

Redox controls on methane formation, migration and fate in shallow aquifers” by Humez et al.

Point-by-point reply to the comments

Dear Editor,

Thank you for having accepted our manuscript submitted to HESS with minor revisions. Please find below our point-by-point reply to the comments by the reviewers with the location of modifications (line and page) as they appear in the revised manuscript.

We hope that these changes address the concerns of the editor and the reviewers in a satisfactory manner.

With best regards on behalf of all co-authors,

Dr. Pauline Humez

From reviewer #1 (Dr. McIntosh)

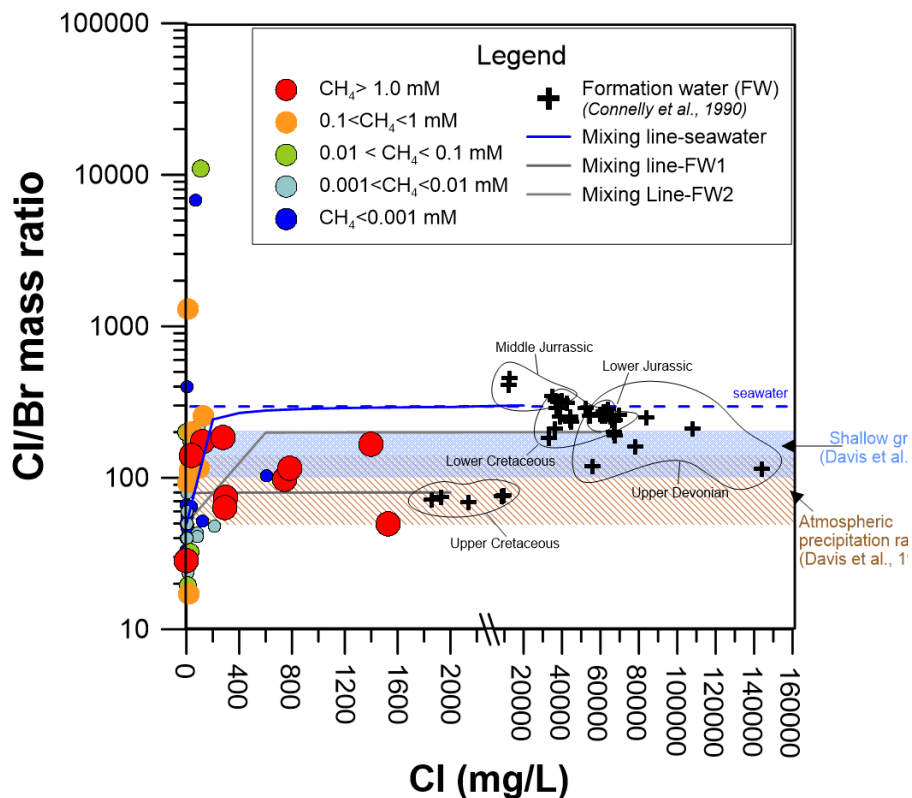
#	Comments/Authors Comments (AC)	Revised manuscript location
1	Pg 2, Line 21, my last name is misspelled: “McInstosh et al., 2014” should be “McIntosh”.	
	AC: This correction has been made in the revised version of our manuscript.	Line 21 p 2
2	Pg 2, Line 28, our companion paper to McIntosh et al. (2014), “Hamilton et al. (2015) Hydrogeology Journal” compared water chemistry to methane concentration in groundwater in Ontario, similar to your study. Please include this reference.	
	AC: This reference has been added to the revised version of our manuscript.	Line 22 p 2
3	Pg 3, Lines 20-23: Methanogenic systems can also have high d13C-CH4 values from closed system CO2-reduction, where most of the CO2 pool has been depleted, and d13C values of CH4 and CO2 become increasingly more positive (up to -50+ per mil in some cases). This is another way that d13C-CH4 values can seem “artificially” high, but still be methanogenic. See Bates et al. (2011) Chemical Geology. I would add a sentence on this here and point out that d13C-CO2 and/or d13C-DIC values can help distinguish these relatively positive d13C-CH4 values from methanogenic vs. thermogenic gas sources.	
	AC: This is a very good point. To address this we have added the following statements to the revised version of our manuscript: “Methanogenic systems can also be characterized by high $\delta^{13}\text{C}_{\text{CH}_4}$ values and thus by a pseudo-thermogenic methane isotope signature as a results of CO₂ reduction in a closed system. Under such circumstances, $\delta^{13}\text{C}$ values of CH₄ and CO₂ both increase as the CO₂ pool becomes progressively depleted (Whiticar et al., 1986; Whiticar, 1999; Bates et al., 2011). Hence, $\delta^{13}\text{C}_{\text{CO}_2}$ and $\delta^{13}\text{C}_{\text{DIC}}$ values constitute an additional parameter that can help to distinguish whether elevated $\delta^{13}\text{C}$ values of methane are associated	Line 21-29 p 3

	with biogenic methane formation or thermogenic gas sources (Whiticar et al., 1986; Whiticar, 1999)".	
4	Pg 7, Line 28, change "adding" to "summing". AC: This correction has been made in the revised version of our manuscript.	Line 2 p 8
5	Pg 9, Line 17, "closed" should be "close" to the LMWL. AC: This correction has been made in the revised version of our manuscript.	Line 23 p 9
6	Pg 10, section 3.6. Did you measure d13C-C2 values? If so, include. This could help identify microbial oxidation of higher chain hydrocarbons, microbial production of ethane, or mixing with thermogenic gas. AC: This is a very good point. The C isotope ratios of ethane have been measured on 19 samples. The plot of concentrations versus $\delta^{13}\text{C}$ values of ethane is displayed below. For a sub-set of samples there is a trend of increasing $\delta^{13}\text{C}$ values with decreasing ethane concentration, consistent with ethane oxidation. All these samples are from the "mixed category", and we feel that this observation does not add much new insight into the existing discussion. Hence, we prefer not to add this information unless the editor and/or reviewer strongly suggest otherwise.	No changes have been made in the revised manuscript
7	Pg 11, Line 6+, we also found the highest methane values in groundwater in Ontario associated with Na-Cl and Na-HCO3 type waters (see McIntosh et al., 2014; Hamilton et al., 2015). AC: Thank you for pointing this out. We have added the above references to this statement.	Line 15 p 11
8	Figure 5: Plot end-members on the other plots in Figure 5. AC: The end-members have been integrated in the new version of Figure 5.	See revised Fig. 5
9	Pg 11, Line 14, Needs further explanation. Does your data (e.g. lack of Br??) differentiate between these two sources of Na-rich waters: brines versus cation exchange? Reader is left wondering which of these processes is important here,	

which can have implications for fluid migration vs. in-situ water-rock reactions.

AC: The Br concentrations have been analyzed and we have created a plot of Cl/Br ratios vs Cl concentrations (see plot below) with current seawater shown as a dashed blue line. The Cl/Br ratios of our samples are in the range of ratios previously reported for other groundwater samples from Canada and United States (blue area in Figure below according to Gue et al. 2015; Davis et al., 2004). We stated in the original manuscript that Na-rich waters are due to either mixing with deep saline water and/or cation exchange. The plot below provides further initial insights by suggesting that samples with elevated Na and Cl concentrations appear to be mixtures between low TDS groundwater and high TDS formation waters (see crosses in Figure below, Connelly et al. 1990). We also observed that 7 of 8 samples with Cl concentration > 200 mg/L fall in the highest methane concentration category of > 1.0 mM and belong to category #4. This provides an initial indication that in select cases elevated methane in groundwater may be associated with admixture of deeper saline water rather than associated with Na-bicarbonate water type due to cation exchange processes. However, this hypothesis about methane migration pathways requires significant further testing based on regional hydrodynamic and geochemical data, which we intend to conduct in 2016. At this point, we feel that we have insufficient clear evidence to further discuss these options, and therefore have delete the sentence “Typically, Na-dominated groundwaters are linked to two principal processes: (1) mixing with deep saline fluids (Na-Cl types) and (2) cation exchange (Na-HCO₃ type).”.

Some text was deleted to simply this discussion (line 21, p 11)



10	Pg 12, Line 9, your results of finding high CH ₄ only when [SO ₄] ^{<} 1 mM is consistent with what has been observed in deeper coalbed methane and organic-rich shale microbial gas systems. I would make this link by adding a sentence and reference to that literature. See Schlegel et al. (2011) or other reference.	
	AC: Thank you for this valuable suggestion. Following your suggestion we have added the following statement and reference to the revised version of our manuscript: “This result is consistent with what has been observed in deeper coalbed methane and organic-rich shale microbial gas systems (Schlegel et al., 2011).”	Line 16-17 p 12
11	Pg 13, Line 2, add “CH ₄ ” to your list of isotopic compositions investigated.	
	AC: This has been changed in the revised version of our manuscript.	Line 9 p 13
12	Pg 13, Section 4.3.1. The question mark for group 3 in Figure 8 needs an explanation– i.e. what does the “question mark” represent?	
	AC: This question mark has been removed from Fig. 8 in the revised version of our manuscript to avoid confusion.	See revised Fig. 8
13	Pg 14, Line 20+, This part needs clarification because you go on to say, and show in Figure 11, that there are some samples with evidence of methane oxidation, whereas you say here that there is no evidence of oxidation. Be more specific here, or simply remove statement and save for later when you discuss the higher d ¹³ C-CH ₄ values.	
	AC: This statement has been deleted in the revised version of our manuscript.	Line 27 p 14
14	Pg 15, Line 27, “has not yet been oxidized” – be more specific: e.g. there is no evidence of methane oxidation because the elevated d ¹³ C-CH ₄ values are not associated with low d ¹³ C-DIC values, as expected for methane oxidation.	
	AC: As suggested, this sentence has been added in the revised version of our manuscript as “since there is no evidence of methane oxidation such as low δ¹³C_{DIC} values as expected for methane oxidation”.	Line 4-5 p 16
15	Pg 16, Line 8, change “imparting a d ¹³ C value” to “imparting a relatively high d ¹³ C value”.	
	AC: This modification has been integrated in the revised version of our manuscript.	Line 15 p 16
16	Pg 16, Line 10, “It is also possible that post-sampling degradation of low-methane samples occurred.” Be more specific. What could have happened (physically) and how might that have changed the isotopic values?	
	AC: The statement “post-sampling degradation of low-methane samples occurred” refers to potential slow diffusive gas loss from sampling containers resulting in ¹³C enrichment in the residual methane. This has been clarified in the revised version of the manuscript.	Line 18-19 p 16
17	Pg 17, Line 23, change “interpretation of gas composition” to “interpretation of natural gas composition.”	
	AC: This correction has been added to the revised version of our manuscript.	Line 1 p 18
18	Pg 18, Line 3: add an “and” between “d ¹³ C, dD”.	
	AC: This has been corrected in the revised version of our manuscript.	δD removed from the discussion

		in agreement with the comment #21
19	Pg 18, Line 7, be more specific about this statement. For example, could add text at the end of the sentence: “such as the introduction of deeply-sourced thermogenic gases into shallow aquifers.” AC: We have followed the advice of the reviewer and have added this statement to the revised version of our manuscript.	Line 14-15 p 18
20	Figure 3, part b, It’s not clear what the “(C(-4)) becomes stable” label represents. Need to explain in the figure caption and/or text. AC: This note has been removed from Fig. 3 in the revised version of our manuscript to avoid confusion.	See revised Fig. 3
21	Figure 11, part b, there wasn’t much (if any?) discussion of the d2H-CH4 values in the text - add. BTW - I’m no longer using this plot in my own research because I’ve found that it is misleading; the dD-CH4 values are low in these western, higher latitude regions not because of a shift in metabolic pathway to “methyl type fermentation”, but rather because of isotopic exchange between the 2H in the CH4 and H2O (shown in several studies now). Recent microbial studies from several coalbed methane and black shale systems show that both CO2 reduction and acetoclastic methanogenesis are typically important/present. I have a paper in review on this topic that will hopefully be published soon. In the meantime, see Bates et al. (2011) Chemical Geology for a reference. AC: We fully agree with the reviewer’s reasoning especially since the $\delta^2\text{H}$ values of groundwater in Alberta are very negative. We therefore have removed Figure 11b and the associated discussion in the text.	Fig. 11b and associated discussion removed from the text
22	Figure 12, part b, need to subscript “CH4” in the y-axis label. AC: This subscript has been added in the y-axis label of Fig. 12 in the revised version of our manuscript.	See revised Fig. 12

From reviewer #2

#	Authors Comments (AC)	
1	We have changed the units in Figures 6, 7 and 9 to mmol/L as suggested;	See Revised Figs. 6, 7 and 9
2	The ordinate label for Figure 12 has been refined by using mmol/L as a unit for sulfate concentration and by changing the number increments on the x-axis of both graphs ($\Delta x = 1$ mmol/L for the sulfate concentration axis and $\Delta x = 2$ ‰ for $\delta^{13}\text{C}_{\text{DIC}}$ axis).	See revised Fig. 12
3	Table 2 has been changed to show significant digits consistent with the stated measurement uncertainty of the respective parameters;	See Table 2

Additional

Authors Comments (AC)	
We have also made additional minor changes in other figures to ensure consistency with respect to units as used throughout the text.	See revised Fig. 3a; Fig. 4c; Fig. 10a

Redox controls on methane formation, migration and fate in shallow aquifers

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Abstract. Development of unconventional energy resources such as shale gas and coalbed methane has generated some public concern with regard to the protection of groundwater and surface water resources from leakage of stray gas from the deep subsurface. In terms of environmental impact to and risk assessment of shallow groundwater resources, the ultimate
15 challenge is to distinguish: (a) natural in-situ production of biogenic methane, (b) biogenic or thermogenic methane migration into shallow aquifers due to natural causes, and (c) thermogenic methane migration from deep sources due to human activities associated with the exploitation of conventional or unconventional oil and gas resources. This study combines aqueous and gas (dissolved and free) geochemical and isotope data from 372 groundwater samples obtained from 186 monitoring wells of the provincial Groundwater Observation Well Network (GOWN) in Alberta (Canada), a province with a long record of
20 conventional and unconventional hydrocarbon exploration. We investigated whether methane occurring in shallow groundwater formed in-situ, or whether it migrated into the shallow aquifers from elsewhere in the stratigraphic column. It was found that methane is ubiquitous in groundwater in Alberta and is predominantly of biogenic origin. The highest concentrations of biogenic methane (> 0.01 mM or > 0.2 mg/L), characterized by $\delta^{13}\text{C}_{\text{CH}_4}$ values < -55 ‰, occurred in anoxic Na-Cl, Na-HCO₃ and Na-HCO₃-Cl type groundwaters with negligible concentrations of nitrate and sulfate suggesting that
25 methane was formed in-situ under methanogenic conditions for 39.1% of the samples. In only a few cases (3.7 %) was methane of biogenic origin found in more oxidizing shallow aquifer portions suggesting limited upward migration from deeper methanogenic aquifers. 14.1% of the samples contained methane with $\delta^{13}\text{C}_{\text{CH}_4}$ values > -54 ‰, potentially suggesting a thermogenic origin, but aqueous and isotope geochemistry data revealed that the elevated $\delta^{13}\text{C}_{\text{CH}_4}$ values were caused by microbial oxidation of biogenic methane or post-sampling degradation of low CH₄ content samples rather than migration of
30 deep thermogenic gas. A significant number of samples (39.2%) contained methane with predominantly biogenic C isotope ratios ($\delta^{13}\text{C}_{\text{CH}_4} < -55$ ‰) accompanied by elevated concentrations of ethane and sometimes trace concentrations of propane. These gases observed in 28.1% of the samples, bearing both biogenic ($\delta^{13}\text{C}$) and thermogenic (presence of C₃)

characteristics, are most likely derived from shallow coal seams that are prevalent in the Cretaceous Horseshoe Canyon and neighboring formations in which some of the groundwater wells are completed. The remaining 3.7% of samples were not assigned because of conflicting parameters in the datasets or between replicates samples. Hence, despite quite variable gas concentrations and a wide range of $\delta^{13}\text{C}_{\text{CH}_4}$ values in baseline groundwater samples, we found no conclusive evidence for deep thermogenic gas migration into shallow aquifers either naturally or via anthropogenically-induced pathways in this baseline groundwater survey. This study shows that the combined interpretation of aqueous geochemistry data in concert with chemical and isotopic compositions of dissolved and/or free gas can yield unprecedented insights into formation and potential migration of methane in shallow groundwater. This enables the assessment of cross-formational methane migration and provides an understanding of alkane gas sources and pathways necessary for a stringent baseline definition in the context of current and future unconventional hydrocarbon exploration and exploitation.

