



New Constraints on Methane Fluxes and Rates of Anaerobic Methane Oxidation in a Gulf of Mexico Brine Pool via In Situ Mass Spectrometry

Citation

Wankel, Scott D., Samantha B. Joye, Vladimir A. Samarkin, Sunita Rajesh Shah, Gernot Friederich, John Melas-Kyriazi, and Peter R. Girguis. 2010. New constraints on methane fluxes and rates of anaerobic methane oxidation in a Gulf of Mexico brine pool via in situ mass spectrometry. *Deep Sea Research Part II: Topical Studies in Oceanography* 57(21-23): 2022-2029.

Published Version

doi:10.1016/j.dsr2.2010.05.009

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1 **New constraints on methane fluxes and rates of anaerobic**
2 **methane oxidation in a Gulf of Mexico brine pool via *in situ***
3 **mass spectrometry**

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45

46 **Abstract**

47 Deep sea biogeochemical cycles are, in general, poorly understood due to the
48 difficulties of making measurements *in situ*, recovering samples with minimal
49 perturbation and, in many cases, coping with high spatial and temporal heterogeneity. In
50 particular, biogeochemical fluxes of volatiles such as methane remain largely
51 unconstrained due to the difficulties of accurate quantification *in situ* and the patchiness
52 of point sources such as seeps and brine pools. To better constrain biogeochemical fluxes
53 and cycling, we have developed a deep sea *in situ* mass spectrometer (ISMS) to enable
54 high-resolution quantification of volatiles *in situ*. Here we report direct measurements of
55 methane concentrations made in a Gulf of Mexico brine pool located at a depth of over
56 2300m. Concentrations of up to 33mM methane were observed within the brine pool,
57 while concentrations in the water directly above were three orders of magnitude lower.
58 These direct measurements enable the first accurate estimates of the diffusive flux from a
59 brine pool, calculated to be $1.1 \pm 0.2 \text{ mol m}^{-2} \text{ yr}^{-1}$. Integrated rate measurements of
60 aerobic methane oxidation in the water column overlying the brine pool were $\sim 320 \text{ } \mu\text{mol}$
61 $\text{m}^{-2} \text{ yr}^{-1}$, accounting at most for just 0.03% of the diffusive methane flux from the brine
62 pool. Calculated rates of anaerobic methane oxidation were 600 to 1200 $\mu\text{M yr}^{-1}$, one to
63 two orders of magnitude higher than previously published values of AOM in anoxic
64 fluids. These findings suggest that brine pools are enormous point sources of methane in
65 the deep sea, and may, in aggregate, have a pronounced impact on the global marine
66 methane cycle.

67

67 **Introduction**

68 **1.1. Global importance of methane**

69 The marine methane cycle has been the subject of much investigation in recent
70 years, in large part due to burgeoning interest and concern over deep ocean methane
71 hydrates. The deep ocean methane reservoir represents an enormous and dynamic pool
72 of carbon likely exceeding reserves of conventional oil and gas (Collett and Kuuskraa,
73 1998). In deep ocean regions, characterized by low temperatures, high pressure and
74 sufficient methane concentration, methane exists largely in the solid form of a gas
75 hydrate (Kvenvolden, 1993). Methane seeps and associated gas hydrates have been
76 identified along many passive and active continental margins (Kvenvolden and Lorensen,
77 2008). Because the destabilization of hydrates is sensitive to increases in temperature or
78 decreases in pressure, it has been postulated that increases in mean global temperatures
79 might trigger a release of methane into the ocean and atmosphere. A significant release
80 of methane into the atmosphere could ultimately lead to a catastrophic greenhouse effect;
81 this mechanism has been invoked as an explanation for past deglaciation events (Dickens,
82 2003; Sloan *et al.*, 1992; Zachos *et al.*, 2001).

83 Despite recent, numerous studies of methane hydrates, modern fluxes of methane
84 from the deep sea into surface waters and ultimately the atmosphere are very poorly
85 constrained. Estimates of methane flux have been aided, to some degree, by recent
86 advances in our understanding of marine microbiological influences on the global
87 methane cycle. Aspects of the marine methane cycle remain largely unconstrained due to
88 limitations in methods and technologies that enable accurate assessment of methane

89 concentration and flux – as well as rates of biological methanogenesis or methanotrophy.
90 Pressure and temperature have a pronounced effect on methane solubility. As such,
91 upon retrieval of methane-saturated waters or hydrate-rich sediments from the deep
92 ocean, methane rapidly outgasses to the atmosphere. Thus, it has been challenging to
93 constrain flux and microbial activity *in situ*, under environmentally relevant conditions.
94 Because previous data have shown that methane oxidation, both aerobic and anaerobic,
95 are the largest methane sinks in marine environments (Reeburgh, 2007), understanding
96 what controls methane oxidation -including concentration and abiotic flux- are paramount
97 to understanding global methane dynamics.

98 To better constrain the methane flux in chemically reducing environments – and
99 ultimately to quantify the influence of biotic and abiotic processes on the methane cycle –
100 we employed a newly developed *in situ* mass spectrometer (ISMS) to conduct direct
101 measurements of methane concentration which - in concert with shipboard
102 microbiological measurements - were used to generate more robust estimates of diffusive
103 flux and net methane oxidation rates in a newly discovered brine pool in the Gulf of
104 Mexico.

