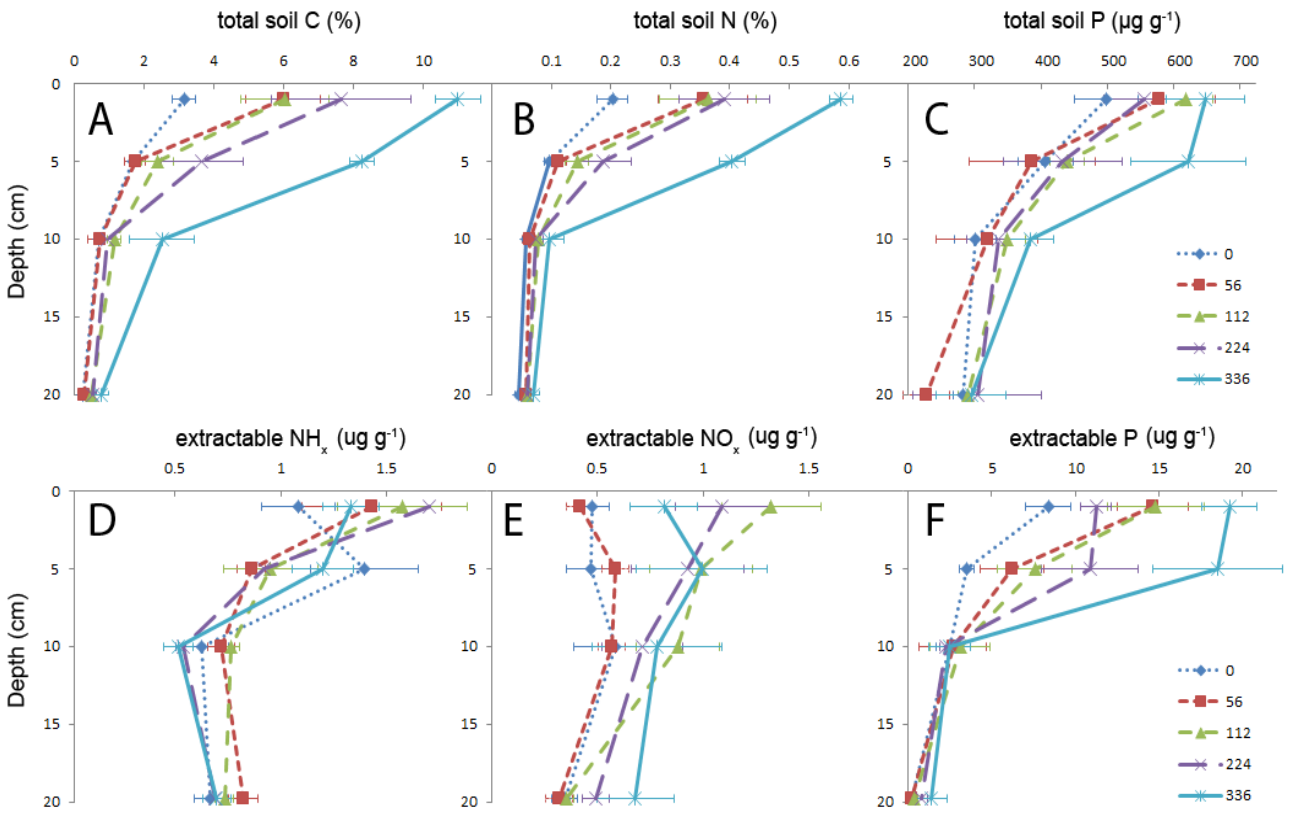


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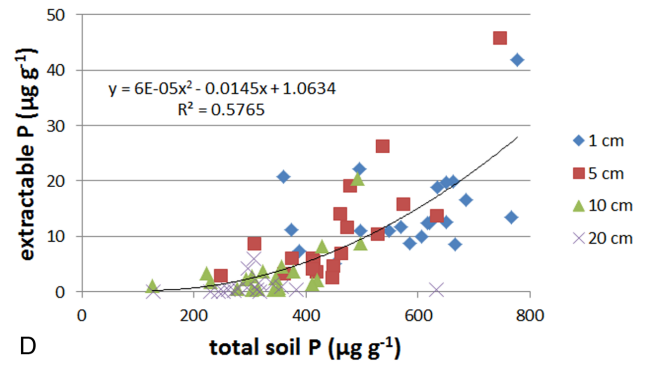
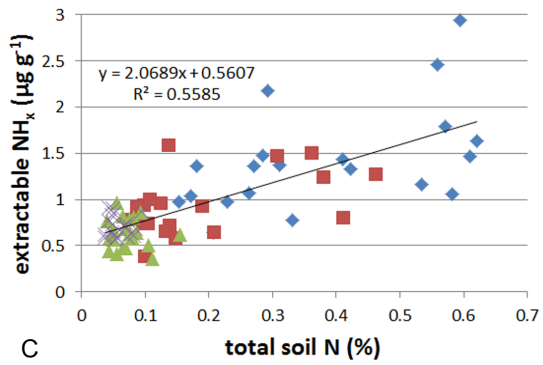
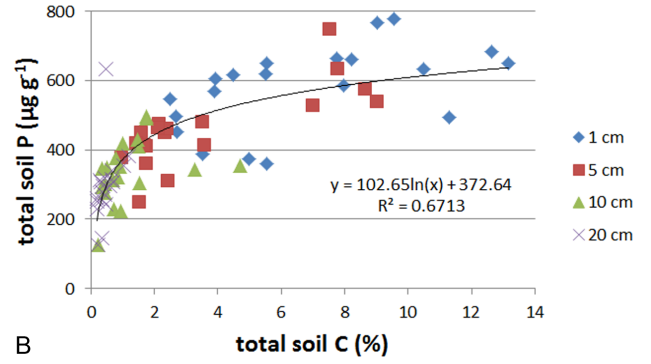
