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# High temperature generation and equilibration of methane in terrestrial geothermal systems: evidence from clumped isotopologues

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## Abstract

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Fluids emanating from geothermal areas contain trace quantities of methane and other simple hydrocarbons. These hydrocarbons are thought to derive from thermal cracking of organic matter dissolved in circulating meteoric or seawater or found in pre-existing organic-rich sedimentary rocks, but an abiotic origin has also been proposed. We measured the relative abundances of four CH<sub>4</sub> isotopologues (<sup>12</sup>CH<sub>4</sub>, <sup>13</sup>CH<sub>4</sub>, <sup>12</sup>CH<sub>3</sub>D, and <sup>13</sup>CH<sub>3</sub>D) in hydrothermal gases discharged by steam vents and geothermal wells from Iceland and Nisyros island (Greece) in order to investigate the origin of methane. Measured methane samples yielded consistently low  $\Delta^{13}$ CH<sub>3</sub>D values ( $^{13}$ CH<sub>3</sub>D abundance relative to stochastic) of 0.82 to 1.77‰, which correspond to high apparent temperatures of isotopologue equilibrium ( $T_{\Delta^{13}CH_2D}$  = 278–490°C). Hydrothermal well fluids from the Krafla and Námafjall geothermal fields in Iceland yielded the lowest Δ<sup>13</sup>CH<sub>3</sub>D values, and thus the highest  $\Delta^{13}\text{CH}_3\text{D-based}$  temperatures averaging  $438^{+55}_{-45}~^{\circ}\text{C}.$  Those samples also show the most pronounced departures in  $\delta D_{CH4}$  and  $\delta^{13}C_{CH4}$  values expected for isotopic equilibrium with respect to  $\delta D_{H2O}$  and  $\delta^{13}C_{CO2}$ . In contrast, CH<sub>4</sub> samples from natural steam vents in other Iceland locations and in Nisyros have slightly higher  $\Delta^{13}CH_3D$  values (with  $T_{\Delta^{13}CH_2D}=$  $351^{+42}_{-35}$  °C) and have  $\delta D_{CH4}$  and  $\delta^{13}C_{CH4}$  values that are consistent with those expected for isotopic equilibrium with both H<sub>2</sub>O and CO<sub>2</sub>. The short fluid residence times (1-50 years) in systems that are exploited for geothermal energy, such as Krafla and Námafjall, combined with the proximity of a hot magma chamber, favor the preservation of kinetic signals. The initial disequilibrium  $\delta D_{CH4}$ and  $\delta^{13}C_{CH4}$  values are consistent with a thermogenic origin from immature organics dissolved in hydrothermally heated groundwater, but an abiotic origin cannot be excluded. The high apparent  $\Delta^{13}$ CH<sub>3</sub>D-based temperatures at Krafla and Námafjall could therefore represent nonequilibrium signals associated with either pyrolysis or abiotic generation of CH<sub>4</sub> in a superheated vapor or supercritical water phase (>374°C), considered to exist in the roots of the system above the magmatic heat source. Isotopologue equilibration calculations demonstrate that under such conditions (e.g. >400°C) kinetic signals would be erased in days to months, implying rapid migration and quenching of CH<sub>4</sub> into the overlying subcritical (<300°C) hydrothermal reservoir fluids. In systems with longer fluid residence times such as Nisyros, equilibrium isotopologue distributions at temperatures of ~350°C are consistent with long fluid residence times on the order of >100 years. Our calculations further reveal that CO<sub>2</sub>–CH<sub>4</sub> isotopic equilibration requires unreasonably long fluid residence times, suggesting that any apparent <sup>13</sup>C equilibrium may be coincidental.

Keywords: Methane; Isotopologues; Geothermal systems; Hydrogen isotope equilibration; Superheated vapor

## 1. Introduction

 Most methane at the Earth's surface is ultimately derived from biological activity, either through microbial methanogenesis or by the thermal degradation of organic matter. A small fraction of Earth's CH<sub>4</sub> may be produced abiotically, including that formed in hydrothermal systems (Sherwood Lollar et al., 1993; McCollom and Seewald, 2006; Proskurowski et al., 2008; Klein et al., 2019; Reeves and Fiebig, 2020). Since methane is the simplest hydrocarbon (CH<sub>4</sub>) and is often ascribed with an important role in the development of early life (e.g. Kasting et al., 2001; Ueno et al., 2006), it is of interest to characterize its various formation pathways, which can ultimately help in understanding its source on other planetary bodies (Etiope et al., 2011).