105 **2. Geologic Setting**

106 Along the continental shelf in the Gulf of Mexico, massive reservoirs of liquid
107 and gaseous hydrocarbons lie buried beneath kilometers of sediment accumulated from
108 the Mississippi River drainage basin. Due to compression and dewatering of the
109 overlying sediments, underlying evaporite deposits have undergone plastic deformation
110 resulting in salt-diapir driven tectonic activity (Kennicutt *et al.*, 1988). The resulting
111 system of fractures and faults provides conduits for the emission of hydrocarbons to the

112 seafloor via seepage (Roberts and Carney, 1997). Hydrocarbon seeps are often
113 characterized by abundant chemosynthetic based macro- and microfaunal communities
114 including tubeworms, mussel beds, and bacterial mats which thrive on the reduced
115 organic compounds emanating from below (Fisher *et al.*, 2007; MacDonald *et al.*, 1990;
116 MacDonald *et al.*, 2003). In addition, hypersaline brine fluids seep from the seafloor in
117 many locations (MacDonald *et al.* 1990; Joye *et al.* 2005, 2009). Previous studies have
118 provided insight into the geochemical composition of these brine pools, though to date
119 volatile flux and net rates of methane oxidation remain poorly constrained due to the
120 challenges in quantification resulting from off-gassing (at *in situ* pressures and
121 temperatures relevant here, methane saturation is $\sim 174 \text{ mmol kg}^{-1}$ (Duan and Mao,
122 2006)). Accurate sampling of fluids with high gas content using conventional methods
123 (e.g., Niskin Bottles) has thus proven impractical for volatiles.

124 The brine pool characterized in this study (AC601) is located in the Alaminos
125 Canyon lease block 601 (26° 23.53 N; 94° 30.85 W; Roberts *et al.* this issue; (Roberts *et*
126 *al.*, 2007). The brine pool is estimated to be $\sim 250\text{m}$ in diameter and approximately 2334
127 meters below sea surface. This brine pool was visited during expeditions on board the
128 *RV Ronald H. Brown* using the *DSV Jason II* during expeditions from May 6 through
129 June 4, 2006 and June 3 through July 6, 2007 (see Roberts *et al.*, this issue), for further
130 description of the expeditions and site locations). Discrete geochemical measurements of
131 this brine pool were made during the 2006 and 2007 expedition, while deployment of the
132 ISMS was carried out during the 2007 expedition.

133 **3. Methods**

134 **3.1. *Fundamentals of Membrane Inlet Mass Spectrometry/ISMS***

135 Over the past five decades, the use of membrane inlet mass spectrometry (MIMS)
136 has proven to be a powerful tool for measuring complex mixtures of dissolved gases in
137 both industrial and laboratory settings (Johnson *et al.*, 2000; Ketola *et al.*, 2002). MIMS
138 represents an optimal technique for mixed environmental gas analysis, having a high
139 degree of sensitivity and precision, with minimal sample perturbation (Kana *et al.*, 1994).
140 MIMS has been used over the past several decades to measure and monitor a wide range
141 of dissolved gases in aquatic and terrestrial environments, including studies of bacterial
142 mats, peat bogs, estuarine sediments, forest soils, and tree canopies to name a few
143 (Hemond, 1991; Kana *et al.*, 1998; Lloyd *et al.*, 1986; Lloyd *et al.*, 1998). MIMS has
144 also been used to study metabolite flux during shipboard high-pressure experiments (e.g.,
145 (Girguis *et al.*, 2002; Girguis *et al.*, 2000). MIMS has also emerged as an important tool
146 for analyzing dissolved gases in seawater, (e.g. dissolved gases in surface waters
147 analyzed continuously shipboard;(Kaiser *et al.*, 2005; Tortell, 2005a; Tortell, 2005b) and
148 more recently for *in situ* marine surface waters (Bell *et al.*, 2007; Camilli and Hemond,
149 2004; Kaiser *et al.*, 2005; Tortell, 2005b, c).

150 The recent adaptation of MIMS to *in situ* environmental analyses demonstrates
151 the utility of such an instrument operating while underway at sea – allowing the continual
152 monitoring of many gas species in real time. This approach allows highly accurate
153 monitoring of spatially explicit biogeochemical changes. For example, changes in O₂/Ar
154 indicate changes in net community production in ocean surface waters in different
155 regions of the eastern equatorial Pacific (Kaiser *et al.*, 2005). Additionally, N₂/Ar has

156 been used to identify areas of active denitrification (e.g., N₂ production) in seasonally
157 oxygen-depleted bottom waters via MIMS in Saanich Inlet (Tortell, 2005b). Underway
158 shipboard trace gas analysis has also shed light on dynamics of dimethylsulfide (Tortell,
159 2005a).

160 Given the apparent utility of real-time quantification by MIMS, we aimed to
161 develop a MIMS that would achieve comparable performance in waters deeper than 1000
162 meters. Currently, investigation of deep water samples still generally requires the
163 collection of individual samples and shipboard analysis (e.g., Tortell, 2005b), risking
164 contamination and/or degassing. Furthermore, individual sample collection and analysis
165 can often result in delays between sampling and data interpretation. Our understanding
166 of deep-sea biogeochemistry would greatly benefit from real-time, *in situ* dissolved gas
167 analysis. Here we present results from a real-time *in situ* membrane inlet mass
168 spectrometer designed to A) operate at depths up to 450 atmospheres of pressure, B)
169 provide real-time data to the user (when used on human occupied submersibles or
170 remotely operated vehicles), and C) enable sampling with high spatial and temporal
171 resolution using an *in situ* pumping system.

172 **3.2. *ISMS Design and Calibration***

173 This ISMS consisted of three primary sub-systems: 1) a high-pressure membrane
174 inlet (Fig. 1a) with a small volume seawater pumping system; 2) a quadrupole mass
175 spectrometer (Fig. 1e, f) and oil-less vacuum pumping system (Fig. 1 d, g); and 3) an
176 underwater housing (either a 2000 meter-rated aluminum 3300 alloy housing, or a 4500
177 meter rated 6AL-4V titanium housing, both of which are approximately 120 cm in length
178 and 24 cm in diameter). The membrane inlet assembly consisted of a circular sheet

179 (0.625 in. diameter) of polydimethylsiloxane (PDMS) membrane structurally backed by
180 an integrated woven fiber (Franatech, Germany). This pliable membrane material was
181 supported by a sintered stainless steel frit (5 μ m pore size, Applied Porous Materials,
182 Tariffville, CT), which was in turn supported by the titanium body of the inlet housing.
183 Sample water was pumped through the inlet housing assembly (2ml internal volume) at a
184 flow rate of \sim 3ml/min using a small solenoid pump (The Lee Company, Westbrook, CT),
185 which was controlled by an adjustable timing circuit located inside of the pressure
186 housing.