Keywords. Methane • Alberta • groundwater • stable isotopes • geochemistry • redox processes

1 Introduction

Development of unconventional energy resources such as shale gas and coalbed methane is often accompanied by concerns of some landowners and parts of the public that shallow groundwater could be affected by leakage of stray gas from the deep subsurface. To address unambiguously such concerns, it is essential to assess the natural occurrence of methane and its spatial distribution, the variability of methane concentrations and the sources of methane in shallow groundwater prior to unconventional energy development to establish a baseline. In the last 5 years, an increasing number of publications have addressed the questions of occurrence and sources of methane in shallow groundwater in natural gas producing regions (Osborn et al., 2011a and b; Warner et al., 2013; Darrah et al., 2012, 2014; Jackson et al., 2013; Molofsky et al., 2013; Siegel et al., 2015; Vengosh et al., 2013; Brantley et al., 2014; Baldassare et al., 2014; McPhilips et al., 2014; McIntosh et al., 2014; Vidic et al., 2013; [Hamilton et al., 2015](#)). These studies have contributed a wealth of baseline data for gas occurrences in shallow groundwater in many regions of North America. Some of these studies reported that elevated methane concentrations in shallow aquifers were correlated with geology, especially the occurrence of low-sulfur coal deposits, and topography, since groundwater from wells in valleys tended to have higher methane concentrations (Mathes and White, 2006; Molofsky et al., 2013, Etiope et al., 2013). In other cases, methane concentrations were correlated with groundwater types with elevated methane concentrations predominantly reported in sodium chloride or sodium bicarbonate groundwater types (Molosky et al., 2013; McPhilips et al., 2014). Where thermogenic gas was found in shallow groundwater, it is however not always clear to what extent this occurred inadvertently as a result of human activities or due to natural flowpaths.

Assessment of the aqueous geochemistry and the redox conditions in the aquifers affected by elevated methane concentrations can reveal whether methane formed in-situ, or whether it formed elsewhere in the stratigraphic column and

migrated into the shallow aquifer. In terms of potential environmental impact and risk assessment focusing on shallow groundwater resources, the ultimate challenge is to distinguish: (a) natural in-situ production of biogenic methane in methanogenic aquifers, (b) biogenic or thermogenic methane naturally migrating into shallow aquifers, and (c) predominantly thermogenic methane from deep sources migrating due to human activities associated with exploitation of conventional or unconventional oil and gas resources.

In-situ formation of methane in shallow aquifers requires highly reducing conditions. According to the 'redox ladder' concept, microbial formation of CH₄ can only occur after dissolved oxygen is consumed, denitrification has removed nitrate, and bacterial sulfate reduction has progressed towards completion (Appelo and Postma, 2005; Barker and Fritz, 1981; Darling and Goody, 2006; Whiticar, 1986). Hence, analysing a variety of water chemistry parameters (e.g. the redox couples Fe³⁺/Fe²⁺, NO₃⁻/NO₂⁻, CO₂/CH₄, SO₄²⁻/H₂S etc.) can provide important clues to whether in-situ formation of methane within a shallow aquifer is possible. In addition, biogenic methane formed in shallow aquifers is characterized by very negative δ¹³C and δ²H values (e.g. Whiticar, 1999). Therefore, isotope analyses on methane, higher alkanes (where present) and other dissolved groundwater constituents such as dissolved inorganic carbon (δ¹³C_{DIC}), nitrate (δ¹⁵N_{NO3}, δ¹⁸O_{NO3}) and sulfate (δ³⁴S_{SO4}, δ¹⁸O_{SO4}) can provide important additional insights about redox conditions and methane formation pathways in shallow aquifers.

Combined geochemical and isotopic analyses on groundwater and its dissolved or free gases phases thus have the potential to determine whether in-situ methane formation is possible or if gas migration must have occurred. The latter would be for instance the case if biogenic methane is found in aerobic or non-methanogenic aquifer sections. Alternatively, if thermogenic methane with elevated δ¹³C values accompanied by ethane and propane (e.g. Whiticar, 1999) is detected in shallow aquifers, gas migration from deeper geological formations into shallow aquifers must be postulated. In these cases, it is desirable to determine the depth of the gas source and its natural or anthropogenic migration pathways. It is equally important to identify apparent (or pseudo-) thermogenic methane characterized by elevated δ¹³C values that are in reality caused for instance by microbial oxidation of biogenic methane enriching the remaining methane in ¹³C (Barker and Fritz, 1981), rather than by migration of thermogenic gas from deep geological sources. Methanogenic systems can also be characterized by high δ¹³C_{CH4} values and thus by a pseudo-thermogenic methane isotope signature as a result of CO₂-reduction in a closed system. Under such circumstances, δ¹³C values of CH₄ and CO₂ become both increasingly more positive as the CO₂ pool becomes progressively depleted (Whiticar et al., 1986; Whiticar, 1999; Bates et al., 2011). Hence, δ¹³C_{CO2} and/or δ¹³C_{DIC} values constitute an additional parameter to distinguish elevated δ¹³C_{CH4} values of methane from biogenic versus thermogenic gas sources (Whiticar et al., 1986; Whiticar, 1999).

Alberta is a province in Western Canada with a long history of conventional and unconventional energy exploitation. Conventional oil and natural gas have been produced from numerous reservoirs in the province since the first natural gas find in 1883 and more than 400,000 oil and natural gas wells have been drilled (e.g. Breen, 1993). More recently,

unconventional natural gas has been exploited from rather shallow coalbed deposits (200-800 m below ground), predominantly in the Horseshoe Canyon Formation in and east of the Edmonton to Calgary corridor (ECC) in the southeastern part of the province with a peak activity occurring between 2006 and 2010. In the last decade, shale gas exploration and exploitation have also commenced in the Triassic Montney and the Devonian Duvernay Formations in the northwestern part of the province typically at depths exceeding 1.5 km. Therefore, assessment of the occurrence and the sources of methane in shallow groundwater is of key importance for two reasons: 1) to assess whether previous oil and gas exploitation has caused negative impacts on shallow groundwater due to stray gas contamination; and 2) to establish a baseline against which potential future impacts of stray gas migration on shallow aquifers, or the lack thereof, can be determined.

A baseline study was conducted between 2006 and 2014 investigating the occurrence of methane in shallow groundwater of Alberta (Canada) obtained from provincial monitoring wells. The objective was to determine, based on comprehensive aqueous geochemical and isotopic data evaluations of groundwater samples and their dissolved and free gases, the distribution and sources of methane in shallow groundwater. An additional goal was to characterize the hydrochemical environment in which methane was formed or transformed through redox processes to evaluate whether methane occurring in shallow groundwater in Alberta at baseline conditions has formed in-situ and under which geochemical conditions, or whether methane had formed elsewhere and has migrated into the shallow aquifers.

2 Background: Materials and Methods

2.1 Study site GOWN network

The Groundwater Observation Well Network (GOWN) of the Alberta Government initiated in 1955 (Alberta Research Council, 1956) and taken over by Alberta Environment (AENV) in 1982 is comprised of groundwater monitoring wells completed in various shallow aquifers throughout the province (Fig. 1). A recent comprehensive monitoring program collects water level information together with geochemical and isotopic data (H, O, C, S) since 2006 in order to record potential impacts on quantity and quality of groundwater in Alberta. The GOWN consists currently of over 250 active observation wells with many wells located in coalbed methane (CBM) production areas in the southeastern part of the province, while few wells exist in the shale gas development regions in the northwest as shown by the unequal spatial distribution of the well locations in Fig. 1. Since 2006, groundwater samples and dissolved gas and free gas samples from GOWN wells have been routinely obtained where possible for chemical and isotopic analyses. A first assessment of the gas geochemical dataset has been reported by Humez et al. (2015, 2016). This study evaluates both aqueous and gas geochemical and isotopic data of the GOWN monitoring program of shallow groundwater samples. GOWN wells are drilled into aquifers either within surficial deposits from the last major glaciation or reach sedimentary bedrock of usually Paleogene or Cretaceous age. The depth of the GOWN wells varies from 26 m to 250 m with an average of 60 m below ground surface (bgs) accessing groundwaters from different shallow aquifers (Fig. 1). The wells have typically stainless steel

casing with diameters varying from 32 mm to 254 mm with an average of 109 mm and they typically have short stainless steel or PVC screens only in the target aquifer formation.

The aquifer lithologies vary considerably comprising mostly fractured mudstone, sandstone and siltstone beds or lenses, pre-glacial sand, or surficial sandy and gravelly lacustrine or moraine deposits (Dawson et al., 1994, Fig. 1). The regional Upper Cretaceous-Paleogene stratigraphy differentiates many sandy clastic depositions including the (i) Lower Campanian Milk River Formation (and equivalents), (ii) Middle to Upper Campanian Belly River (Judith River) Group (and equivalents), (iii) Upper Campanian to Lower Maastrichtian Horseshoe Canyon Formation (and equivalents), (iv) Upper Maastrichtian to Lower Paleocene Scollard Formation (and equivalents), and (v) the Middle to Upper Paleocene Paskapoo Formation (and equivalents), which comprise the major shallow aquifers in the study area. Four sedimentary units with more fine-grained materials comprise the (i) the Lower Campanian Pakowki Formation (upper Lea Park Formation), (ii) the Middle Campanian Bearpaw shales, (iii) the Maastrichtian Battle Shales Formation, and (iv) the upper part of the Scollard Formation (Dawson et al., 1994), which are typically classified as aquitards (Fig. 1).

The Quaternary deposits include the Muriel Lake Formation that is composed of silt, sand and gravel of glacio-fluvial origin. Among the stratigraphic intervals containing coal zones with CBM potential are the Lower Cretaceous Mannville Group (e.g. Mannville Group coals), and the Belly River (e.g. McKay Coal, Taber Coal, Lethbridge Coal zones), Horseshoe Canyon (e.g. Drumheller Coal zone, a primary CBM target), and Scollard Formations (e.g. Ardley Coal zone). Thin coal seams occur also throughout the Paskapoo Formation. More information about these geological formations can be found in Meyboom (1960), Rosenthal et al. (1984), Hamblin (1998), Hamblin (2004), Dawson et al. (1994), Lyster and Andriashek (2012), Grasby et al. (2008), and Prior et al. (2013).

2.2 Samples and laboratory techniques

Between 2006 and 2014, a total of 372 groundwater samples were obtained from 186 GOWN wells accessing various shallow aquifers throughout Alberta. Many wells were sampled repeatedly, either on the same day as replicates for sampling and analytical quality control, or at greater time intervals to assess temporal water quality variations. All samples, including the replicates, are considered as individual samples in this paper. The database contains both aqueous and gaseous geochemical data for 372 samples. An electrical charge balance criteria for cations and anions of $\pm 10\%$ was applied for the aqueous geochemical data. Since calcium concentrations were not reported for a number of samples 35% of the groundwater samples were excluded leaving 242 samples (criteria #1, Table 1). Eight additional samples were discarded because no gas composition data were reported (criteria #2, Table 1). Among the 234 remaining samples, 150 samples had dissolved and free gas information, and 80 samples had only dissolved gas analyses reported. Four samples had only information on free gas without dissolved gas concentrations being reported. Among these 234 samples, 9 samples had no chemical data reported. Hence, a total of 225 samples have been evaluated as they have information on gas composition associated with balanced major ion chemistry i.e. Ca, Mg, Na, K, SO₄, NO₃, Cl, DIC (dissolved inorganic carbon) required to evaluate the

water type of the investigated samples (criteria #3, Table 1). 135 of 225 samples (60%) contained ion chemistry, gas analyses and carbon isotopic data for methane in free (n= 100) or dissolved (n= 2) gas phases or both (n= 33) (Table 1).

2.2.1 Major and Minor Ion analysis

To collect samples representative of aquifer conditions, the groundwater wells were purged until the field parameters pH, redox, dissolved oxygen, temperature, and electrical conductivity stabilized. Alberta Innovates Technology Futures (AITF) conducted the major and minor ion chemistry analyses on filtered samples (0.45µm) that were acidified to pH < 2 for cation analysis and non-acidified for major anion determination. ICP-MS analysis was used to determine cation concentrations while titration for alkalinity and ion chromatography were used to determine anion concentrations. The detection limits are indicated in Table 2 and concentrations are expressed in mol/L (M).

2.2.2 Gas composition

A detailed description of the sampling equipment and procedures for free and dissolved gas samples is given in Humez et al. (2015). The composition of free and dissolved gas samples was determined in the laboratory by gas chromatography yielding concentrations for oxygen, carbon dioxide, methane and higher alkane chain compounds (such as ethane) with measurements conducted by AITF with uncertainties of ± 5% of the analytes. Gas composition data for free gas samples is reported in parts per million by volume (ppmv) and for dissolved gases expressed in mol/L (M) or mg/L to ensure comparability with other studies. The gas dryness parameter defined as the ratio between methane / higher n-alkanes was also determined.

2.3 Isotopic analyses

$\delta^{13}\text{C}_{\text{CH}_4}$, $\delta^2\text{H}_{\text{CH}_4}$, $\delta^{13}\text{C}_{\text{DIC}}$, $\delta^{34}\text{S}_{\text{SO}_4}$, $\delta^{18}\text{O}_{\text{SO}_4}$, $\delta^{15}\text{N}_{\text{NO}_3}$, $\delta^{18}\text{O}_{\text{NO}_3}$, $\delta^2\text{H}_{\text{H}_2\text{O}}$ and $\delta^{18}\text{O}_{\text{H}_2\text{O}}$ values were analyzed in the Isotope Science Laboratory at the University of Calgary. Stable isotope ratios are reported in the internationally accepted delta notation (‰) relative to VPDB for $\delta^{13}\text{C}$ values, VSMOW for $\delta^2\text{H}$ and $\delta^{18}\text{O}$ values, VCDT for $\delta^{34}\text{S}$ values and N_2 in air for $\delta^{15}\text{N}$ values. All carbon and hydrogen isotope analyses on methane and CO_2 were conducted on a ThermoFisher MAT 253 isotope ratio mass spectrometer (IRMS) coupled to a Trace GC Ultra + GC Isolink (ThermoFisher). The precision for carbon isotope analyses was better than ± 0.5‰ for hydrocarbons and better than ± 0.2‰ for carbon dioxide. The precision for hydrogen isotope analysis of hydrocarbons was better than 3%. Water isotope analyses were performed by Off-axis Cavity Ringdown Spectroscopy using a Los Gatos Water Isotope Analyzer (DLT-100). Precision was better than ± 2 ‰ for $\delta^2\text{H}$ and ± 0.2 ‰ for $\delta^{18}\text{O}$.