The commonly proposed pathway for abiotic production of methane involves the reduction of  $CO_2$  by  $H_2$ , often referred to as the Sabatier reaction (e.g. Giggenbach, 1997, 1987; Klein et al., 2019; McCollom and Seewald, 2006) :

$$CO_2 + 4H_2 \rightleftharpoons CH_4 + 2H_2O$$
 (1)

For example, high concentrations of  $CH_4$  found in many serpentinization environments have been attributed to this reaction, where the reducing conditions (high  $fH_2$ ) imposed by serpentinization reactions provide a thermodynamic drive for  $CH_4$  formation, with mantle  $CO_2$  or dissolved carbonate of seawater or meteoric water origin as the source of inorganic carbon (Shock, 1992; Klein et al., 2019). Hydrolysis-induced oxidation of ferrous iron drives the evolution of serpentinization systems towards high  $fH_2$  after prolonged water-rock interaction, and is favored in host rocks with high Fe (II) contents and increasing fluid pH (Leong and Shock, 2020).

Geothermal systems are also characterized by dynamic multiphase reactions and metastable equilibria with evolving fluid pH and precipitation of secondary minerals, which have important controls on redox conditions and  $fH_2$  (Giggenbach, 1987; Arnórsson et al., 2007; Stefánsson, 2017). However, the generally lower Fe (II) content of rocks in geothermal vs. serpentinization systems is less favorable for  $H_2$  generation and thus  $CH_4$  production. Furthermore, the reduction of  $CO_2$  to  $CH_4$  and other hydrocarbons has a high activation energy and is shown to be sluggish under single-liquid phase hydrothermal conditions (McCollom, 2016; Seewald et al., 2006). Nevertheless, based on bulk stable isotope compositions of geothermal  $CH_4$ , in particular on elevated  $\delta^{13}C$  values compared to conventional thermogenic sources, it has been proposed that methane found in geothermal environments is of abiotic origin (Fiebig et al., 2007; Etiope and Sherwood Lollar, 2013). However, observed  $\delta^{13}C_{CH4}$  and  $\delta D_{CH4}$  patterns also follow trends of thermogenic  $CH_4$  formation, such that most (if not all) geothermal methane could derive from the pyrolysis of organic matter brought to depth by circulation of surface derived waters (Des Marais et al., 1981; Fiebig et al., 2019).

This poses the question as to what conditions are most favorable for abiotic methane production. On the one hand, reaction (1) is exothermic, with a  $\Delta H$  of reaction ranging from -250

to -210 kJ/mol between  $100^{\circ}$ C and  $500^{\circ}$ C (based on SUPCRT92; Johnson et al., 1992), such that production of methane is favored at lower (<300°C) temperatures but kinetics become increasingly sluggish. On the other hand, higher temperatures promote faster rates of reaction (and hence of CH<sub>4</sub> production), albeit with less thermodynamic drive. In addition to the temperature control, experimental studies suggest that the presence of a vapor phase, as opposed to an entirely aqueous phase, promotes methane production from  $CO_2$  reduction (McCollom, 2016, 2013 and references therein). The faster rate of  $CO_2$  reduction in a gas phase supports the hypothesis that methane in seafloor hydrothermal vents could have been generated abiotically in fluid inclusions within the oceanic lithosphere, and later extracted by circulating hydrothermal fluids (Kelley, 1996; McDermott et al., 2015; Wang et al., 2018; Klein et al., 2019; Labidi et al., 2020). In geothermal systems, the combination of high temperatures with processes that promote the formation of a vapor phase could overcome the kinetic inhibitions associated with abiotic methane.

Chemical and isotope geothermometers can give valuable insight into the thermal evolution of solutes, gases and isotopes in hydrothermal fluids. Chemical geothermometers are based on the fundamental assumption of chemical equilibrium between various gaseous and aqueous species and/or hydrothermal minerals at a particular temperature (e.g., Arnórsson et al., 1983a; Chiodini and Marini, 1998; D'Amore and Panichi, 1980; Giggenbach, 1988; Gunnarsson and Arnórsson, 2000). Isotope thermometers, which rely on the isotopic fractionation between gaseous and/or aqueous species, can preserve more intricate signals of subsurface temperatures and processes, although they can also be affected by kinetic effects (Richet et al., 1977; Giggenbach, 1982; Ohmoto and Lasaga, 1982; Horibe and Craig, 1995; Giggenbach, 1997; Horita, 2001; Fiebig et al., 2004). Other processes may control or affect these apparent equilibria, for example sources of elements (and thus initial isotopic compositions), sluggish reaction kinetics, and secondary processes in the fluid conduit from reservoir to surface like boiling and mineral formation (Arnórsson et al., 2007). Substantial discrepancies are commonly obtained between temperatures calculated from different geothermometers. This may reflect either different closure temperatures of a particular chemical or isotope reaction, different initial degrees of disequilibrium controlled by reaction kinetics, and/or different responses to secondary processes and re-equilibration (Giggenbach, 1987; Chiodini and Marini, 1998; Arnórsson et al., 2007; Stefánsson, 2017).