187 The membrane assembly was connected to a Stanford Research Systems Residual
188 Gas Analyzer (SRS RGA100) via standard vacuum flanges. Within the vacuum system,
189 a pressure of \sim 10e⁻⁵ Torr was maintained by a turbo-molecular pump (model: ATH 31+;
190 Alcatel, France) backed by a diaphragm roughing pump (model: ANDC83.4; KNF-
191 Neuberger, Trenton, NJ). Open source electron impact ionization was carried out with a
192 thoriated iridium wire filament. The mass spectrometer and pumps were protected from
193 membrane failure by a high-pressure / high-vacuum solenoid valve (Circle Seal, Inc.,
194 Corona, CA), which is actuated upon intrusion of water. The entire mass spectrometer
195 assembly was housed in one of the aforementioned housings. 24 VDC power and two
196 independent RS-232 channels (for serial communications with the turbo pump control
197 board and continuous feedback from the RGA analyzer) were supplied via a wet-connect
198 underwater cable (SubConn, Inc., North Pembroke, MA). In this configuration, real time
199 monitoring of fluid chemistry is achievable during submersible or ROV operations,
200 which allows for informed site selection for fluid measurements as well as adaptive
201 sampling of biological specimens.

202 To conduct the benchtop high-pressure calibrations (Fig. 2), model 110A HPLC
203 pumps (Beckman-Coulter, Fullerton, CA) were used to deliver calibrated solutions past
204 the membrane surface at various flow rates and pressures. Hydrostatic pressure against
205 the membrane inlet was manually controlled with a backpressure valve (StraVal Valve,
206 Garfield, NJ) and monitored with high pressure gauges. Various calibration solutions
207 were used including air-sparged DI water or seawater, degassed distilled water and/or
208 seawater equilibrated with gas mixtures of interest (e.g., CH₄), including the use of a high
209 pressure equilibration system for generating very high CH₄ concentrations (Fig. 2). A
210 gas chromatograph (HP 5890 Series II plus with a TCD) outfitted with a custom gas
211 extractor (Childress *et al.*, 1984) designed for quantification of rapidly degassed seawater
212 samples was used to obtain independent analyses of dissolved methane concentrations,
213 while previously described equations of state were used to calculate dissolved methane
214 concentrations at very high pressures (Duan and Mao, 2006) (Fig. 2b). During lab
215 experiments, relative changes in signal intensity were proportional to changes in the
216 permeation of gas through the membrane (either due to changes in permeate
217 concentration or changes in the permeability coefficient). We and others have observed
218 that changes in hydrostatic pressure can have an influence on permeation of gases
219 through membrane materials, in particular PDMS, interpreted to be caused by
220 compression of the membrane pore space through which analyte gas passes (Bell *et al.*,
221 2007). A change in the relationship between dissolved gas concentration and signal
222 intensity was observed during large changes in hydrostatic pressure (Fig. 2a). To account
223 for this response, we conducted calibrations using methane dissolved in seawater over a
224 range of *in situ* pressures and used these results to develop an empirical correction as

225 previously described (Bell et al., 2007). While this approach corrects for implicit
226 changes in membrane behavior, it should be noted that the ISMS dataset presented here is
227 comprised entirely of fluids sampled at a relatively constant depth ($\sim 2330\text{m} \pm 2$ (233
228 bar)) and temperature and as such effects due to differential pressure or temperature
229 among the samples collected were negligible. Based on benchtop calibrations, the
230 accuracy of the ISMS methane concentrations in the configuration described here was \pm
231 11%, primarily due to the correction required for the pressure effects on the PDMS
232 membrane (accuracy is improved through the use of alternate membrane material (e.g.
233 Teflon AF)). Notably, however, the precision of the ISMS measurements is much better
234 than this and, based on benchtop calibrations, is within $\pm 1\%$.

235 **3.3. *Water Column Methane and Oxygen Concentration***

236 To determine brine pool and seawater column methane concentrations, a CTD
237 rosette was lowered into the brine pool, using sonar to identify the brine pool as the
238 rosette approached the bottom. Niskin bottles were tripped during descent to prevent
239 contamination of bottles as gas came out of solution during the rosette's ascent. Two
240 bottles were tripped in the brine pool itself, while two were tripped 1 meter above the
241 brine-seawater interface (the interface was confirmed by the real-time conductivity
242 signal). After securing the rosette on deck, water samples were immediately transferred
243 to 1-liter PET-G bottles using gas-impermeable tubing. Bottles that were tripped in the
244 brine were substantially over-pressured and not suitable for gas quantification (though
245 samples were transferred to the PET-G bottles for rate measurements). A second sample
246 was transferred to a 250 mL BOD bottle for determination of dissolved oxygen using a
247 high sensitivity Orion[®] oxygen electrode. Methane was extracted using an adaptation of

248 the sonication/vacuum extraction technique (Suess *et al.*, 1999) followed by gas
249 chromatography for quantification. Prior to dissolved gas extraction, samples were
250 stored at bottom water temperature (4 °C). Two individual samples were analyzed from
251 each rosette bottle.

252 **3.4. ISMS Deployments and Determining Brine Pool Methane Concentrations**

253 Upon reaching the brine pool, the submersible took care not to disturb the brine-
254 seawater interface. Using the ROV manipulator, the ISMS sample inlet was positioned
255 and held in place until the ISMS response reached steady state from which *in situ*
256 concentrations of CH₄ were calculated. A total of five independent sets of measurements
257 were made, beginning with two just above the brine fluid and three at depths of
258 approximately 5, 20 and 80 cm into the fluid (Figs. 3 and 4).