To determine the isotopic composition of sulfate, dissolved sulfate was converted to barium sulfate (BaSO_4) and subsequently analyzed using a ThermoQuest Finnigan Delta^{plus}XL IRMS coupled with either a Fisons NA 1500 Elemental Analyzer for $\delta^{34}\text{S}_{\text{SO}_4}$ analysis or a HEKAtech HT Oxygen Analyser with Zero-blank autosampler for $\delta^{18}\text{O}_{\text{SO}_4}$ analysis. Precision for $\delta^{34}\text{S}_{\text{SO}_4}$ and $\delta^{18}\text{O}_{\text{SO}_4}$ is ± 0.5 ‰.

The isotopic composition of nitrate was determined on N_2O generated by the denitrifier technique (c.f. Silva et al., 2000;

Sigman et al., 2001; Casciotti et al., 2002) using a Thermo Delta V Plus IRMS coupled with a Finnigan MAT PreCon. Precisions of $\delta^{15}\text{N}_{\text{NO}_3}$ and $\delta^{18}\text{O}_{\text{NO}_3}$ are $\pm 0.3 \text{ ‰}$ and $\pm 0.7 \text{ ‰}$ respectively.

The analytical results for all groundwater samples were further investigated using PHREEQC (Parkurst and Appelo, 1999) to assess geochemical speciations, potential redox values (pe), ionic balance ($< \pm 10 \%$), among others. SPSS 22 was used for determining descriptive statistics such as median, mean, range, standard deviation and to evaluate the correlation between variables. Pearson correlation analysis was conducted where linear trends between two variables existed. When nonlinear relationship between two variables existed or in presence of outliers, Spearman's rho and Kendall's tau tests were used instead (Humez et al., 2016).

3 Results

10 3.1 Field Parameters

During sampling of shallow groundwater in the field, temperature, electrical conductivity, dissolved oxygen content and oxidation-reduction potential (ORP) were determined for all water samples. The average groundwater temperature was $7 \pm 3 \text{ }^\circ\text{C}$, while the average pH value was 7.8. The electrical conductivity of the groundwater samples ranged from 212 to $>16,000 \text{ }\mu\text{S/cm}$ with an average value of $1634 \text{ }\mu\text{S/cm}$. Fourteen groundwater samples had dissolved oxygen concentrations $> 0.06 \text{ mM}$ ($> 2 \text{ mg/L}$), 56 water samples had dissolved oxygen concentrations ranging between 0.01 and 0.06 mM (0.5 and 2.0 mg/L), and 129 samples had dissolved oxygen concentrations $<0.01 \text{ mM}$ ($<0.5 \text{ mg/L}$). For water samples with redox potential reported, 70 samples had Eh values $< 0 \text{ mV}$ ($\text{Eh} = 0.059 \cdot \text{pe}$ (Volt) with $\text{pe} = \log(e^-)$).

3.2 Major ion concentrations and hydrochemical water type classification

20 Major ion chemistry of groundwater and methane concentrations were determined for 225 groundwater samples from shallow aquifers and results are summarized in Table 2. Chloride concentrations ranged from 0.01 to 68.01 mM with mean and median values of 2.81 mM and 0.40 mM ($n=225$). Sulfate concentrations ranged from 4.55 μM to 74.16 mM with mean and median values of 2.83 mM and 0.72 mM ($n=225$) respectively. DIC concentrations ranged from 0.81 mM to 39.07 mM with mean and median values of 12.94 and 12.29 mM ($n= 225$) respectively. Nitrate concentrations ranged from 0.21 μM to 21.2 mM ($n=136$). For 90 samples, NO_3 concentrations have not been reported but the ion balance is acceptable so that NO_3 represents $\ll 10 \%$ of the anions.

The major cations Na, Ca, Mg, K showed a wide range of concentrations (Table 2). Sodium concentrations ranged from 0.03 mM to 165.78 mM with mean and median values of 17.28 mM and 12.76 mM ($n=225$) respectively. Calcium concentrations ranged from 0.01 mM to 9.08 mM with mean and median values of 0.94 mM and 0.40 mM ($n=136$) respectively. For 89 samples, Ca concentrations were not measured but the ion balance is acceptable so that Ca represents $<10\%$ of the cations (set to 0 for the Piper plot). Magnesium concentrations ranged from 2.26 μM to 12.45 mM with mean and median values of

0.60 mM and 0.05 mM (n=225) respectively. Potassium concentrations ranged from 0.01 mM to 0.54 mM with mean and median values of 0.07 mM and 0.05M (n=225) respectively. Total dissolved solids (TDS) were calculated by [summing](#) major ion concentrations. Total dissolved solids (TDS) ranged from 180 to 15,500 mg/L with an average value of 1264 mg/L.

5 The Piper plot in Fig. 2 shows that water types were found to be highly variable ranging from Ca-Mg-HCO₃ to Na-Cl types with the following order Na-HCO₃ (59.1%) > Na-HCO₃-Cl (17.0%) > Ca-HCO₃ (8.0%) > Na-Cl (5.8%) > Ca-Na-HCO₃ (5.3%) > Ca-HCO₃-Cl (2.2%) > Ca-Na-HCO₃-Cl (1.3%) > Ca-Na-Cl (0.9%) > Ca-Cl (0.4%). Elevated concentrations of methane in groundwater were found predominantly in Na-Cl, Na-HCO₃ and Na-HCO₃-Cl water types (see color code in Fig. 2).

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3.3 Methane, ethane and propane occurrence in shallow groundwater

Methane was detected above the limit of detection (DL) in all samples for which free gas analyses were available. The average methane concentration was 265,466 ppmv (n= 147). Twenty-five percent of the samples had methane concentrations
15 > 390,000 ppmv (third quartile, Q₃). In dissolved gas samples, the average methane concentration was 0.43 mM (n=221). Twenty-five percent of the samples had a dissolved methane concentration >0.4 mM (third quartile, Q₃). The highest methane concentration was 3.01 mM (Table 2).

In free gas samples, the average ethane (C₂H₆) concentration was 215 ppmv (n=96) with a maximum value of 3650 ppmv. In dissolved gas samples, the average ethane concentration was 0.60 μM with a maximum of 17.63 μM. In free gas samples, the
20 average detected propane (C₃H₈) concentration was 0.67 ppmv (n=36) with a maximum value of 4.60 ppmv. In dissolved gas samples, the average propane concentration was 0.03 μM with a maximum of 0.90 μM (Table 2).

Figure 3a shows that elevated dissolved methane concentrations were generally found at redox potentials (Eh) below 0 mV. Dissolved CH₄ concentrations and Eh values are weakly inversely correlated (Kendall's *tau* = -0.106 and p < 0.05, Spearman's *rho* = -0.167 and p < 0.05). One triplicate sample from a well located between Calgary and Red Deer had an
25 elevated pe value while the methane concentrations were > 0.5 mM (Fig 3a). The highest methane concentrations in dissolved gas samples occurred at pH values > 7 and at low Eh < 0 mV (Fig. 3b).

A cross-plot of average TDS contents versus water type reveals that the highest TDS values were associated with Na-Cl, Ca-Cl and Na-HCO₃ water types, whereas Ca-HCO₃ waters had the lowest average TDS content (Fig. 4a). A comparison of methane occurrences and water types revealed that dissolved methane occurs predominantly in Na-HCO₃ waters for 133
30 samples out of 221 (Fig. 2, Fig. 4b for dissolved methane). In dissolved gas samples, the highest average methane concentrations of > 1 mM were observed in Na-Cl, Na-HCO₃-Cl and Na-HCO₃ water types, while in all other water types average methane concentrations ranged between 0.07 to 78 μM (Fig. 4c). In free gas samples, the highest average methane concentrations of >260,000 ppmv were also observed in Na-Cl, Na-HCO₃-Cl and Na-HCO₃ waters types, while in all other

water types average methane concentrations ranged between 400 and 70,753 ppmv on average (Fig. 4d). The majority of free gas samples with methane concentrations $> 150,000$ ppmv ($n = 65$ out of 147) and > 0.5 mM ($n=53$ out of 221) in dissolved gas samples were associated with the Na-HCO₃ water-type. However, Fig. 4c,d consistent with Fig. 2 reveal also 5 exceptional samples with high methane concentrations in dissolved gas ($n=5$), free gas, or both phases ($n=3$) occurring in Ca-Na-HCO₃ (#6) and Ca-HCO₃ (#9) water types (see Sect. 4.4).

Ethane was also observed in some dissolved and free gas samples. In dissolved gas samples, ethane concentrations > 0.3 μ M were only observed in groundwater of the Na-Cl, Na-HCO₃-Cl and Na-HCO₃ water types. Only one exception circled in Fig. 4e of a sample with elevated ethane content (0.9 μ M) was observed in a Ca-HCO₃-Cl type sample (see Sect. 4.4). In free gas samples, ethane concentrations >100 ppmv were only observed in groundwater of the Na-Cl, Na-HCO₃-Cl and Na-HCO₃ water types with average concentrations of 79, 603, and 160 ppmv respectively (Fig. 4f). Only one exceptional sample containing ethane in free gas at 73 ppmv was found in Ca-Na-HCO₃ type water (Fig. 4f).

Few samples contained propane with the highest concentration of dissolved propane (0.9 μ M) occurring in the Ca-HCO₃-Cl water type and lower propane concentrations found in the Na-Cl (0.01 μ M, $n= 5$) $>$ Na-HCO₃-Cl (0.006 μ M, $n=9$) $>$ Ca-HCO₃ (0.003, $n=2$) $>$ Na-HCO₃ (0.002 μ M, $n=15$) water types. For free gas samples, propane was found in Na-HCO₃-Cl (1.3 ppmv, $n=8$) $>$ Ca-HCO₃ (0.6 ppmv, $n=3$) $>$ Na-HCO₃ (0.5 ppmv, $n=21$) $>$ Na-Cl (0.3 ppmv, $n=4$) water types.

Hence, there appears to be a correlation between the water type and the number of samples containing elevated methane and ethane concentrations in the shallow groundwaters of Alberta and a consistent relationship between gases in dissolved and free phases as shown in Humez et al. (2016).

20 3.4 Isotopic composition of groundwater

The $\delta^{18}\text{O}$ and $\delta^2\text{H}$ values of groundwater varied from -24.3 to -8.4 ‰ with an average of -18.5 ± 1.9 ‰ and from -190.8 to -94.2 ‰ with an average of -147.4 ± 13.1 ‰, respectively ($n=222$) (Fig. 5). Hydrogen and oxygen isotope values of all water samples plotted close to the local meteoric water lines (LMWL) of Edmonton and Calgary (Peng et al., 2004) suggesting atmospheric recharge of groundwater with at most minor influence of evaporation and water-rock interactions on the isotopic composition of the groundwater.

3.5 Isotopic composition of dissolved constituents in groundwater

3.5.1 Sulfur and Oxygen Isotope Ratios of Sulfates

The $\delta^{34}\text{S}_{\text{SO}_4}$ values in groundwater ranged from -26.6 to $+40.9$ ‰ with a mean value of $+1.8 \pm 12.4$ ‰ ($n=158$). The $\delta^{18}\text{O}_{\text{SO}_4}$ values in groundwater ranged from -17.7 to $+11.2$ ‰ with a mean value of -0.6 ± 6.7 ‰ ($n=138$) (Table 2).

3.5.2 Carbon Isotope Ratios of Dissolved Inorganic Carbon

The partial pressure of CO₂ ($p\text{CO}_2$) was calculated for all samples based on field pH and alkalinity based on geochemical speciation with PHREEQC (Parkurst and Appelo, 1999). P_{CO_2} values ranged between $10^{-8.14}$ and $10^{+0.58}$ atm and the pH values in the samples containing methane ranged from 6.5 to 10.2 with a mean value of 7.9 ± 0.8 (n=225). The $\delta^{13}\text{C}_{\text{DIC}}$ values ranged from -30.8 ‰ to elevated values of $+21.2$ ‰ with an average of -10.8 ± 8.7 ‰ (n=221). The highest $\delta^{13}\text{C}_{\text{DIC}}$ values of $+21.2$ ‰, $+17.7$ ‰, $+15.3$ ‰ and $+14.3$ ‰ occurred in samples with elevated methane concentrations in dissolved and free gases of >1 mM and $>900,000$ ppmv respectively.

3.5.3 Nitrogen and Oxygen Isotope Ratios of Nitrates

10 | Only 24 samples contained sufficient nitrate for isotope analysis (Table 2). The $\delta^{15}\text{N}_{\text{NO}_3}$ values varied from -10.4 to $+21.8$ ‰ with an average of $+7.8 \pm 8.2$ ‰, while $\delta^{18}\text{O}_{\text{NO}_3}$ values ranged from -13.2 to $+25.7$ ‰ with an average of -1.5 ± 12.0 ‰.

3.6 Isotopic composition of methane

Methane in 133 groundwater samples including replicates had a median $\delta^{13}\text{C}$ value of -66.2 ‰ with a minimum value of -92.8 ‰, a maximum of -20.5 ‰, and a mean value of -64.6 ± 14.9 ‰ in free gas phase (Table 2). The median $\delta^{13}\text{C}$ value of methane in dissolved gas samples was -65.6 ‰ with a minimum value of -85.5 ‰, a maximum of -35.8 ‰ and a mean value of -65.0 ± 10.5 ‰ (n=35) (Table 2). Fifty-eight groundwater samples had a median $\delta^2\text{H}_{\text{CH}_4}$ value of -291.5 ‰ with a minimum value of -437.1 ‰ and a maximum of -80.9 ‰ in free gas phase (Table 2).

4 Discussion

4.1 Geochemical constraints of methane-containing groundwater

20 | Figure 5 shows that aqueous geochemistry results of the groundwater samples can be explained by three end-member compositions and their respective mixtures. The first group is described by (i) groundwater samples with low chloride, sulfate, sodium, DIC concentrations and low to intermediate $\delta^{18}\text{O}_{\text{H}_2\text{O}}$ and $\delta^2\text{H}_{\text{H}_2\text{O}}$ values, representing a freshwater (TDS < 2000 mg/l) end-member (plotting close to the origin in Fig. 5, named end-member #1). Samples belonging to this group had generally low methane concentrations of <0.001 mM and a wide range of nitrate concentrations. The second group
25 | represents (ii) groundwater samples with low chloride concentrations and low $\delta^{18}\text{O}_{\text{H}_2\text{O}}$ and $\delta^2\text{H}_{\text{H}_2\text{O}}$ values but high sodium, sulfate and DIC concentrations (blue shading in Fig. 5). Samples belonging to this group had TDS > 4000 mg/L but had mostly low methane concentrations. These samples were predominantly obtained from GOWN wells located to the east of the Edmonton-Lethbridge corridor. Only one water sample in this group had an elevated nitrate concentration of > 4 mM and

two other water samples had a nitrate concentration < 0.02 mM. These three samples were taken from wells completed in the Horseshoes Canyon, Bearpaw and Belly River Formations. The samples from this group (ii) appear to be impacted by sulfide oxidation in tills and cation exchange resulting in elevated Na, DIC and sulfate concentrations (Grasby et al., 2010). The third identified group is (iii) groundwater with elevated chloride, sodium and DIC concentrations, elevated $\delta^{18}\text{O}_{\text{H}_2\text{O}}$ and $\delta^2\text{H}_{\text{H}_2\text{O}}$ values, and TDS values between 2000 and 4000 mg/L, but with negligible nitrate (< 0.002 mM) and sulfate concentrations (< 1 mM) (green shading in the Fig. 5). This group is composed of Na-HCO₃ and Na-HCO₃-Cl water-types and contains the groundwater with the highest methane concentrations. The samples in this group were obtained predominantly from wells completed in coal-bearing geological formations (e.g. Belly River and Horseshoes Canyon Formations).