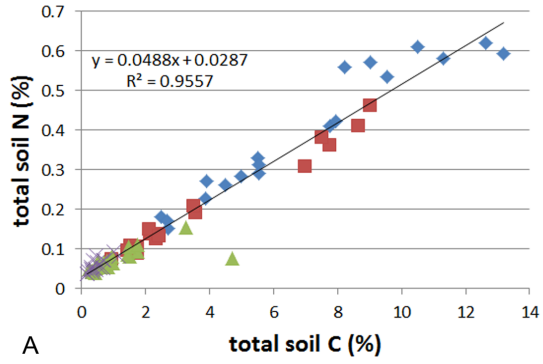


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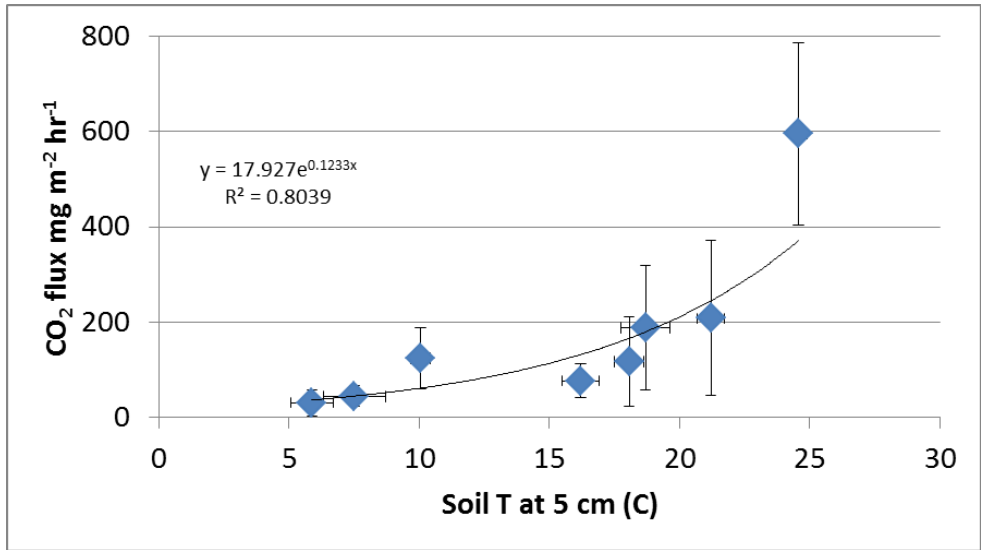


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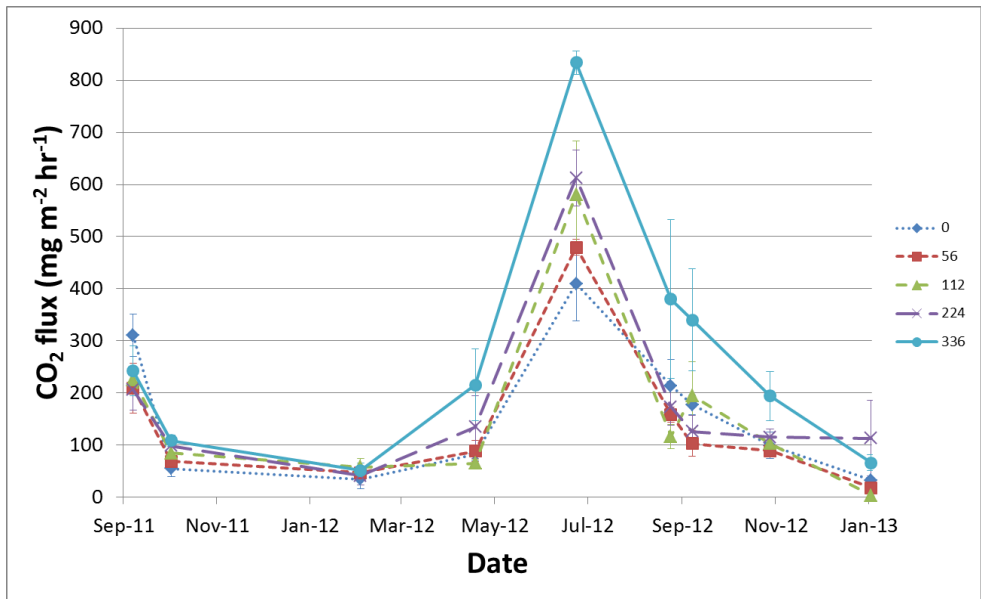