259 **3.5. Methane Oxidation Rates**

260 Aerobic methane oxidation occurs according to the following stoichiometry:



262 Accordingly, aerobic methane oxidation rates were determined by incubating samples
263 with C³H₄ and tracking the production of ³H₂O (Carini *et al.*, 2005; Valentine *et al.*,
264 2001). Typically, triplicate live and dead (Hg killed; i.e. samples were amended with
265 HgCl₂ to arrest all biological activity) samples from each depth were incubated for 36
266 hours at *in situ* temperatures. Unreacted C³H₄ tracer was removed by purging samples
267 with water-saturated CH₄ and the oxidation product, ³H₂O, was quantified by liquid
268 scintillation counting (Carini *et al.*, 2005).

269 In general, marine anaerobic methane oxidation in hydrocarbon seeps and brine
270 pools is coupled to sulfate reduction, with the net reaction:



272 Anaerobic methane oxidation rates were also determined by incubating samples with
273 $^{14}\text{CH}_4$ and tracking the production of $^{14}\text{CO}_2$ (as in Joye *et al.*, 1999; Valentine *et al.*,
274 2001). Triplicate live and dead (Hg killed) samples from the surface (~20 cm) and sub-
275 surface (~100 cm) brine were incubated for 48 hours at *in situ* temperatures. Unreacted
276 $^{14}\text{CH}_4$ tracer was removed by purging with water-saturated CH_4 and the $^{14}\text{CO}_2$ oxidation
277 product was quantified following acid extraction and trapping on a phethylamine wick,
278 followed by liquid scintillation counting (Carini *et al.*, 2005).

279 3.6. *Sulfate Reduction Rates*

280 Samples for sulfate reduction were collected into gas tight glass tubes, amended
281 with radiotracer ($^{35}\text{SO}_4^{2-}$) and incubated for 24 hours (as in Joye *et al.*, 2004; Orcutt *et al.*,
282 2005). For each depth, triplicate samples were incubated along side controls (killed at
283 time zero). After incubation, samples were transferred from the tubes to 50 ml centrifuge
284 tubes and mixed with 20% zinc acetate. Samples were processed and rates calculated as
285 presented in Orcutt *et al.* (2005).

286 3.7. *Major Ion chemistry*

287 Concentrations of major ions (SO_4^{2-} , Cl^- , dissolved inorganic carbon (DIC),
288 dissolved organic matter (DOC and DON) and dissolved inorganic nitrogen (NH_4^+ , NO_2^-
289 and NO_3^-) were determined using previously reported methods (see (Joye *et al.*, 2004)
290 and Joye *et al.* this volume and references therein).

291 4. Results

292 4.1. *General geochemical composition of brine pool AC601*

293 Geochemical data on the waters collected from the brine pool are given in Table
294 1. The waters of the brine pool were anoxic ($O_2 < 2\mu\text{M}$) with a pH of ~ 6.3 . Salinities
295 were substantially elevated above seawater at 82 and 92 for the 20 and 100 cm depths,
296 respectively, with chloride measuring 1366 and 1533 mM at each depth. Water from
297 both depths was highly enriched in dissolved inorganic carbon (DIC; 11.2 mM at 20 cm,
298 12.8 mM at 100 cm). Sulfate concentrations were lower than seawater, decreasing with
299 depth into the pool (Fig 4). Dissolved inorganic nitrogen was dominated by very high
300 NH_4^+ (1750 and 2195 μM , 20 and 100 cm, respectively), with NO_3^- disappearing sharply
301 in the top meter of the brine. The dissolved organic matter content was also high with a
302 low C:N of ~ 4.6 suggesting the importance of autochthonous production within the brine
303 waters.

304 4.2. *Water Column and Brine Pool CH_4 Oxidation and SO_4^{2-} Reduction Rates*

305 Aerobic methane oxidation rates measured in the water column above the brine
306 pool from depths of 300m to 2313m (or heights above the brine pool from 0 to 2013m)
307 ranged from 0.00 to $6.33 \pm 0.9 \text{ pmol L}^{-1} \text{ d}^{-1}$ (hereafter pM d^{-1}) (Fig 4). The aerobic
308 methane oxidation rate in the sample taken from directly above ($\sim 3\text{m}$) the brine pool
309 (2328m) was significantly higher, $129.6 \pm 18.2 \text{ pM d}^{-1}$, than rates at any other depth.
310 Using the methane concentrations determined via gas chromatography and the
311 empirically derived oxidation rates, we calculate an integrated methane oxidation rate in
312 the water column above the brine pool of $320 \mu\text{mol m}^{-2} \text{ yr}^{-1}$

313 Within the brine pool, two samples (20cm and 100cm) were retrieved and used
314 for sulfate reduction and anaerobic oxidation of methane (AOM) rate measurements
315 (Table 2). Methane concentrations measured in the bottles used for the rate
316 measurements were 454 and 1320 μM , respectively (Table 1), giving rates at these depths
317 of 78.8 ± 7.6 and $62.1 \pm 13.1 \text{ nmol L}^{-1} \text{ d}^{-1}$, respectively. Sulfate concentrations in the
318 brine were depleted relative to seawater with concentrations of 20 and 16mM at 20cm
319 and 100cm, respectively. Sulfate reduction rates were 107 and 50 nmoles per liter per
320 day (hereafter nM d^{-1}) at 20cm and 100cm, respectively, and were comparable to the rates
321 of AOM on a per mole basis. As mentioned above, these oxidation rates were measured
322 using water taken from CTD rosette bottles, which, when sampling gas-charged waters,
323 are subject to outgassing and gas phase exchange during recovery. Thus, these rates are
324 considered to be conservative estimates of anaerobic methane oxidation. *In situ* methane
325 oxidation rates are expected to be higher as methane concentrations increase (i.e., on a
326 first order basis up to k_{max}) and are calculated below.