The groundwater compositions investigated in this study can thus be explained by mixing between the high TDS Na-HCO₃ and Na-HCO₃-Cl end-member #3 and freshwater end-member #1 and/or mixing between the freshwater end-member #1 and end-member #2 (Fig. 5).

Results presented in Figs. 2 and 4 show that water samples with elevated methane and ethane concentrations are predominantly associated with groundwaters of the Na-HCO₃ water-type. This is consistent with the observations made in groundwater baseline studies of [McIntosh et al. \(2014\)](#) and [Hamilton et al. \(2015\)](#) in south-western Ontario (Canada), [Molofsky et al. \(2013\)](#) in Susquehanna County (Northeastern Pennsylvania, USA) and [McPhillips et al. \(2014\)](#) in Chenango country (Central New York State, USA). In [these](#) studies, elevated methane concentrations (dissolved CH₄ > 1 mg/L or > 0.06 mM) in groundwater were predominantly found in sodium-chloride (Na-Cl) or sodium-bicarbonate (Na-HCO₃) water types, while calcium-bicarbonate (Ca-HCO₃) type groundwater had typically very low methane concentrations or no methane at all. This is consistent with the majority of the observations made in this study.

We observe that groundwater samples with elevated methane and ethane concentrations **are associated** predominantly **with** end-member #3 with negligible nitrate and sulfate concentrations (green shading in Fig. 5). Therefore, the redox status of the various groundwaters was further investigated, to better constrain the conditions that may facilitate the formation and occurrence of methane and other alkane gases in shallow groundwater bodies.

4.2 Redox-sensitive parameters and the distribution of methane in groundwater samples

As groundwaters evolve from highly oxidized to highly reducing conditions, they undergo a sequence of redox reactions including O₂ consumption, denitrification, Mn- and Fe-reduction, bacterial sulfate reduction followed by methanogenesis (Appelo and Postma, 2005). Since Mn and Fe concentration data were not available in our data set, the groundwater samples collected in this study were classified into four categories depending on the concentrations of terminal electron acceptors (TEAPs), such as O₂, NO₃, Mn, Fe, and SO₄, participating successively in redox reactions:

- 1) oxidized;
- 2) denitrified but still sulfate containing;
- 3) undergoing bacterial sulfate reduction, and;

4) methanogenic

Figure 6a shows that typically samples with elevated oxygen concentrations > 0.01 mM ($P_{O_2} < 10^{-1.5}$ atm) do not contain methane with the exception of 16 of 221 samples. In-situ methane formation is not possible in oxygen-containing groundwater (Chapelle, 2001). Only one sample had an elevated O_2 content of 0.2 mM in concert with a dissolved methane concentration of 0.7 mM (Fig 6a) potentially indicating methane migration from more reducing aquifer portions (see [section 4.4](#)).

Figure 6b shows that samples containing nitrate (> 0.006 mM) did not contain any dissolved methane. A significant inverse correlation was found between nitrate and methane concentrations (Kendall's $\tau = -0.167$ and $p < 0.05$, Spearman's $\rho = -0.228$ and $p < 0.05$, $n=133$) suggesting that nitrate-containing groundwaters are not suitable for methane formation or conservation. Figure 6c shows that only groundwater samples with sulfate concentrations lower than 1 mM contained elevated dissolved methane. A significant inverse correlation was found between sulfate and methane concentrations (Kendall's $\tau = -0.320$ and $p < 0.05$, Spearman's $\rho = -0.463$ and $p < 0.05$, $n=221$) suggesting that methane formation does not commence while sulfate is still present at concentration > 1 mM. An alternate explanation could be that methane migrated into the aquifers containing O_2 , NO_3 and/or SO_4 thereby creating more reducing conditions and consuming oxygen from all these species. We suspect that noticeable amounts of methane in dissolved and free gas samples were only observed after sulfate had been removed presumably by bacterial (dissimilatory) sulfate reduction to levels < 1 mM (Fig. 6c). [This result is consistent with what has been observed in deeper coalbed methane and organic-rich shale microbial gas systems \(Schlegel et al., 2011\)](#). These observations are consistent with the redox ladder concept and are further illustrated in Fig. 7.

Figure 7 shows a cross-plot of nitrate and sulfate concentrations with concentrations of dissolved methane displayed as colored circles revealing that groundwater samples containing elevated concentrations of nitrate (>0.006 mM) and intermediate to high sulfate (1 to 1000 mM) concentrations did not contain significant amounts of dissolved methane (<0.01 mM) (light grey shading, Fig. 7). The presence of nitrate and sulfate indicates that neither complete denitrification nor complete bacterial sulfate reduction has occurred in these aquifers and hence in-situ methane formation would be in contradiction to the redox ladder concept. In this group, only four samples had non-negligible methane concentrations of $0.06 < CH_4 < 1.2$ mM (circled in Fig. 7; see [section 4.4](#)). This is consistent with the hypothesis that methane migrated into aquifers containing NO_3 and SO_4 .

Another group of groundwater samples is characterized by elevated (> 1 mM) sulfate concentrations, low nitrate, and negligible amounts of dissolved methane (<0.01 mM) (hatched area, Fig. 7). Nearly all these samples had negligible methane concentrations except five samples (circled in Fig. 7 within the hatched area) with elevated methane concentrations > 0.01 mM. There are two possible reasons which could explain the coexistence of methane with sulfate concentrations slightly above 1 mM; either these groundwater samples never contained nitrate but the presence of sulfates indicates that bacterial sulfate reduction has not occurred yet and hence methanogenesis has not commenced; alternatively, methane may have migrated into some of these aquifers and was oxidized through denitrification explaining the lack of nitrate.

A third group of samples has negligible concentrations of nitrate (< 0.006 mM) and sulfate (< 1 mM) and contains the vast majority of samples with methane concentrations >0.1 mM including those with the highest methane content of >1.2 mM (dark grey shading, Fig. 7). This suggests that both denitrification and bacterial sulfate reduction have occurred creating redox conditions favorable for in-situ methanogenesis. These conditions were predominantly observed in Na-HCO₃ and Na-HCO₃-Cl water-types with one exception in Ca-HCO₃ water (circled dot in dark grey shading, Fig. 7).

4.3 Stable isotopes constraints

To further test the hypotheses of occurrences of redox processes such as denitrification, bacterial sulfate reduction, methanogenesis, and potentially methane oxidation, we investigated the isotopic compositions of nitrate, [sulfate](#), [DIC](#) and [methane](#).

4.3.1 Nitrogen and Oxygen Isotope Ratios of Nitrates

The isotopic composition of nitrate indicates predominantly the sources of this nutrient (Kendall, 1998). In one group of groundwater samples with both low $\delta^{15}\text{N}_{\text{NO}_3}$ ($< +10$ ‰) and $\delta^{18}\text{O}_{\text{NO}_3}$ values (< 0 ‰) accompanied by low nitrate concentrations (<0.06 mM) nitrate appears to be derived from nitrification of soil organic matter (Fig. 8b) (Kendall, 1998). These groundwaters are characterized by mainly low methane concentrations (<0.05 mM) except for two samples with methane concentrations of 0.2 mM and 1.3 mM. A second group of groundwater samples had low $\delta^{18}\text{O}_{\text{NO}_3}$ values of nitrate (-10 ‰ $< \delta^{18}\text{O}_{\text{NO}_3} < +5$ ‰) but elevated $\delta^{15}\text{N}_{\text{NO}_3}$ values ($> +10$ ‰, Fig. 8b). These samples were also associated with the highest nitrate concentrations of up to 21.2 mM suggesting nitrate contamination most likely from manure spreading (Rock and Mayer, 2004). These samples belong to various groundwater types and had negligible methane concentrations <0.01 mM. The third group is described by elevated $\delta^{15}\text{N}_{\text{NO}_3}$ ($> +5$ ‰) and $\delta^{18}\text{O}_{\text{NO}_3}$ values ($> +20$ ‰), negligible nitrate concentrations (<200 μM) and methane concentrations up to 0.3 mM. Such an isotopic signature could theoretically be sourced from NO₃-containing mineral fertilisers (Kendall et al., 1998) but in this case elevated nitrate concentrations would be expected. During denitrification in a closed system, it is expected that as nitrate concentrations decrease the remaining nitrate becomes progressively enriched in ¹⁵N and ¹⁸O (Mariotti et al., 1988; Boettcher et al., 1990). Plotting nitrate concentrations versus $\delta^{15}\text{N}$ values of nitrate containing samples provides some evidence that these samples may have been affected by denitrification (Fig. 8a). Hence, isotope analyses revealed different sources of nitrate and processes such as mixing between nitrification-derived and manure-derived end-members, but only little indication of denitrification. Only 24 of 225 samples had sufficient nitrate to conduct isotope analyses and no methane was observed in samples with elevated nitrate concentrations (Fig. 8a).

4.3.2 Sulfur and Oxygen Isotope Ratios of Sulfates

During bacterial (dissimilatory) sulfate reduction (BSR) in a closed system, it is expected that sulfate concentrations decrease while ^{34}S and ^{18}O become progressively enriched in the remaining sulfate (Fritz et al., 1989). Plotting sulfate concentrations versus $\delta^{34}\text{S}_{\text{SO}_4}$ values (Fig. 9a) reveals that samples with the highest sulfate concentration have $\delta^{34}\text{S}_{\text{SO}_4}$ values between 0 and -10‰ . This, together with $\delta^{18}\text{O}_{\text{SO}_4}$ values $< +4\text{‰}$ suggests that these groundwater samples derive their sulfate predominantly from pyrite oxidation (Fig. 9b, Grasby et al., 2010). Many samples with lower sulfate concentrations also show $\delta^{34}\text{S}$ and $\delta^{18}\text{O}$ values of sulfate $< 0\text{‰}$ suggesting that oxidation of sulfide minerals is the sulfate source (Fig. 9b), creating a mixing trend in Fig. 9a between a low- SO_4 content groundwater with high- SO_4 concentration samples. There is, however, also a trend of increasing $\delta^{34}\text{S}_{\text{SO}_4}$ values $> +15\text{‰}$ with decreasing sulfate concentrations (Fig. 9a) suggesting that bacterial sulfate reduction has occurred in these aquifers with the highest $\delta^{34}\text{S}$ value of $+56.4\text{‰}$ at a sulfate concentration of 0.03 mM . This is also confirmed by elevated $\delta^{18}\text{O}$ values of sulfate (Fig. 9b). The group of samples displaying evidence of BSR (Fig. 9b) contains many samples with elevated methane concentrations ($>0.1\text{ mM}$).

4.3.3 Carbon isotope ratios of inorganic carbon

The isotopic composition of DIC is indicative of sources of carbon and processes that have generated or affected DIC (Mook, 2000). Fig. 10a reveals that the $\delta^{13}\text{C}$ values of DIC of most samples varied between -20 and -10‰ , suggesting that the majority of the DIC is derived from a combination of oxidation of organic carbon and carbonate dissolution (Clark and Fritz, 1997). Most samples in this category have methane concentrations $<0.1\text{ mM}$. Fig. 10a further reveals a second group of samples with $\delta^{13}\text{C}$ values of DIC $> -8\text{‰}$ and reaching values as high as $+20\text{‰}$. The positive $\delta^{13}\text{C}$ values are clear evidence for in-situ biogenic methanogenesis within the aquifer, during which ^{12}C is preferentially allocated to methane while the remaining CO_2 and subsequently DIC becomes enriched in ^{13}C (Barker and Fritz, 1981). All samples in this category had methane concentrations $> 1.2\text{ mM}$.

Fig. 10b shows that all samples with elevated sulfate concentrations contained DIC with low $\delta^{13}\text{C}_{\text{DIC}}$ values (average -13.8‰). In contrast, samples with the lowest sulfate concentrations were accompanied by the highest $\delta^{13}\text{C}_{\text{DIC}}$ values of up to $+21.2\text{‰}$ and the highest methane concentrations, while samples with methane concentrations between 0.1 and 1.2 mM plot in between (Fig. 10b). This strongly supports the hypothesis that BSR needs to proceed towards completion prior to commencement of in-situ methanogenesis with the aquifer and generation of elevated methane concentrations in the aquifer.

4.4 Evidence for in-situ formation, for migration, and for oxidation of methane

4.4.1 Classification criteria

Using the information described above, we evaluated whether geochemical conditions in the aquifers were suitable for in-situ methane generation, or whether the geochemical conditions suggest that methane must have migrated into the aquifer.

5 This was achieved for 135 samples that had sufficient aqueous and gas geochemistry data including the following information and parameters:

- (1) Water type derived from the Piper diagram based on balanced major ion chemistry,
- (2) Redox parameters such as dissolved oxygen, sulfate, and nitrate concentrations,
- (3) Gas composition in dissolved and/or free gas samples,
- 10 (4) Isotope values of methane in free and/or dissolved gas samples,
- (5) Geological formation in which the groundwater wells were completed.

4.4.2 In-situ biogenic methane generation (category #1)

Category #1 contains samples with $\delta^{13}\text{C}_{\text{CH}_4} < -55 \text{ ‰}$ and a high dryness parameter >1000 indicating biogenic methane. Aqueous geochemistry data were consistent with methanogenic conditions (no nitrate, sulfate concentrations negligible) and
15 no traces of propane were detected (Table 3). This category contains 53 of 135 samples (39 %) yielding clear evidence that biogenic methane was generated in-situ under methanogenic aquifer conditions (Fig. 11). Elevated methane and ethane concentrations were found usually in aquifers completed in the coal- and shale-bearing geological formations (e.g. Horseshoes Canyon and Belly River Group Formations). These samples are classified as CH₄-type A in Table 3, and Fig. 13.

20 The remaining 82 samples that did not fall into category #1 show at least one of the following characteristics:

- 1) Presence of traces of propane in dissolved and/or free gas (n = 31 for free gas, n=22 for dissolved gas);
- 2) Dryness parameter < 500 (n=23);
- 3) Elevated methane concentrations ($>0.01 \text{ mM}$), while oxygen ($>0.01 \text{ mM}$), sulfate ($>1 \text{ mM}$), and/or nitrate ($>0.006 \text{ mM}$) concentrations are not negligible (n = 6);
- 25 4) A carbon isotope ratio that may suggest thermogenic methane ($\delta^{13}\text{C}_{\text{CH}_4} > -55 \text{ ‰}$) (n = 24).

These characteristics may indicate that methane has migrated and potentially has undergone oxidation.

4.4.3 Migration of biogenic methane into more oxidizing aquifer sections (category #2)

30 Category #2 contains samples with $\delta^{13}\text{C}_{\text{CH}_4} < -55 \text{ ‰}$, elevated dryness parameter >1000 , and no traces of propane, indicating biogenic methane. However, Table 3 reveals that methane was detected in groundwater with either elevated sulfate

concentrations (> 1 mM, $n=3$) or elevated nitrate concentrations (> 0.006 mM, $n=2$). This is inconsistent with conditions suitable for in-situ methanogenesis. Therefore it is postulated that biogenic methane had migrated from more reducing sections of the aquifer into sections with more oxidizing conditions and has not yet been oxidized [since there is no evidence of methane oxidation such as low \$\delta^{13}\text{C}_{\text{DIC}}\$ values as expected for methane oxidation, likely](#) due to [short](#) residence times with respect to the rather slow turnover of microbial [anaerobic methane oxidation \(AOM\)](#) (Jorgensen et al., 2001). This category contains 4 % of the samples (5 of 135 samples) and is listed as CH_4 -type B in Table 3 and Fig. 13.