Geothermometers based on the relative concentrations and  $^{13}$ C fractionation between CH<sub>4</sub> and CO<sub>2</sub> converge to similar temperatures (often ~300-350°C) for several geothermal areas, including those found in Yellowstone, New Zealand and Nisyros (Craig, 1953; Hulston and McCabe, 1962; Giggenbach, 1982; Giggenbach, 1997; Fiebig et al., 2004), which has led to the suggestion that abiotic CH<sub>4</sub> is produced through reaction (1). However, since the  $\delta$ D and  $\delta$ <sup>13</sup>C systematics of hydrothermal CH<sub>4</sub> and higher n-alkanes in several locations can also be explained by open-system thermal cracking of marine or terrestrial organic matter (Fiebig et al., 2019), observed equilibrium  $^{13}$ C fractionations in some systems might be fortuitous. This is explained by the maturation trend of organic matter, where thermal cracking of mature organic matter produces CH<sub>4</sub> with higher  $\delta$ <sup>13</sup>C, in apparent isotopic equilibrium with mantle CO<sub>2</sub>. Moreover, evidence for the natural occurrence of reaction (1) in geothermal systems, in particular with regards to the existence of an efficient catalyst under hydrous conditions, is still lacking. It follows that chemical equilibrium between CO<sub>2</sub>-CH<sub>4</sub>-H<sub>2</sub> may not be attained as often as has been

previously suggested (e.g. Giggenbach, 1997), owing to the complex reaction pathways and intermediates involved in the abiotic reduction of  $CO_2$  (Seewald et al., 2006), written in its simplified version as reaction (1).

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Measurements of the relative abundances of methane isotopologues (12CH<sub>4</sub>, 13CH<sub>4</sub>, <sup>12</sup>CH<sub>3</sub>D, <sup>13</sup>CH<sub>3</sub>D and <sup>12</sup>CH<sub>2</sub>D<sub>2</sub>) can be used to calculate the temperatures that correspond to internal isotopologue equilibria (Ono et al., 2014; Stolper et al., 2014a; Wang et al., 2015; Stolper et al., 2017; Young et al., 2017). The metrics  $\Delta^{13}CH_3D$  and  $\Delta^{12}CH_2D_2$  represent the excess abundance of each doubly-substituted isotopologue (13CH<sub>3</sub>D and 12CH<sub>2</sub>D<sub>2</sub>, respectively) relative to a stochastic distribution, and become smaller with increasing temperature. These clumped isotopologue fractionations can be compared to isotopic fractionations between CH<sub>3</sub>D vs. HDO and <sup>13</sup>CH<sub>4</sub> vs. <sup>13</sup>CO<sub>2</sub> to examine the degree of equilibrium or disequilibrium among these three independent isotope/isotopologue systems (Wang et al., 2018; Labidi et al., 2020). Previous applications of the clumped CH<sub>4</sub> isotopologue geothermometer reported apparent ∆<sup>13</sup>CH<sub>3</sub>Dbased temperatures in marine hydrothermal vents ranging from 160°C to 360°C (Wang et al., 2018; Labidi et al., 2020). Compared to seafloor vents, methane from terrestrial fumaroles measured to date shows higher apparent equilibrium temperatures, ranging from 350 to 580°C (Douglas et al., 2017). We note that the measurements of Douglas et al. (2017) were reported as  $\Delta_{18}$  values, which differ slightly from  $\Delta^{13}$ CH<sub>3</sub>D since it incorporates both methane isotopologues of mass 18 ( $^{13}CH_3D$  and  $^{12}CH_2D_2$ ). Accurate conversion between  $\Delta_{18}$  and  $\Delta^{13}CH_3D$  values requires the measurement of <sup>12</sup>CH<sub>2</sub>D<sub>2</sub>. Since <sup>13</sup>CH<sub>3</sub>D is approximately 50 times more abundant than  $^{12}\text{CH}_2\text{D}_2$ , however, the value of  $\Delta_{18}$  is expected to be close to  $\Delta^{13}\text{CH}_3\text{D}$  for natural methane samples. The use of these different proxies depends on the analytical procedure, since in early clumped CH<sub>4</sub> studies by mass spectrometry <sup>13</sup>CH<sub>3</sub>D and <sup>12</sup>CH<sub>2</sub>D<sub>2</sub> peaks were not well resolved (Stolper et al., 2014b). Spectroscopic measurements (as those performed here) are not affected by this overlap. The relatively low  $\Delta_{18}$  values measured for CH<sub>4</sub> from terrestrial fumaroles are of interest because their corresponding high apparent temperatures often exceed the critical temperature of pure water (T<sub>c</sub> = 374°C), which would imply generation or equilibration of CH<sub>4</sub> in a supercritical water or superheated vapor phase (above critical temperature but below critical pressure of water; Fig. 1). Such fluids have been suggested to exist at the contact between hydrothermal systems and magmatic heat sources (Hayba and Ingebritsen, 1997; Scott et al., 2015; Heřmanská et al., 2019). Alternatively, low  $\Delta^{13}CH_3D$  values could also be explained by kinetic effects during thermal cracking of organic matter (e.g. Shuai et al., 2018a), highlighting gaps in our understanding of the origins of geothermal CH<sub>4</sub>.