327 **4.3. *Water Column and Brine Pool Methane Concentrations***

328 Methane concentrations measured in the water column (Fig 4) directly above the
329 brine pool ranged from 30nM to 70nM at depths between 2000 and 300m, representing
330 concentrations that were 15 to 32 times that of atmospheric equilibrium and underscoring
331 the transport of methane from below. At depths below 2000m, closer to the brine pool,
332 concentrations increased sharply and ranged from 111 nM to 24 μM . Methane
333 concentrations, as measured by the ISMS approximately 5 cm above the brine
334 fluid/seawater interface near the shore of the pool and 1 cm above the brine
335 fluid/seawater interface in the center of the brine pool were 180 and 590 μM ,

336 respectively. Approximately 1m above the brine surface, the ISMS-measured methane
337 concentration was approximately $\sim 35 \mu\text{M}$, which is in general agreement with the
338 methane concentrations measured in the CTD rosette at this depth ($24 \mu\text{M}$, well below
339 the saturation of methane at one atmosphere, so these particular Niskin measurements are
340 not compromised by off gassing, see Fig. 4). Concentrations at depths of 5, 20 and 80cm
341 into the brine fluid near the center of the pool were orders of magnitude higher (Fig 4)
342 with values of 14.3, 20.3 and 33.3 mM, respectively ($\pm 2\%$), and more than an order of
343 magnitude in excess of concentrations measured with Niskin sampling (see Table 1).

344 **5. Discussion**

345 **5.1. *In situ rates of brine pool anaerobic methane oxidation***

346 The *in situ* rates of AOM reported here exceed values in other anoxic waters by at
347 least one to two orders of magnitude. Our measured rates of AOM from two sampling
348 depths allowed calculation of first order rate constants of 0.063 and 0.017 yr^{-1} from
349 depths of 20cm and 80cm, respectively. Coupling the *in situ* methane concentration
350 measurements from the ISMS to the aforementioned rate constants yields estimates of the
351 actual rates of AOM of 1285 ± 125 and $572 \pm 121 \mu\text{M yr}^{-1}$ at the depths of 20 and 80cm
352 in the brine pool, respectively. To the best of our knowledge, these AOM rates are by far
353 the highest documented in an anoxic water body. Whereas there has been some evidence
354 that AOM is inhibited by high chloride concentrations (e.g.,(Joye *et al.*, 2009; Oren,
355 2002)), our data (Tables 1 and 2) suggest that moderately high salinities may in fact not
356 be inhibitory to AOM and that coupled sulfate reduction and AOM may yet play an
357 important role in many Gulf of Mexico hydrocarbon/brine environments.

358 Many studies have measured AOM in deep sea environments with high
359 concentrations of methane, and the highest rates are generally found within sediments,
360 particularly hydrate-influenced sediments (e.g., (Devol, 1983; Girguis *et al.*, 2003; Joye
361 *et al.*, 2004; Reeburgh, 1980)). Indeed, far fewer studies have measured AOM occurring
362 in anoxic water columns, and the rates reported in these studies are generally much lower
363 than sediment rates (as is the case with most biogeochemical processes primarily due to
364 microbial density being substantially higher in sediments). Rates of AOM measured in
365 the anoxic waters of the Cariaco Basin ($\sim 1.5 \mu\text{M yr}^{-1}$; Ward *et al.*, 1987), Saanich Inlet
366 ($7.3 \mu\text{M yr}^{-1}$; Ward *et al.*, 1989) and the Black Sea ($0.6 \mu\text{M yr}^{-1}$; Reeburgh *et al.*, 1991)
367 were all orders of magnitude lower than those observed in this study. Joye et al (1999)
368 measured rates as high as $17.5 \mu\text{M yr}^{-1}$ in the bottom waters of alkaline, saline Mono
369 Lake. Notably, these rates were measured during a period when the lake waters were
370 well mixed. More recent data collected during a period of extended meromixis in Mono
371 Lake exhibit substantially higher rates of AOM (up to $365 \mu\text{M yr}^{-1}$) (Joye *et al.*,
372 submitted).

373 The rates of AOM presented here are also consistent with the extremely high *in*
374 *situ* concentrations occurring at these depths. Turnover times of methane in the anaerobic
375 brines were on the order of 16 to 58 years, more than long enough to maintain supply of
376 SO_4^{-2} via diffusion. Such long turnover times also might suggest that supply to the
377 overlying water via diffusion is likely to be a substantial methane sink relative to removal
378 by AOM (or aerobic oxidation at the brine-seawater interface) in similar brine pool
379 environments. Indeed, given that the rate constants were lower than many other
380 comparable environments, the rates presented represent a conservative estimate and, as

381 previously mentioned, the rate constants would likely increase at the higher methane
382 concentrations found *in situ*.

383 **5.2. *Estimates of CH₄ flux from the brine pool***

384 Research on deep-sea fluxes and transformations of biological compounds is
385 constantly challenged by the need to sample at extreme temperatures, depths and
386 pressures. Measurement of these compounds *in situ* provides more rigorous constraints
387 on their fluxes and transformation rates. In the context of the current study, the *in situ*
388 mass spectrometer allowed direct measurement of methane concentrations in a gas-
389 charged brine pool. These concentration measurements were used to calculate a diffusive
390 flux of methane from the brine pool into the overlying water column of $1.1 \pm 0.2 \text{ mol m}^{-2}$
391 yr^{-1} , illustrating the magnitude of methane flux from benthic environs into the overlying
392 mixed layer (discussed in detail below).