4.4.4 Apparent or pseudo-thermogenic methane in shallow aquifers (category #3)

Category #3 contains samples with $\delta^{13}\text{C}_{\text{CH}_4} > -55$ ‰ (Fig. 11) but without detectable higher alkanes. Such high $\delta^{13}\text{C}$ values can either indicate a thermogenic gas source or may be caused by methane oxidation. An increase of $\delta^{13}\text{C}_{\text{CH}_4}$ values was observed with decreasing $\delta^{13}\text{C}_{\text{DIC}}$ values (all < -5 ‰) and sulfate concentrations decreased with increasing $\delta^{34}\text{S}_{\text{SO}_4}$ values (-10 ‰ $< \delta^{34}\text{S}_{\text{SO}_4} < +15$ ‰) (Fig. 12). The very low methane concentrations and the absence of higher alkanes do not support a significant flow of thermogenic gas from deep geological formations below the aquifers. Instead, the data indicate that biogenic methane has been oxidized within the aquifers, possibly coupled with bacterial sulfate reduction (Fig. 12). This process enriches ^{13}C in the remaining methane (Barker and Fritz, 1981) imparting a [relatively high](#) $\delta^{13}\text{C}$ value that can be misinterpreted as indicating a thermogenic gas signature. The occurrence of methane oxidation is further confirmed by a cross-plot of $\delta^{13}\text{C}$ values of methane and those of CO_2 (Fig. 11). It is also possible that post-sampling degradation of low-methane samples occurred [e.g. potential slow diffusive gas loss from sampling containers resulting in \$^{13}\text{C}\$ enrichment in the residual methane](#). Hence we conclude that all the samples in category #3, corresponding to CH_4 -type D (Table 3), are either affected by methane oxidation or in some cases possibly by increased analytical uncertainty due to low methane concentrations. The elevated $\delta^{13}\text{C}$ values of methane are therefore not indicative of leakage of thermogenic methane from deeper portions of the stratigraphic column into shallow aquifers. Category #3 contains 13 % of the samples (17 of 135).

4.4.5 Thermogenic-biogenic mixed gas origin (category #4)

Category #4 contains samples with non-negligible concentrations of higher alkanes (e.g. ethane and propane) and low dryness parameter values. For all these samples, aqueous geochemistry results suggested methanogenic conditions with no nitrate and negligible sulfate concentrations. This category contains 40 % of the samples (55 of 135) and is further subdivided into 3 sub-categories (Table 3).

Sub-category #4.1 contains two samples with $\delta^{13}\text{C}_{\text{CH}_4}$ values > -55 ‰ and with very low methane concentrations and traces of propane. This suggests that mixed thermogenic and biogenic gas may have migrated into overlying aquifers and may have undergone partial methane oxidation as supported by Fig. 11 where these two samples plot in the methane oxidation field.

These samples were derived from groundwater of the Paskapoo Formation and hence are also classified as CH₄-type D (Table 3).

Sub-category #4.2 contains 38 samples with elevated ethane concentrations and traces of propane. All samples from sub-category #4.2 were obtained from groundwater wells completed in coal-bearing formations (e.g. Horseshoe Canyon and Belly River Formations). Five samples had $\delta^{13}\text{C}_{\text{CH}_4}$ values > -55 ‰ while 33 samples had $\delta^{13}\text{C}_{\text{CH}_4}$ values < -55 ‰. Cheung et al. (2010) reported that gases derived from the Horseshoe Canyon Formation in southeastern Alberta contained considerable amounts of ethane up to 4000 ppm in addition to methane with an average $\delta^{13}\text{C}$ value of -54.0 ± 4.1 ‰. This suggests that gases from the coal-bearing Horseshoe Canyon Formation contain a minor thermogenic gas component (Cheung et al., 2010). Hence the minor thermogenic gas components detected in 38 samples of this sub-category #4.2 (28.1%) appear to be mainly derived from shallow coal-bearing sedimentary units such as the Horseshoe Canyon Formation in which many of the groundwater wells are completed, and hence in-situ gas is sampled. This in-situ gas is referred to CH₄-type A* (Table 3, Fig. 13), since it is predominantly biogenic with only traces of thermogenic components in this mixed gas. An alternate explanation for the occurrence of ethane and propane is their microbial formation via ethanogenesis and propanogenesis (Hinrich et al., 2006). We consider this less likely since the microorganisms responsible for biological ethane and propane formations have not yet been identified.

Sub-category #4.3 contains samples with $\delta^{13}\text{C}_{\text{CH}_4}$ values < -55 ‰ and a dryness parameter of < 500 or traces of propane. This sub-category contains 15 samples obtained from wells all completed in non-CBM formations. This indicates a gas that is composed of biogenic methane mixed with smaller portions of thermogenic gas was found in shallower stratigraphic units such as the Paskapoo Formation and in surficial deposits, suggesting that mixed gas has migrated upwards. These samples account for 11.1% of the investigated groundwater and are listed as CH₄-type C in Table 3, and Fig. 13.

No water samples in category #4 required admixture of deep ($>1,000$ m) thermogenic gases to explain the chemical and isotopic characteristics of dissolved and free gas samples.

4.4.6 Samples with inconclusive data sets (category #5)

Five samples (4 %) could not be assigned to any of the above categories due to conflicting parameters in the aqueous or gas chemical and isotopic data sets or between replicate samples.

Conclusion

Analysis of water types suggested that methane occurs predominantly in Na-HCO₃ or Na-(HCO₃)-Cl type waters possibly indicating prolonged water-rock interaction or mixing with less mobile saline water. Taking into account the hydrochemical conditions in methane-bearing aquifers allows for a refined analysis of methane sources and a differentiation between biogenic in-situ production of methane within aquifers versus migration of biogenic or thermogenic gases into the aquifer. To achieve this we combined redox-sensitive aqueous geochemistry parameters and isotopic compositions of nitrate, sulfate

and DIC with the interpretation of [natural](#) gas composition and isotopic fingerprints. This combined approach allowed for an improved understanding of the occurrence and distribution of methane in shallow aquifers than using carbon isotope fingerprinting and dryness parameters alone.

Low $\delta^{13}\text{C}$ values of methane combined with a high dryness parameter and methanogenic conditions indicated by aqueous geochemistry provided clear evidence for in-situ biogenic production of methane in 39% of the investigated samples (CH₄-type A, Fig. 13).

High dryness and biogenic C isotope signatures coexisting with elevated sulfate, nitrate and/or oxygen concentrations, and isotopic compositions of nitrate and sulfate indicating ongoing sulfate reduction and denitrification, point to a second type of biogenic gases having formed in an anoxic milieu before migrating into more oxidizing aquifer sections (CH₄-type B, 3.7% Fig. 13).

Samples with apparent thermogenic gases based on $\delta^{13}\text{C}$ values were often characterized by no detectable higher alkanes. This suggests that these samples contained biogenic gases that had been partly oxidized, which leads to a shift towards elevated “pseudo-thermogenic” $\delta^{13}\text{C}$ values (CH₄-type D, 14.1 % Table 3). It is of key importance to identify the occurrence of “pseudo-thermogenic” gas signatures during monitoring of potential environmental impacts from unconventional [energy](#) resource development to prevent false conclusions [such as the migration of deeply-sourced thermogenic gases into shallow aquifers](#).

For 28.1 % of the samples, ethane and sometimes propane coexisted with biogenic methane (low $\delta^{13}\text{C}$), [which is](#) typical for in-situ gases produced in coal seams in which the groundwater wells are completed (CH₄-type A*, Table 3, Fig. 13).

Migration of mixed gas, composed predominantly of biogenic methane with traces of propane, [into non-CBM aquifers](#) was detected for 11.1% of the samples (CH₄-type C, Fig. 13).

A large majority of gases (67.4 %) obtained from the GOWN network were found to be in-situ gases either derived from in-situ formation in coalbeds (28.1 %) or produced microbially within aquifers with methanogenic conditions (39.3%). We conclude that combining hydrochemistry, in particular redox-sensitive species and their isotope ratios, with gas concentration ratios and carbon isotope signatures of alkanes and CO₂ constitutes an excellent approach to accurately assess methane formation and migration revealing additional insights compared to approaches based on gas composition and isotope ratios only.

Author contribution

P. Humez and B. Mayer prepared the manuscript and implemented the geochemical and multi-isotopic approaches to determine and quantify the origin and fate of methane in the groundwater systems. M. Nightingale, A. Kingston, S. Taylor and G. Bayegnak analyzed, compiled and delivered the geochemical and isotopic databases. V. Becker, M. Nightingale, A. Kingston, S. Taylor contributed significantly to the interpretation of the geochemical and isotopic data. G. Bayegnak

(AEMERA) provided expertise on the regional monitoring program while R. Millot and W. Kloppmann (BRGM) provided expertise with evaluation of geochemical processes.

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10 | Baseline" project [entitled "environmental baseline conditions for impact assessment of unconventional gas exploitation: advancing geochemical tracer and monitoring techniques"](#). [We also thank the reviewers for their constructive comments and suggestions.](#)

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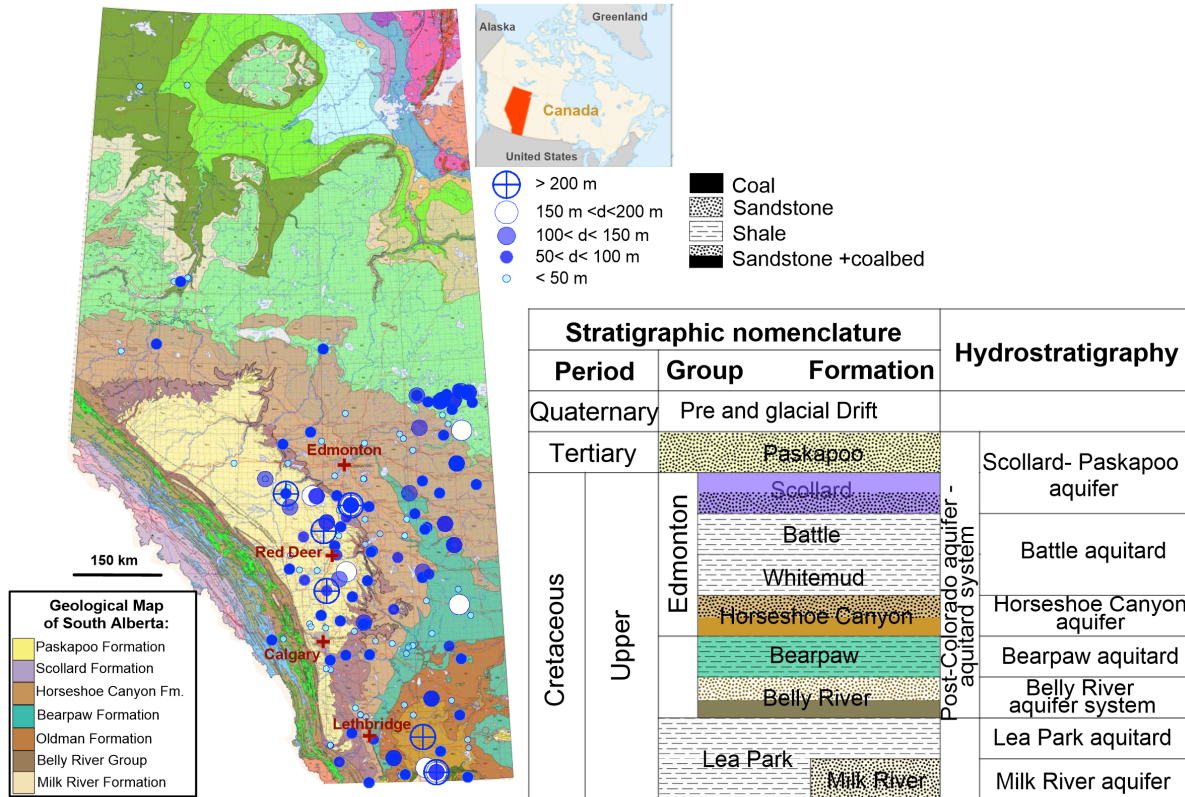


Figure 1: Location and depths of 186 water wells from the GOWN monitoring program used in this study shown on a geological bedrock map of Alberta (Alberta Geological Survey). Also shown is a stratigraphic column for Southern Alberta (modified from Bachu, 1999).

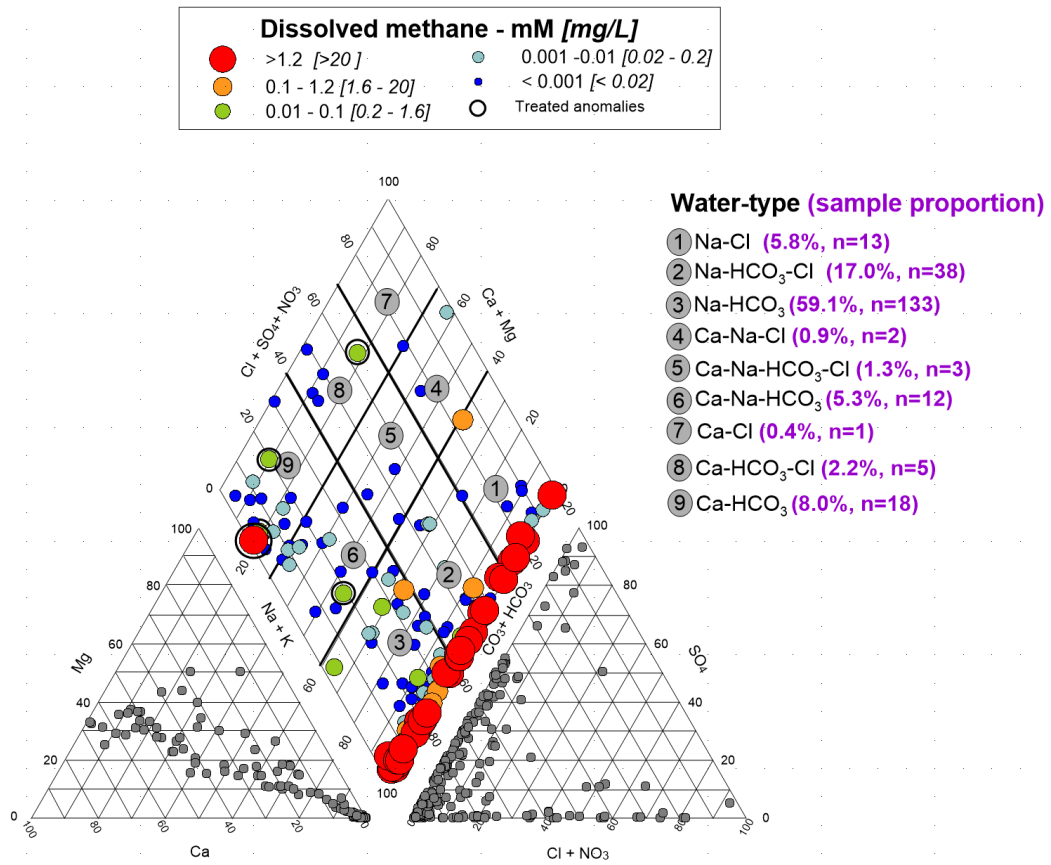


Figure 2: Piper diagram and water type classification of groundwater samples from the GOWN network (n = 221).