In this study, we measured the relative abundances of CH<sub>4</sub> isotopologues (<sup>12</sup>CH<sub>4</sub>, <sup>13</sup>CH<sub>4</sub>, <sup>12</sup>CH<sub>3</sub>D, and <sup>13</sup>CH<sub>3</sub>D) in steam vents and hydrothermal well fluids from Iceland and Nisyros island (Greece) in order to constrain the origin of methane in terrestrial geothermal settings. In addition, measurements of <sup>12</sup>CH<sub>2</sub>D<sub>2</sub> from two Nisyros samples were reported in Gonzalez et al. (2019). Our sample set covers a wide range of geothermal environments, characterized by a variable range of hydrothermal fluid reservoir temperatures, variable fluid residence times and ascent durations to surface. Some of the systems studied, for example Krafla, have also been observed to contain a high-temperature supercritical or superheated fluid reservoir (>400°C) below the conventional subcritical reservoir (~200-300°C). Our results imply that isotope and isotopologue equilibrium might be achieved for CH<sub>4</sub> in long-lived natural geothermal settings, but

significant disequilibria are observed for samples from higher-permeability systems where fluid circulation is accelerated by the operation of active geothermal wells. We discuss the relative rates of isotope/isotopologue exchange to relate isotopologue equilibration with fluid residence times. The measured disequilibrium signals can be explained by a thermogenic origin for methane, but our data does not rule out the possibility of abiotic synthesis.

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# 2. Geological context and sample description

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# 2.1 Iceland

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Iceland is located at the intersection of a hotspot and the mid-Atlantic ridge, giving rise to unusually high heat flow and crustal production rates (Gudmundsson, 2000), reflected by the existence of several active volcanoes. The island is host to over thirty high-temperature geothermal systems (e.g. Ármannsson, 2016), which are located along the active volcanic belts running in a SW-NE trend across the country (Fig. 2A). The development of these systems is thought to be related to the emplacement of intrusive bodies and formation of sub-vertical dykes and fractures leading to high permeability (Arnórsson, 1995), which promotes heating of groundwaters. Decades of geothermal exploration and development have generated a wealth of chemical and isotopic data on Icelandic hydrothermal fluids (Arnórsson et al., 1983a; Ármannsson et al., 1987; Gudmundsson, 2000; Stefánsson et al., 2017; Stefánsson, 2017), and geothermal wells have helped characterize basement geology and alteration assemblages (Ármannsson et al., 1987; Pope et al., 2016). The source water is in most places identified as shallow groundwater of meteoric origin (Stefánsson, 2017), although the hydrothermal systems along the Reykjanes peninsula are dominated by seawater. Surface manifestations in high-temperature areas are usually steam vents (fumaroles), mudpots and highly altered ground. Wells sunk in geothermal fields have typical depths of ~1.5-3 km and reservoir temperatures between 200 and 300°C (Arnórsson, 1995; Ármannsson, 2016). Higher temperature reservoirs supercritical/superheated reservoir temperatures have also been observed through the drilling activity of the Iceland Deep Drilling Project, for example at Krafla where drilling into magma (IDDP-1) at a depth of 2 km resulted in 440°C fluids (Elders et al., 2014) and at Reykjanes where the bottom-hole temperature of a ~4.5 km borehole (IDDP-2) was ~600°C (Bali et al., 2020).