393 Specifically, discrete *in situ* measurements taken over a depth profile across the
394 seawater-brine interface provide a context for calculating diffusive geochemical fluxes of
395 methane into the overlying water column. This approach has been used in numerous
396 studies for estimating the mass transfer (e.g., fluxes) of solutes from one region into
397 another. Brine pools such as AC601 are generally very quiescent in nature with fluid
398 advection playing a small role in controlling fluxes (Joye *et al.*, 2005; Joye *et al.*, 2009).
399 In these locations, where diffusion is the dominant mode of mass transfer, Fick's first law
400 is used to make first order estimates of the diffusive flux based on the measured
401 concentration gradients. The range of possible flux values is estimated based on error in
402 the spatial resolution of the gradient (i.e., since high precision positioning of the sampling
403 wand was not possible, we estimate the potential vertical position $\pm 10\text{cm}$). Moreover,

404 we adopted a value of $1.38 \times 10^{-5} \text{ cm}^2 \text{ s}^{-1}$ for the diffusion coefficient of methane in seawater
405 adjusted for the average viscosity of the brine using the Stokes-Einstein relationship
406 (Mao and Duan, 2008; Sahores and Witherspoon, 1970).

407 Even with our lower-bound estimate of diffusive methane flux ($1.1 \text{ mol m}^{-2} \text{ yr}^{-1}$),
408 water column integrated methane oxidation rates (Fig. 4) measured directly above the
409 brine pool ($\sim 320 \text{ } \mu\text{mol m}^{-2} \text{ yr}^{-1}$) indicate that only a very small fraction of methane
410 escaping the brine pool is biologically consumed in the overlying water column (0.02 to
411 0.03%). While it is likely that lateral advection plays a large role in the disconnect
412 between brine pool flux and water column methane oxidation rates above the brine, upper
413 water column concentrations are nonetheless >10 times that of methane in equilibrium
414 with the atmosphere, confirming the transport of methane from depths $>2000\text{m}$ into the
415 mixed layer which easily escapes, un-oxidized, into the atmosphere.

416 Our estimates of diffusive methane flux should be taken as a lower bound on flux
417 from environments such as Gulf of Mexico brine pools. They also underscore the value
418 of *in situ* measurement for constraining methane fluxes. For example, other studies
419 (Lapham *et al.*, 2008; Schmidt *et al.*, 2003) have modeled diffusive and/or advective
420 fluxes from brine seep environments by comparing profiles of a non-conservative solute
421 (e.g., methane) to conservative solutes (e.g., chloride, temperature). Using a 1-D reaction
422 transport model together with chloride and methane profiles, advective methane fluxes of
423 up to $2 \text{ mol m}^{-2} \text{ yr}^{-1}$ were estimated from brine-influenced sediments characterized by a
424 strong advective flow (Lapham *et al.*, 2008). However, these calculated fluxes were
425 based on methane concentrations from sediment cores that had degassed upon collection,
426 and were thus substantially lower than methane concentrations *in situ*. In another study,

427 Solomon et al. (2008) employed osmotic samplers demonstrating that net seafloor
428 methane fluxes range from $0.89 \text{ mol m}^{-2} \text{ yr}^{-1}$ from a mussel bed environment up to 29
429 $\text{mol m}^{-2} \text{ yr}^{-1}$ from a bacterial mat environment. Hereto, because methane concentrations
430 could not be reliably determined at *in situ* pressure and temperature, fluxes were
431 calculated assuming that porewater was in equilibrium with methane hydrate under *in situ*
432 conditions. However, others have shown that methane in sediments around hydrate can
433 be far from saturated (Lapham *et al.*, pers. comm.), which would result in much lower
434 flux estimates. Future studies should aim to couple *in situ* methane measurements with
435 direct fluid flow measurements to better constrain the contribution of advective flux to
436 water column methane flux.

437 **6. Summary and Conclusions**

438 Our calculated *in situ* AOM rates, using empirically derived rate constants, are
439 higher than those previously published by one to two orders of magnitude. The diffusive
440 flux was estimated to range as high $1.8 \text{ mol m}^{-2} \text{ yr}^{-1}$ from the brine pool, while integrated
441 oxidation rates in the overlying 2000m water column could only account for $0.32 \text{ } \mu\text{mol}$
442 $\text{m}^{-2} \text{ yr}^{-1}$. These data suggest that a very large component of the diffusive brine pool
443 methane flux escapes both aerobic and anaerobic oxidation in the water column above the
444 brine pool and may be released into the atmosphere (or at least subject to dispersion via
445 lateral advection). These first *in situ* measurements of methane concentration from a
446 brine pool using the ISMS enabled robust quantification of methane concentrations at *in*
447 *situ* conditions in these gas-charged brines and reflect the strong influence of the
448 surrounding hydrocarbon seep environments. Such integrated approaches – wherein
449 geochemical determinations are coupled with microbiological activity measurements –

450 are the best means of providing a rigorous constraint on methane diffusive fluxes and
451 transformation rates. This will improve our understanding of the role that hydrocarbon
452 seeps may play in the delivery of methane into the ocean and ultimately the atmosphere.
453

453 **Acknowledgements**

454 We are especially grateful to Stephane Hourdez for his immense help with this
455 instrument during this expedition. We are also grateful to Dr. Charles Fisher for his
456 support, as well as the captains and crew of the *RV Ronald H. Brown* and *RV Atlantis*.
457 Special thanks to the pilots and support staff of the *DSV ALVIN* and *DSV JASON II* from
458 Woods Hole Oceanographic Institution for help collecting and processing samples. An
459 extra thanks goes especially to Matthew Heinz and Tito Collasius for their assistance in
460 the machine shop. This research was supported by the U.S. Department of the Interior
461 Minerals Management Service, the National Oceanic and Atmospheric Administration,
462 the David and Lucile Packard Foundation, Harvard University, and the National Science
463 Foundation (MCB-0702504).