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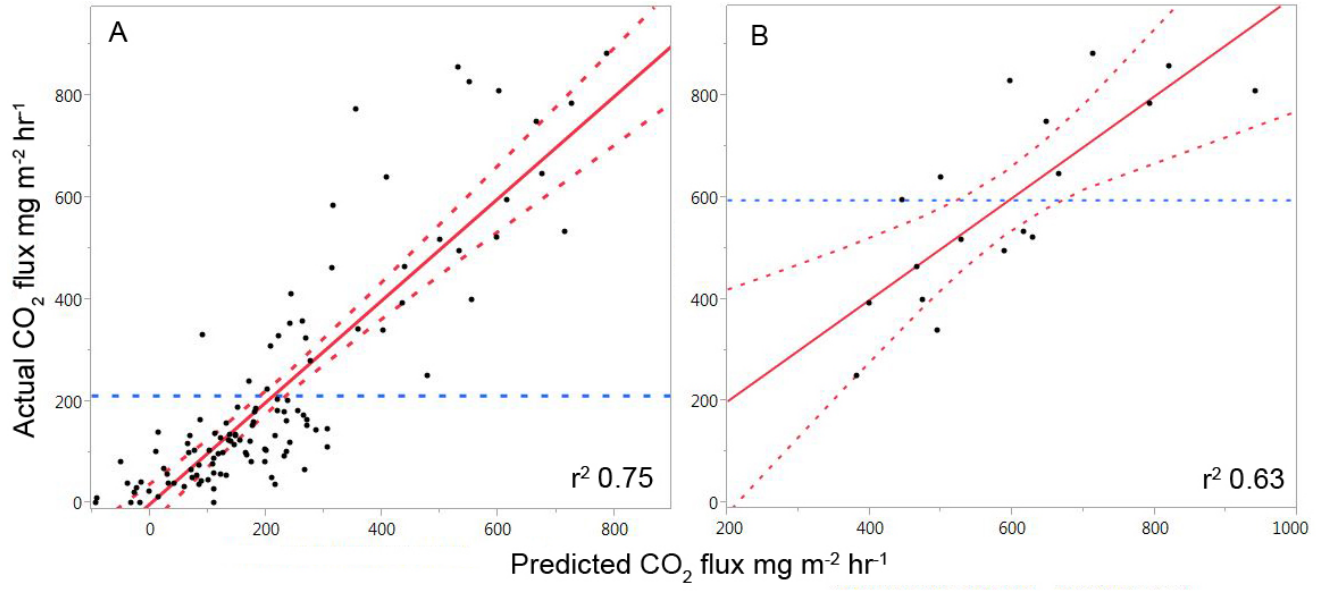


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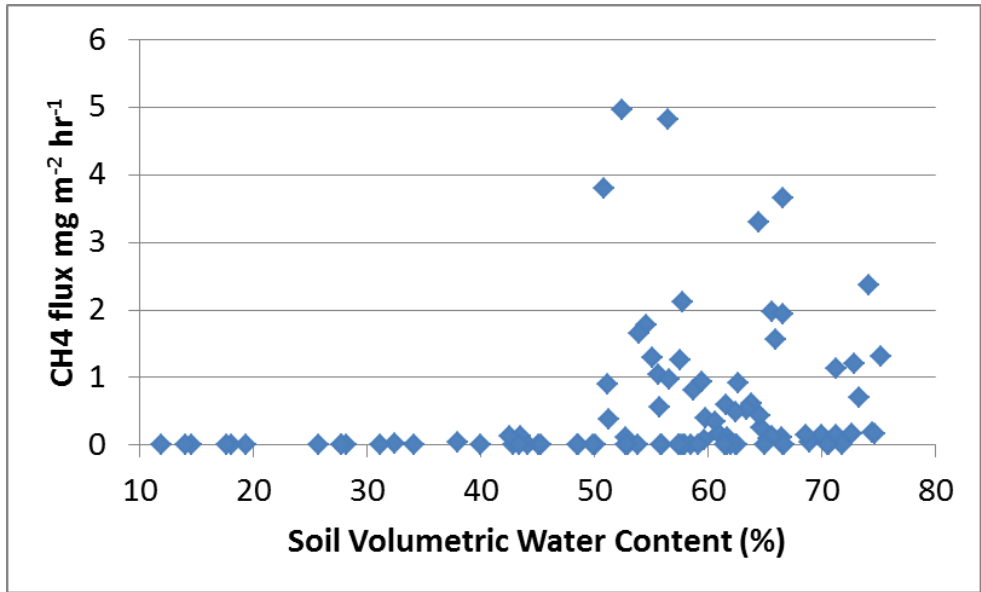


Fig. 11 (174 mm)