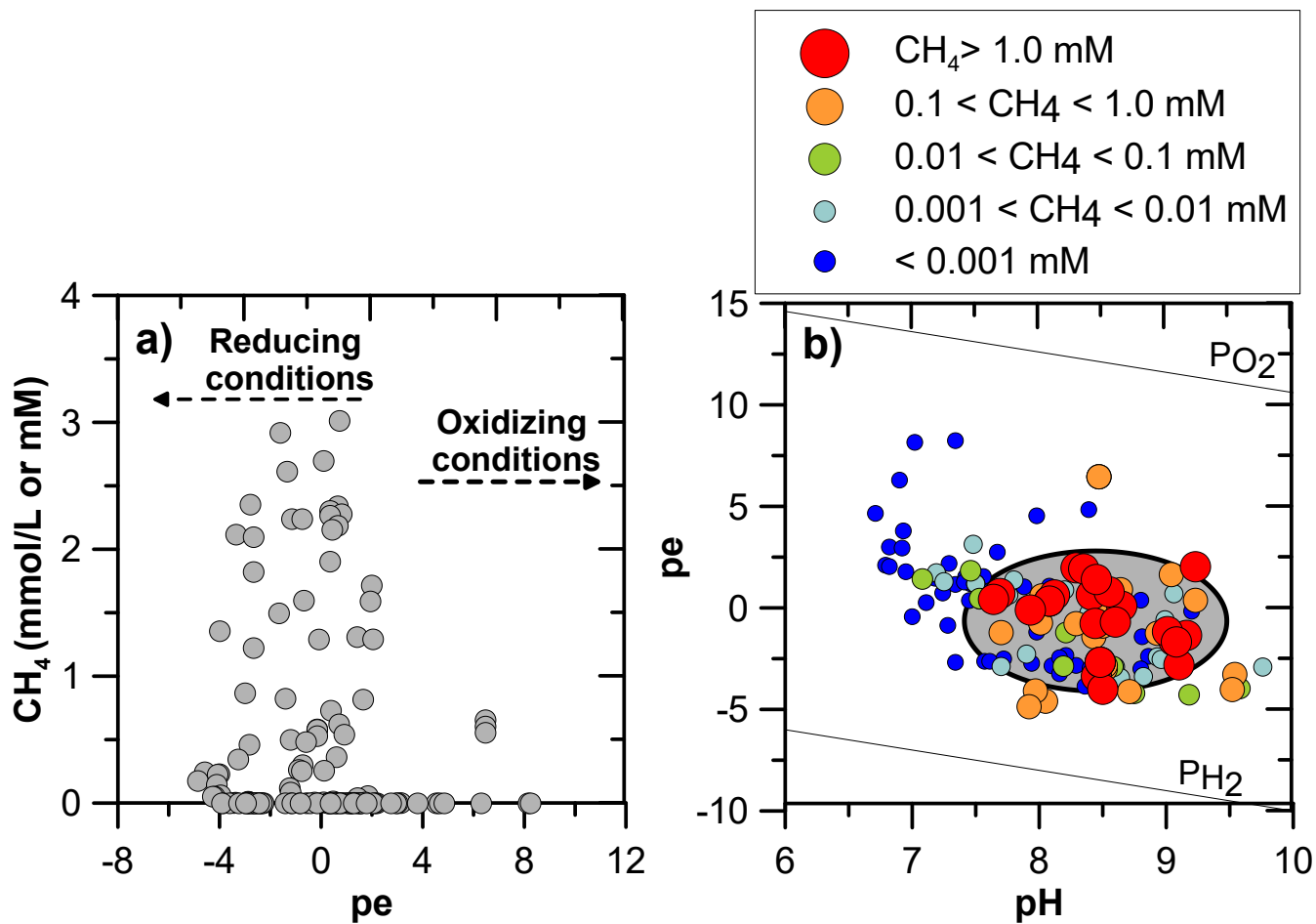


Figure 3: a) Methane concentrations in water samples versus pe b) pe versus pH.

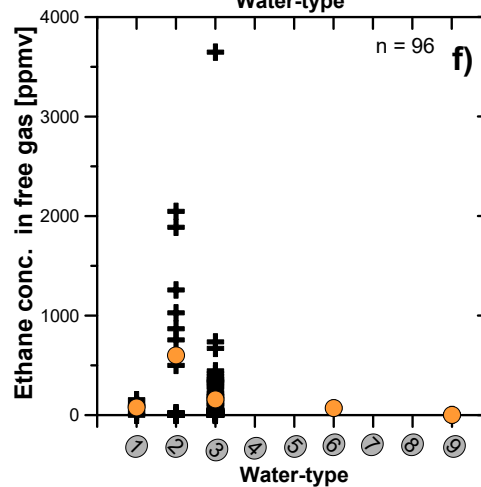
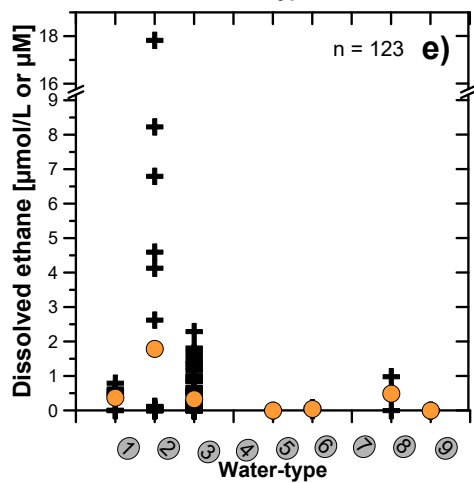
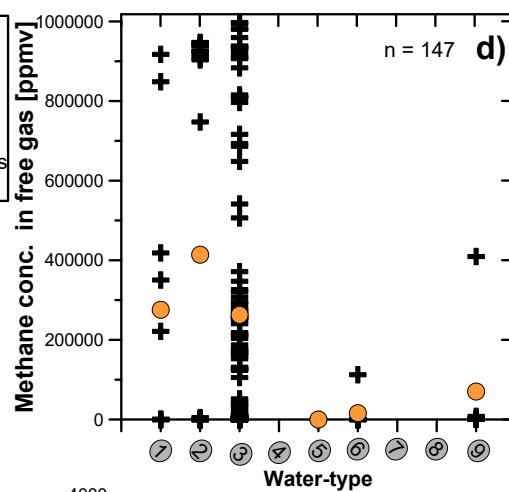
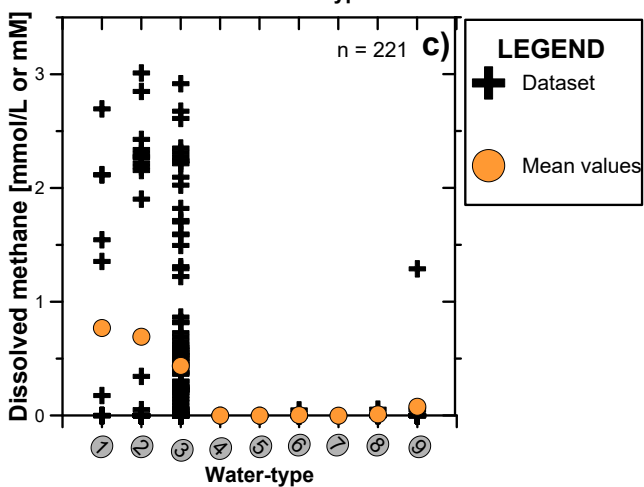
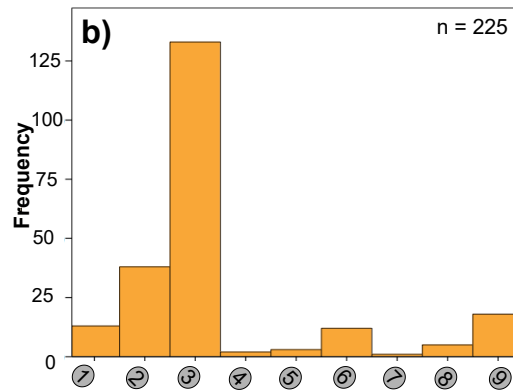
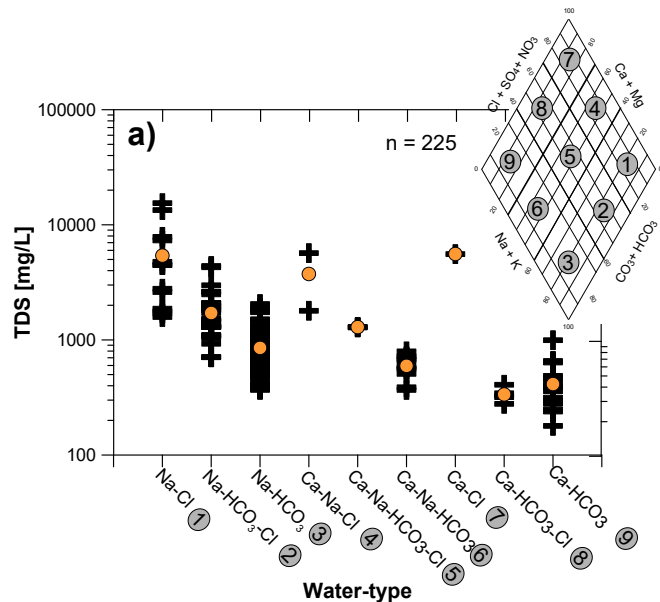


Figure 4: a) Distribution of TDS b) frequency histogram of methane-containing samples, methane and ethane in both dissolved (c, e) and free gas (d, f) phases versus water type classification defined in the Piper diagram.

LEGEND

- $\text{CH}_4 > 1.0 \text{ mM}$
- $0.001 < \text{CH}_4 < 0.01 \text{ mM}$
- $0.1 < \text{CH}_4 < 1.0 \text{ mM}$
- $\text{CH}_4 < 0.001 \text{ mM}$
- $0.01 < \text{CH}_4 < 0.1 \text{ mM}$

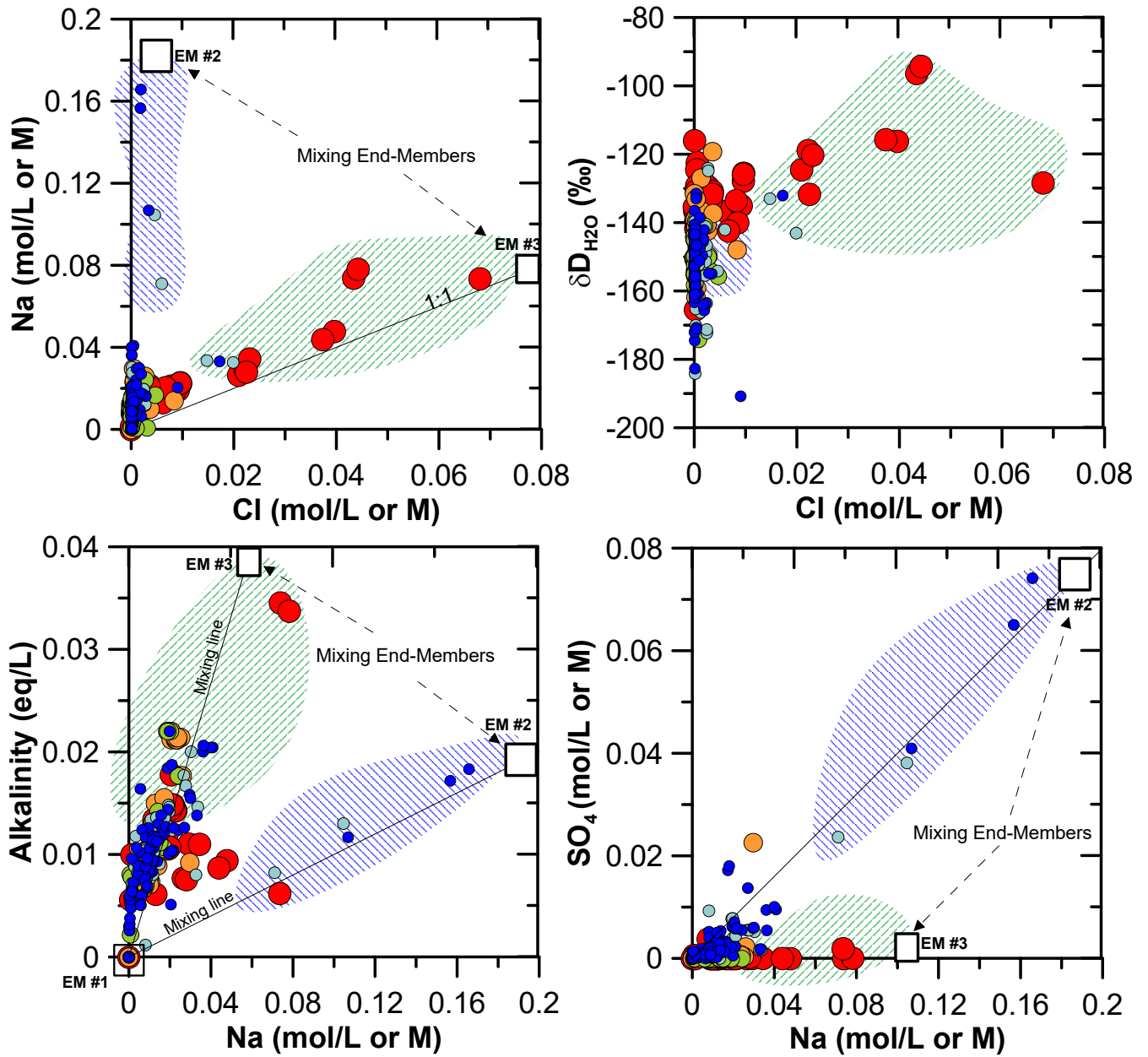
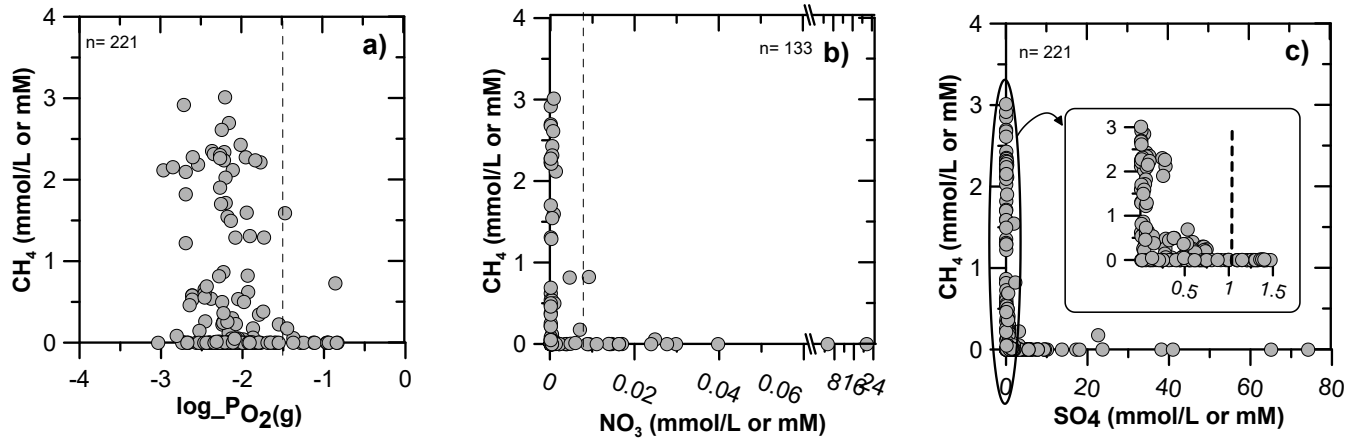


Figure 5: Identification of groundwater geochemistry end-members (shaded colors) and mixing trends indicated with solid lines. Dissolved methane concentrations are indicated by symbol size and color (EM = potential end-member compositions).