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465

465 **Figure captions**

466 **Figure 1:** Schematic of the in situ mass spectrometer. a) membrane inlet housing, b)
467 front end plate of titanium pressure housing, c) high pressure solenoid for isolation of
468 vacuum chamber, d) Alcatel ATH-31+ Turbo Pump, e) vacuum flight tube housing the
469 SRS Quadrupole RGA-200 including ion source, quadrupoles and detectors f) electronics
470 head for controlling and reading spectrometer signals g) KNF Neuberger roughing pump
471 model ANDC-84.3. Sample gas is continuously extracted across the membrane located
472 in (a) into the high vacuum system (d, g), ionized in (e) and analyzed by the detector and
473 electronics control unit housed in (f). The instrument is approximately 1m in length.

474

475 **Figure 2:** a) Normalized response at m/z 15 over a range of hydrostatic pressure for
476 three example fluid temperatures and concentrations, 10°C $1160\mu\text{M}$ CH_4 (grey squares),
477 2°C $800\mu\text{M}$ CH_4 (black triangles) and 14°C $180\mu\text{M}$ CH_4 (grey triangles). Responses to
478 pressure were experimentally fit under a wide range of temperatures and concentrations
479 (as in Bell et al 2007) with values of b' ranging between 0.02 to 0.24 and values of k
480 ranging between 0.84 to 0.94. b) The response of m/z 15 (corrected for pressure effects –
481 see Fig 2a) was linearly proportional to methane concentrations as measured
482 independently by gas chromatography (grey triangles) and as calculated after Duan et al
483 2006 during high pressure calibration measurements (black circles).

484

485 **Figure 3:** Photo from the Pilot Cam of ROV Jason showing the starboard manipulator
486 reaching through the seawater/brine pool interface and sampling at a depth of
487 approximately 80cm.

488

489 **Figure 4:** a) Depth profile of methane concentration and methane oxidation rates in the
490 water column above brine pool AC601. Note log scale. Open circles are concentration
491 measurements made from Niskin bottle samples, while black circles are those made *in*
492 *situ* using the ISMS. b) Close-up of seawater/brine pool interface and profile into the
493 brine fluid. Note the linear scale in contrast to panel a. Measured rates of anaerobic
494 methane oxidation (AOM) at two depths within the brine pool are shown. Note that
495 these, when corrected for in situ CH₄ concentrations, these rates are 30-45 times higher.
496 Sulfate concentrations are depleted in the brine, consistent with its role in AOM.

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Table 1: Major chemical components of brine pool AC601

Depth cm	pH	salinity	oxygen	Concentration mM						Concentration mM															
				DIC	H ₂ S	SO ₄ ⁻²	Cl ⁻	CH ₄ ^a	CH ₄ ^b	NH ₄ ⁺	NO _x	DIN	TDN	DON	DOC	DOC:DON									
5											14.35														
20	6.29	82	<2 μM	11.2	0.00	20	1366	0.454	20.29	1750	3.4	1753.4	1843.5	90.1	423.5									4.7	
80											33.29														
100	6.25	92	<2 μM	12.8	0.25	16	1533	1.320	38.40*	2195	0.3	2195.3	2280.5	85.2	380.0									4.5	

^aConcentrations measured via gas chromatography on samples retrieved with a CTD rosette and Niskin bottles

^bConcentrations measured via *in situ* mass spectrometer

* estimated by regression of the three ISMS data points collected above

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Table 2: Rates of sulfate reduction and anaerobic methane oxidation in Gulf of Mexico brine pool AC601

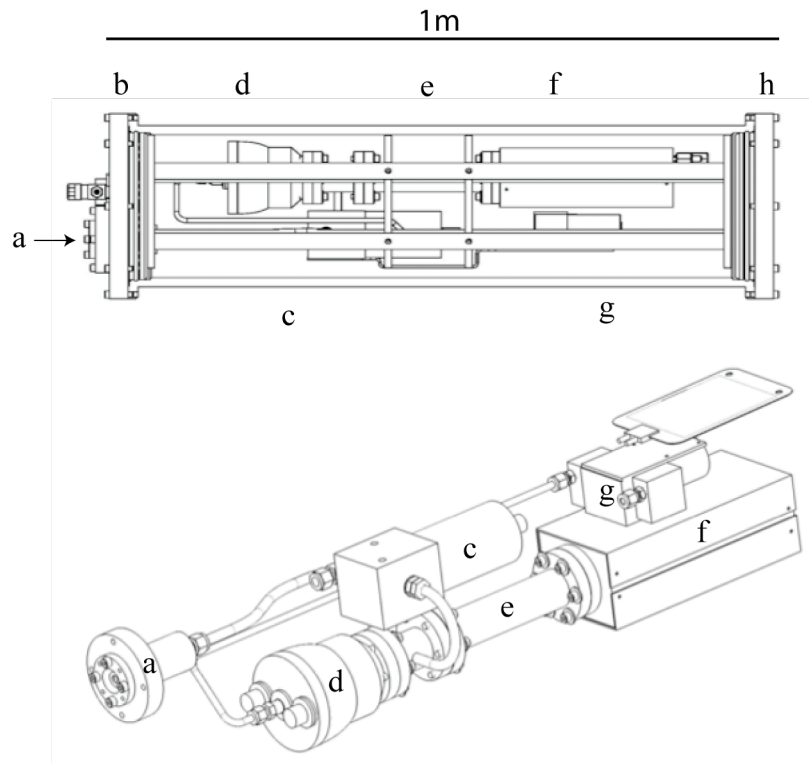
Depth cm	Rate nM/d		
	Sulfate Reduction	Anaerobic Methane Oxidation*	Anaerobic Methane Oxidation [^]
20	107.1 ± 14.6	78.8 ± 7.6	3502.0 ± 340
100	49.8 ± 8.4	62.1 ± 13.1	1807.4 ± 330

* measured using water samples collected via CTD rosette and Niskin bottles

[^] estimated rates corrected for measured concentrations *in situ* and using rate constants measured from shipboard incubations of brine pool water collected via Niskin bottles

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Wankel Figure 1

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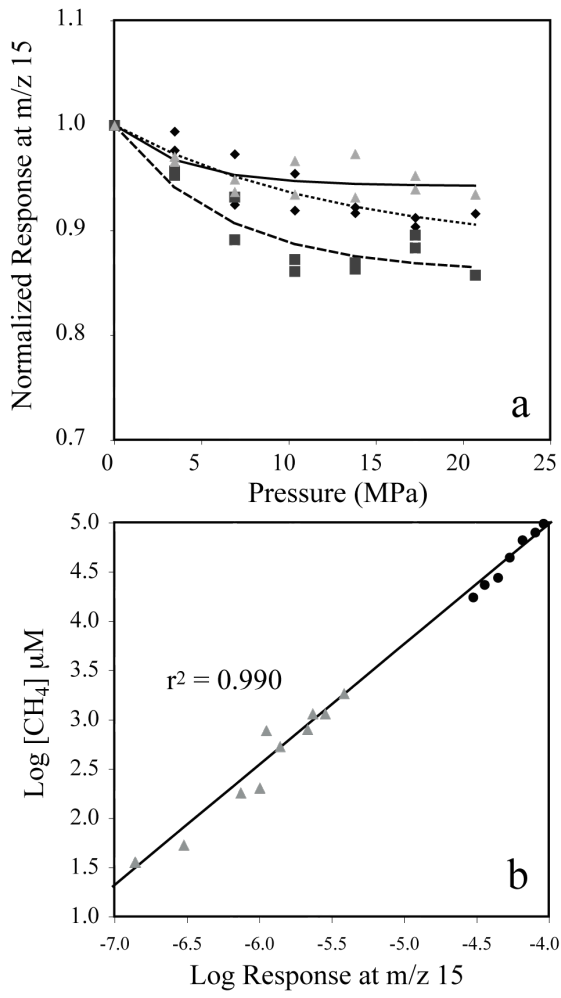
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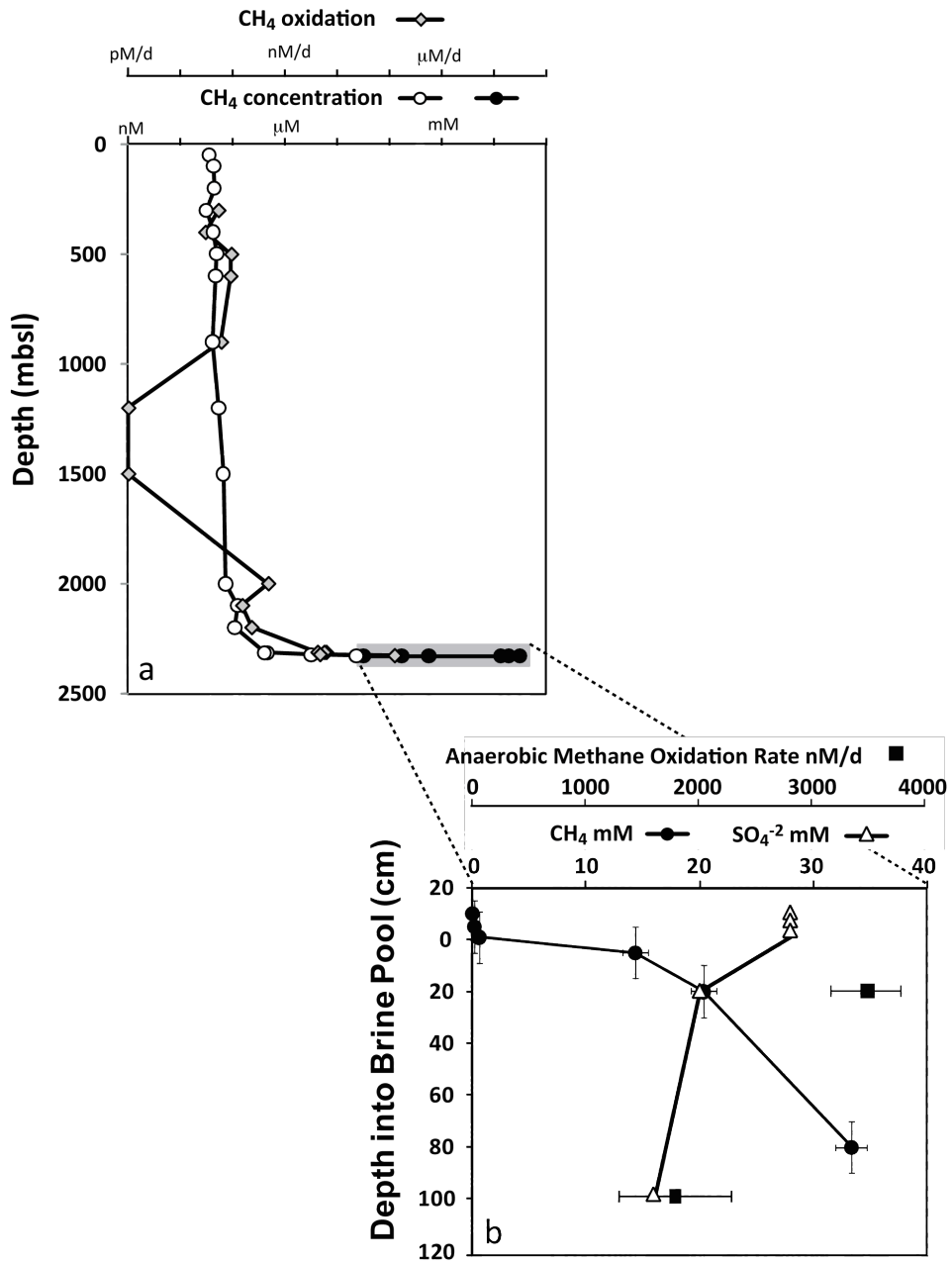
Wankel Figure 2

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Wankel Figure 3



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Wankel Figure 4