5 Figure 6: Redox diagram and binary relationship between methane/nitrate and sulfate concentration in the GOWN water samples.

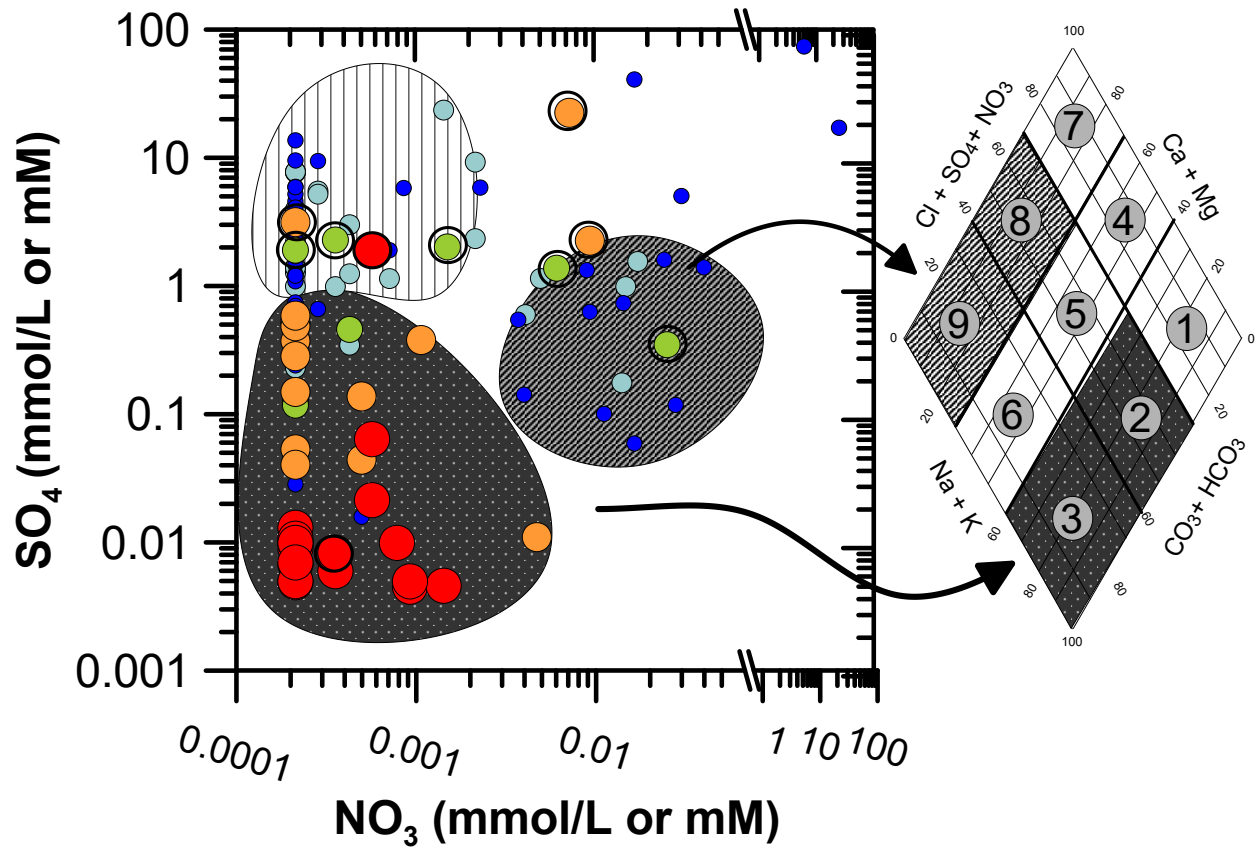
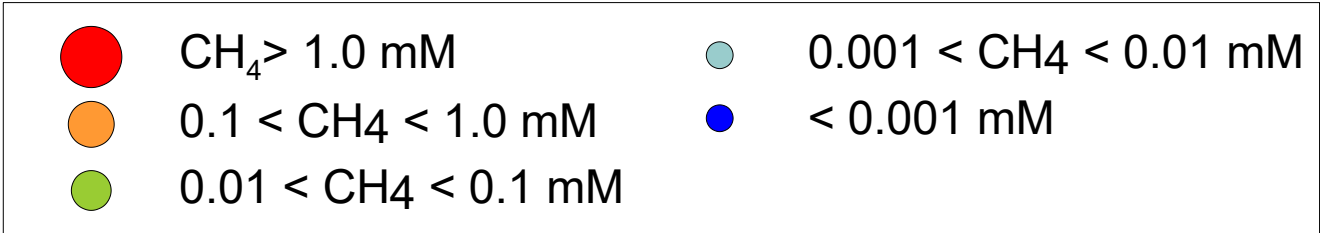


Figure 7: Sulfate versus nitrate concentrations in water samples and methane distribution (n=133) indicated by size and color of symbols. Shaded areas relate to water types.

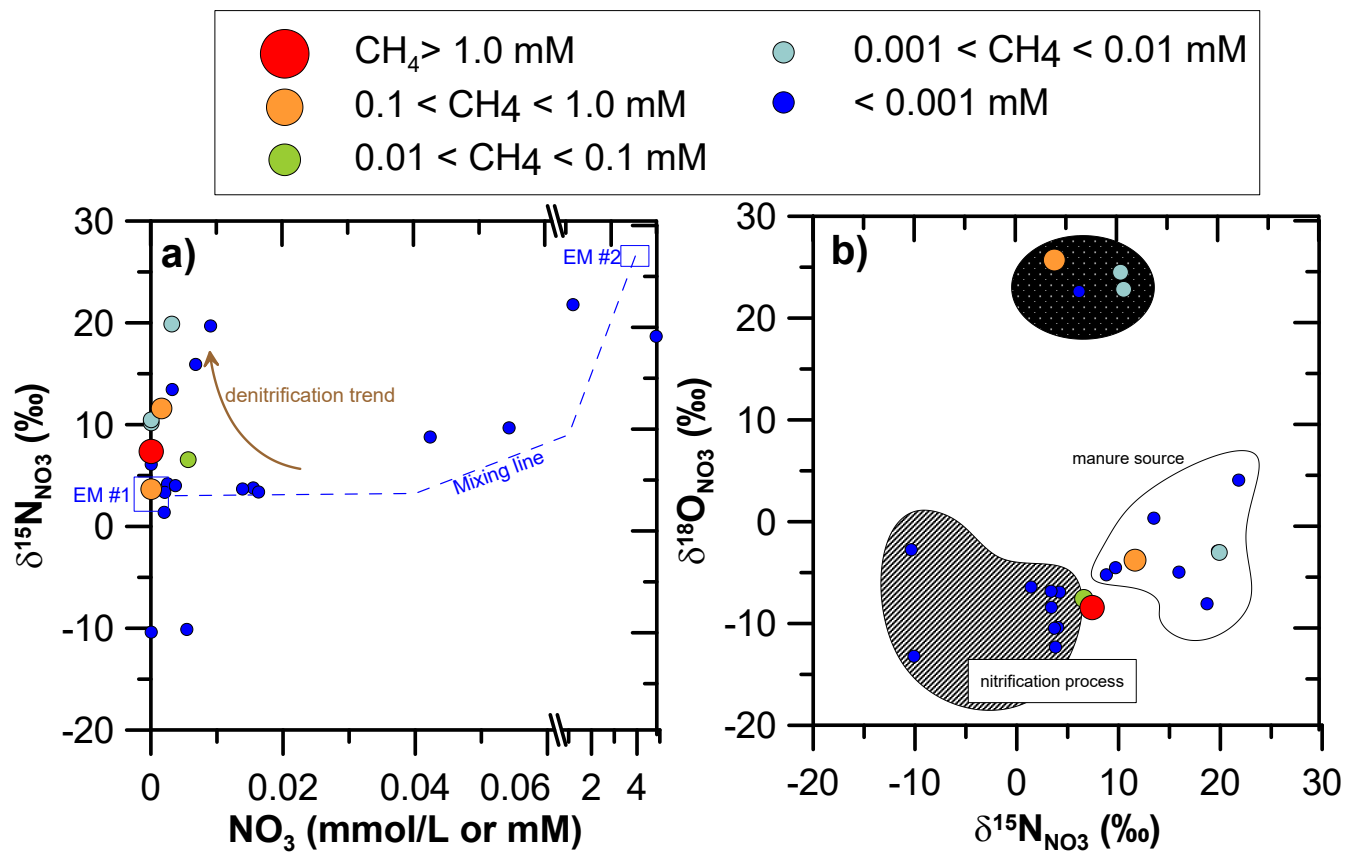


Figure 8: Cross plot of $\delta^{15}N_{NO_3}$ and $\delta^{18}O_{NO_3}$ values with N- NO_3 concentrations (mM) and identification of three groups of data according to nitrate origin and formation processes. Dissolved methane concentration is reported as colored symbols.

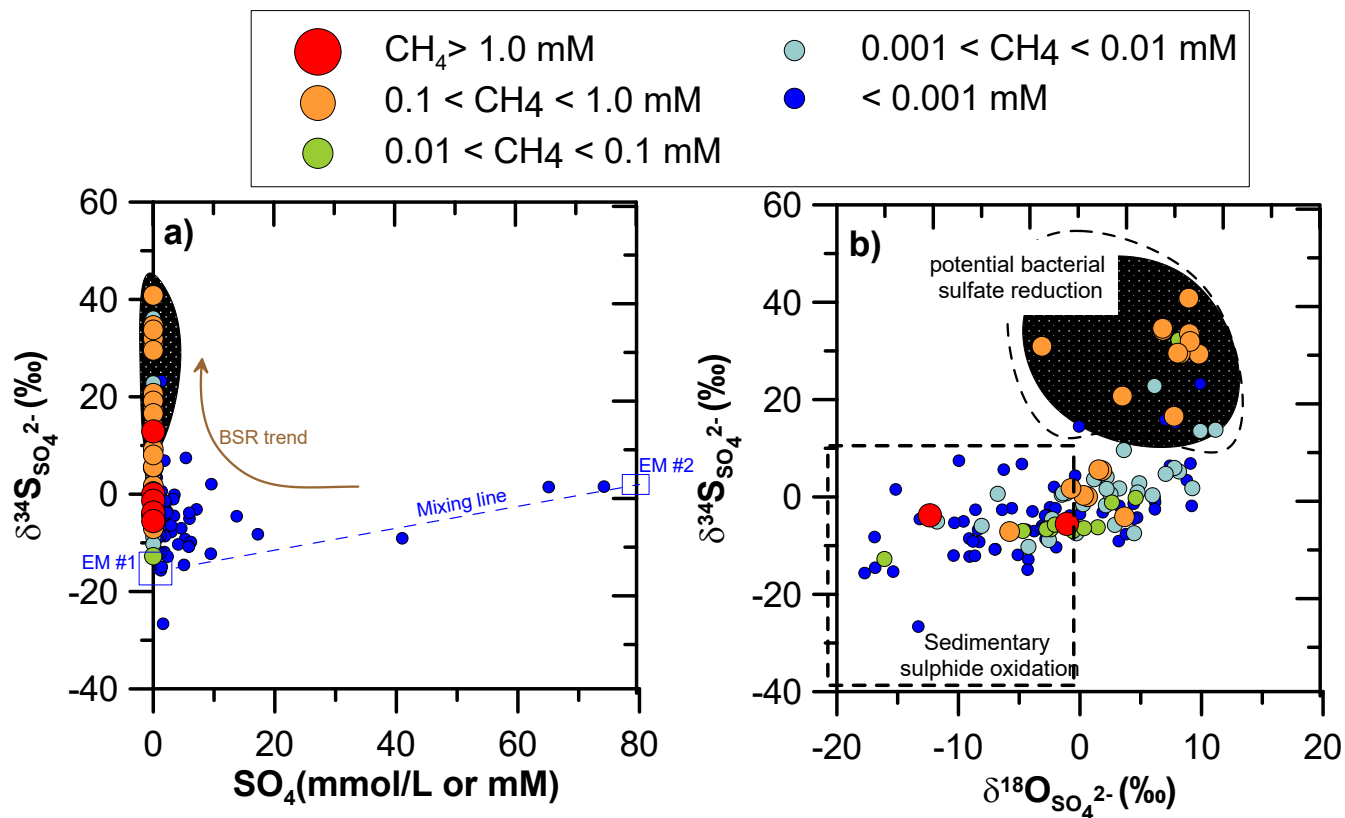


Figure 9: Cross plot of $\delta^{34}\text{S}$ and $\delta^{18}\text{O}$ values and sulfate concentrations (mM) and identification of groups of data according to sulfate origin and formation processes. Dissolved methane concentrations are reported as colored symbols.

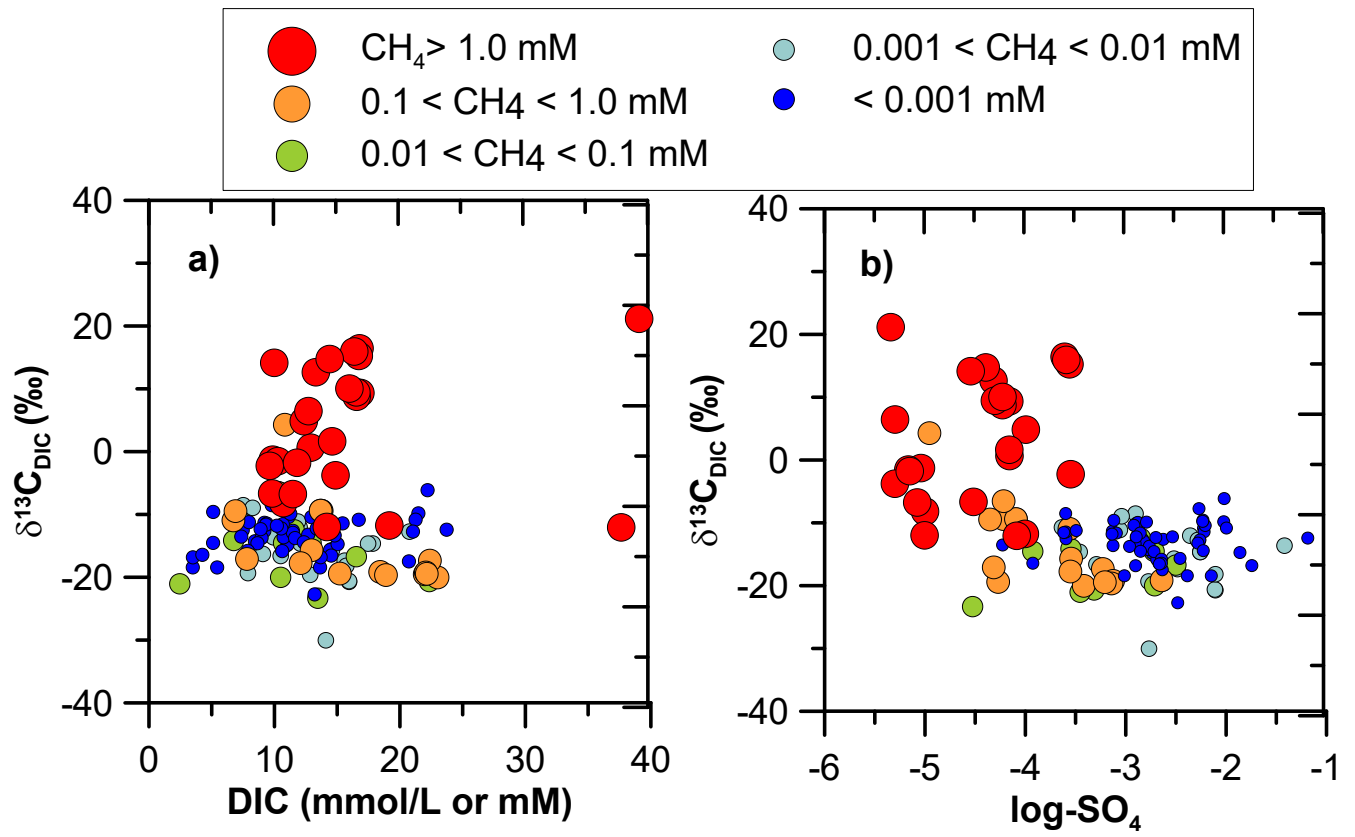


Figure 10: $\delta^{13}\text{C}$ values of DIC and methane concentrations versus a) DIC concentrations and b) sulfate concentrations (\log -concentration).

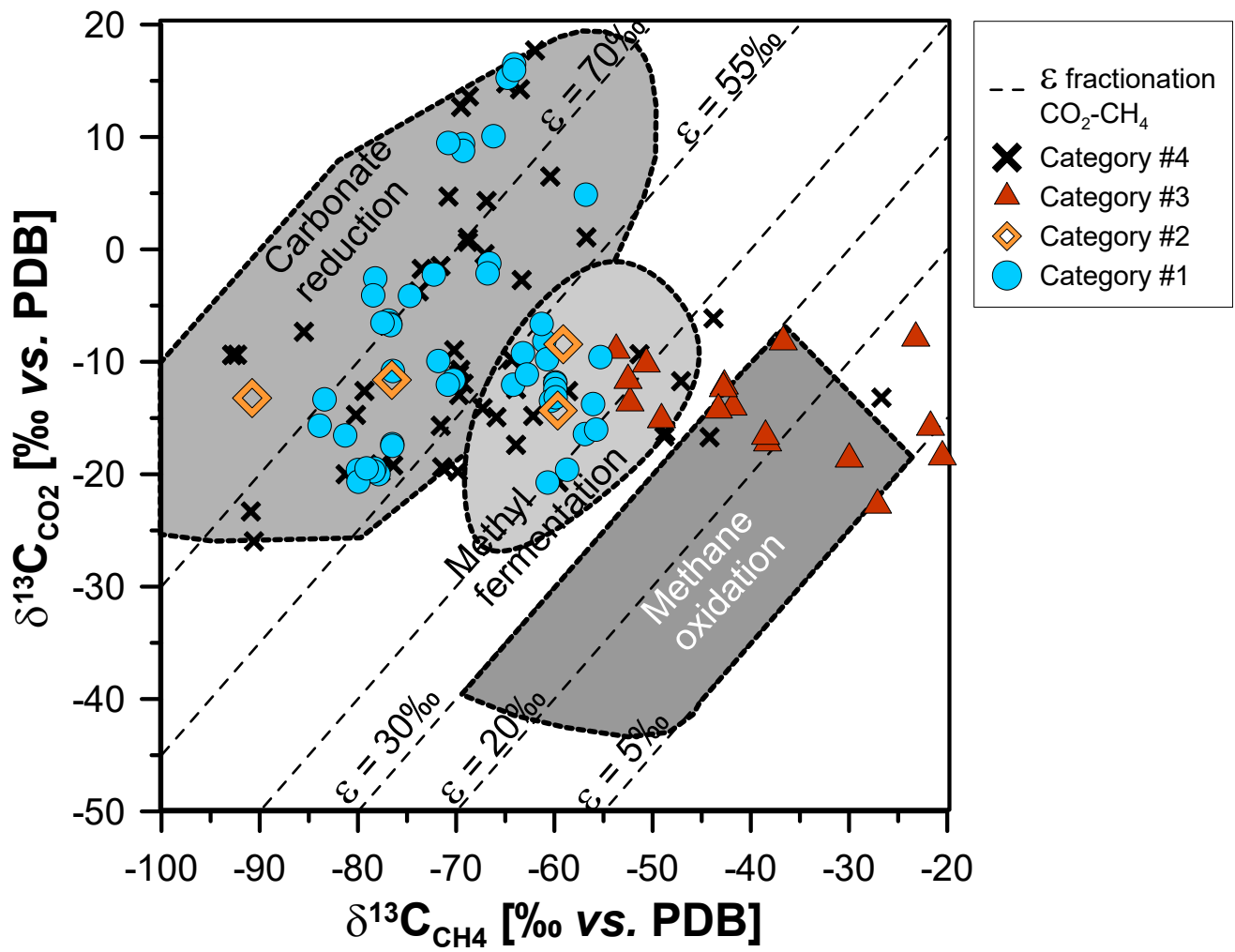


Figure 11: $\delta^{13}\text{C}$ values of methane and CO_2 revealing methane formation and consumption pathways, confirming that category #3 samples plot near the methane oxidation field adapted from Whiticar (1999).

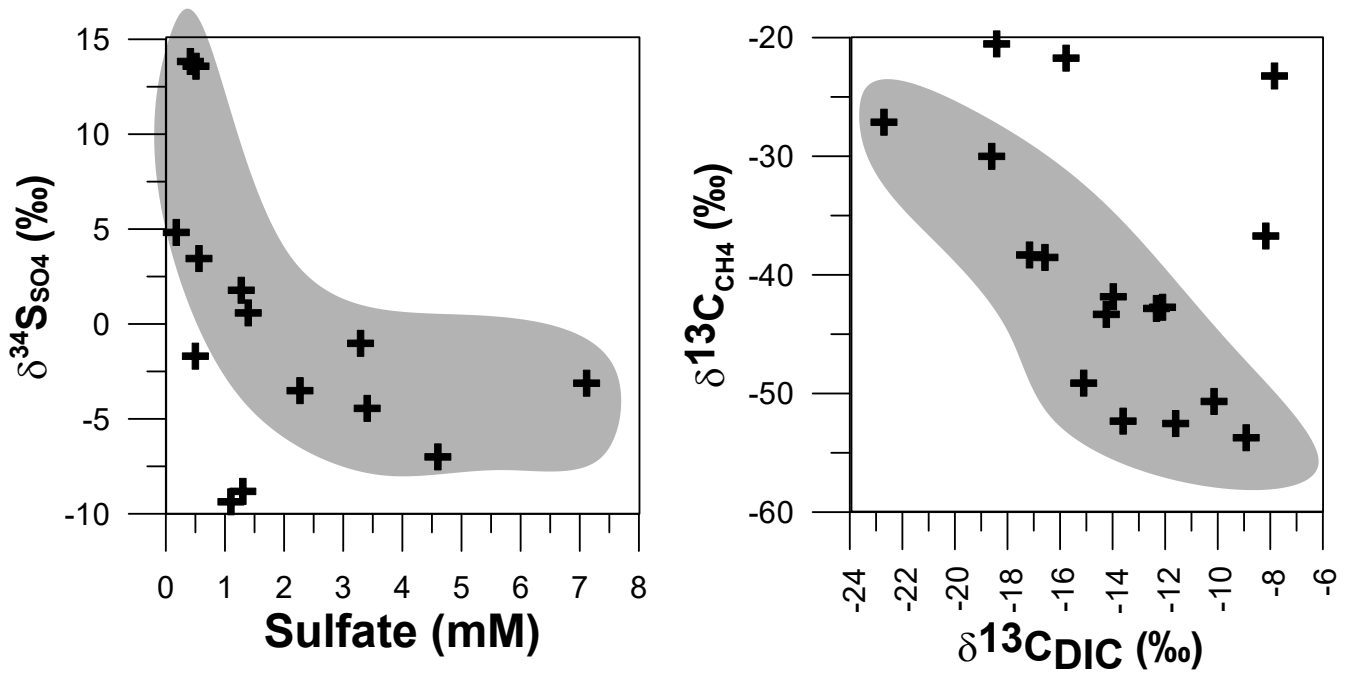
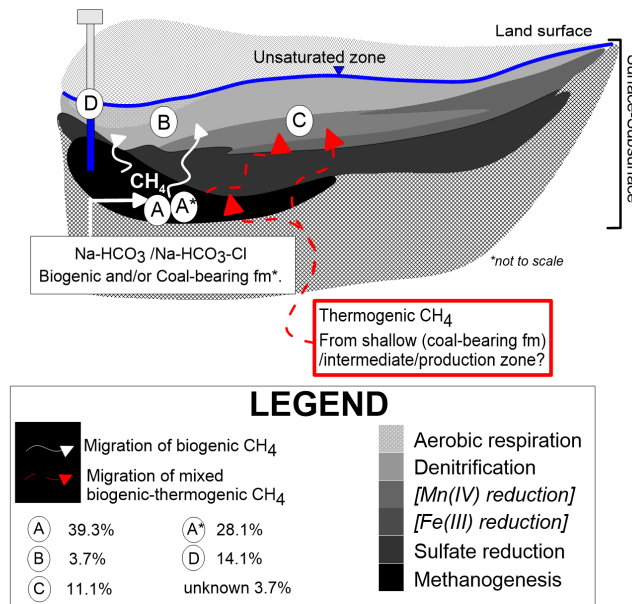


Figure 12: $\delta^{34}\text{S}$ versus sulfate concentrations and $\delta^{13}\text{C}$ of methane versus DIC for samples in category #3 supporting that methane oxidation coupled with bacterial sulfate reduction is responsible for elevated $\delta^{13}\text{C}$ values of methane.



5 Figure 13: Geochemical and multi-isotopic approach applied in this study to classify the methane type/occurrence considering the redox zoning constraint.

Table 1: Summary table of samples in the investigated data set (IB = ionic balance).

	Total	Criteria #1 (C#1) = -10% <IB < +10%	C#2 = C#1+ gas info	C#3= C#1 + C#2 + chemistry
Free gas conc. only	8	4	4	4
Dissolved gas conc. only	134	80	80	78
Free + dissolved gas conc.	187	150	150	143
No analysis available	43* 50**	8* 9**	/	/
Isotope ratios free gas only	141	105	105	100
Isotope ratios dissolved gas only	3	2	2	2
Isotope ratios free + dissolved gas	36	34	34	33
No isotope ratio information	192	101	93	90
Sum	372	242	234	225
* for gas analyses ** for chemistry analyses		Retained subsets for discussion		

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Table 2: Descriptive statistics for concentrations of major and minor species and for the isotopic composition of methane, nitrate, sulfate and DIC. Detection limits are also shown (DL).

	Unit	N	Min	Max	Range	Median	Mean	Stdev	DL
CH_{4,aq}	mM	221	0.00 [2.18e-6]	3.01	3.01	0.00 [2.77e-3]	0.43	0.79	6.25e-7
CH_{4,g}	ppmv	147	0.29	9.98e5	9.98e5	3.83e4	2.65e5	3.55e5	0.05
C₂H_{6,aq}	μM	123	0.00 [3.34e-4]	17.83	17.83	0.06	0.60	1.95	3.33e-4
C₂H_{6,g}	ppmv	96	0.08	3650	3650	38	215	499	0.05
C₃H_{8,aq}	μM	32	0.00 [3.18e-4]	0.90	0.90	0.00 [1.57e-3]	0.03	0.16	2.27e-4
C₃H_{8,g}	ppmv	36	0.05	4.60	4.55	0.26	0.67	0.91	0.05
O₂	mM	199	0.01	0.25	0.25	0.01	0.02	0.04	1.56e-3
DIC	mM	225	0.81	39.07	38.25	12.29	12.94	5.59	8.20e-3
Ca	mM	136	0.01	9.08	9.06	0.40	0.94	1.26	7.49e-3
NO₃	μM	136	0.21	2.13e4	2.13e4	0.21	203	1.87e3	0.03
K	mM	225	0.01	0.54	0.53	0.05	0.07	0.07	5.12e-3
Mg	mM	225	0.00 [2.26e-3]	12.45	12.44	0.05	0.60	1.47	4.12e-6
Na	mM	225	0.03	165.78	165.75	12.76	17.28	20.03	1.30e-2
Cl	mM	225	0.01	68.01	68.00	0.40	2.81	7.82	8.46e-3

SO ₄	mM	225	0.00 [4.55e-3]	74.16	74.15	0.72	2.83	8.00	2.44e-4
$\delta^{13}\text{C}_{\text{CH4-FG}}$	‰	133	-92.8	-20.5	72.3	-66.2	-64.6	14.9	f(CH ₄)
$\delta^2\text{H}_{\text{CH4-FG}}$	‰	58	-437.1	-80.9	356.1	-291.5	-280.8	54.4	f(CH ₄)
$\delta^{13}\text{C}_{\text{CH4-DG}}$	‰	35	-85.5	-35.8	49.7	-65.6	-65.0	10.5	f(CH ₄)
$\delta^{15}\text{N}_{\text{NO3}}$	‰	24	-10.4	21.8	32.2	7.0	7.8	8.2	f(NO ₃)
$\delta^{18}\text{O}_{\text{NO3}}$	‰	24	-13.2	25.7	38.9	-5.1	-1.1	12.0	f(NO ₃)
$\delta^{34}\text{S}_{\text{SO4}}$	‰	158	-26.6	40.9	67.5	-1.6	1.8	12.6	f(SO ₄)
$\delta^{18}\text{O}_{\text{SO4}}$	‰	138	-17.7	11.2	28.9	-0.6	-0.6	6.7	f(SO ₄)
$\delta^{13}\text{C}_{\text{DIC}}$	‰	221	-30.8	21.2	52.0	-12.3	-10.8	8.8	f(DIC)
Range = maximum – minimum; f(X) : function of X concentration									

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10 Table 3: Categories classification, boundaries and methane-type (grey shading indicate anomalies, italic = boundaries, N.D non-detected, D.L detection limit)

Category		#1	#2	#3	#4			#5
Sub-cat					4.1	4.2	4.3	
N		53	5	17	2	38	15	5
$\delta^{13}\text{C}_{\text{CH4}}$ (‰)	<-55	X	X			X	X	X
	>-55	X		X	X	X (n=5)		X
Methane (mM)	< 0.01	X		X (max. 5 μM)	X	X	X	X
	> 0.01	X	X					
Ethane (μM)	< 0.002	X	X	N.D	X	X		
	> 0.002					X	X	X
Propane	<i>N.D</i>	X	X	X		X	X	X
	>D.L				X	X	X	
Dryness		>1000	>1000	n.d	>1000	<500	<500	
SO ₄ (mM)	<1	X						
	>1	X	X	X	X	X	X	X
NO ₃ (mM)	<0.006	X						
	>0.006	X	X		X	X	X	X

Redox ladder		Yes	No	Yes	Yes	Yes	Yes	Yes
Geol. formation		various	non-CBM	various	non-CBM	CBM	non-CBM	
CH₄-type		A	B	D	D	A*	C	E
A: In-situ biogenic CH ₄ (39.3%) A*: In-situ CH ₄ from CBM (28.1 %)		B: migration of biogenic CH ₄ into more oxidizing condition (3.7%)		C: Mixed gas origin (11.1%) D: CH ₄ oxidation/ Post-sampling degradation of low-CH ₄ samples (14.1%)		E: Unknown (3.7%)		