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We collected fluid samples from steam vents (vapor) and geothermal wells (vapor + liquid) spanning the active volcanic zone in Iceland, from the Reykjanes geothermal well field in the SW, Kerlingarfjöll area in the central highlands, to the Krafla and Námafjall geothermal fields in the NE (Fig. 2A). The source of  $H_2O$  in most samples is meteoric, based on its isotopic composition ( $\delta D$  and  $\delta^{18}O$ ) generally corresponding to local precipitation (Stefánsson et al., 2017), except for one well sample from Reykjanes where the  $\delta^{18}O$  and  $\delta D$  values of the liquid water correspond closely with those of seawater (Stefánsson et al., 2017).

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# 2.2 Nisyros, Greece

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Nisyros (Fig. 2B) is the easternmost island of the Aegean volcanic island arc, and is located about 15 km west of the Turkish coast. With a total diameter of ~8 km, it is host to a 4-km-wide, 250-m-deep central caldera which contains several hydrothermal explosion craters with many

active fumaroles (Di Paola, 1974). The most recent of these craters were formed in a series of hydrothermal eruptions historically recorded in the years 1871, 1873 and 1887, during which large rock fragments were ejected and important quantities of steam discharged (Marini et al., 1993 and references therein). Two geothermal wells were drilled in the early 1980's, up to depths of 1816 and 1547 m, and indicated reservoir temperatures above 300°C (Chiodini et al., 1993). These wells identified the presence of carbonate basement and of shallow and deep permeable zones, with potassic-propylitic mineral assemblages at the bottom of the deepest well suggesting temperatures above 350°C (Marini et al., 1993). High reservoir temperatures were also reflected by the presence of high-enthalpy fluids in this deep permeable zone (Chiodini et al., 1993; Marini et al., 1993). It is thought that the deep aquifer was involved in the 1873 eruption, evidenced by the high Cl content of the discharged brine (Marini et al., 1993). The seawater-dominated geothermal system at Nisyros and the geochemistry of steam emanations have been well characterized (Kavouridis et al., 1999; Brombach et al., 2003; Fiebig et al., 2004; Fiebig et al., 2013). We sampled gas-rich steam from 9 fumaroles spanning four distinct phreatic craters within the central caldera (Fig. 2B). Previous investigations have suggested that CH<sub>4</sub> and CO<sub>2</sub> occur in chemical and isotopic equilibrium at a temperature of ~350°C (Fiebig et al., 2004; Fiebig et al., 2013), but the concentrations and  $\delta^{13}$ C signatures of higher hydrocarbons point to a thermogenic origin, suggesting that observed chemical and isotopic equilibrium may be coincidental (Fiebig et al., 2019).

## 3. Methods

Samples were collected in Iceland in August 2015 and in Nisyros in October 2017. For well samples, the liquid and vapor phases were separated using a Webre separator prior to sampling (Arnórsson et al., 2016). Samples of fumarole vapor were collected by inserting a titanium rod into the vapor outlet. Geothermal vapor was subsequently collected into various gas bottles.

For major gas determination, 125 ml evacuated glass flasks partially filled with ~10-50 ml 50% KOH (Iceland) or 4M NaOH (Nisyros) were used. The non-condensable gases including  $H_2$ ,  $N_2$ ,  $CH_4$  and Ar were analyzed using a gas chromatography (GC) system equipped with a thermal conductivity detector (TCD) whereas the condensable gases including  $H_2S$  and  $CO_2$  were analyzed in the vapor condensate using  $H_2$ -precipitation and modified alkalinity titrations, respectively (Arnórsson et al., 2006; Stefánsson et al., 2007). Separate dry-gas samples were collected for CO followed by GC analysis using a pulsed discharge detector (PDD) (Table 1).

Samples for  $\delta D_{H2O}$  measurements were collected into gas-tight glass bottles after condensation in line using a glass cooling spiral (Cioni and Corazza, 1981). For  $\delta D_{H2O}$  analysis, the hydrogen was extracted by the H<sub>2</sub>-water equilibration method using a Pt-catalyst (Horita, 1988) followed by analysis using a Finnegan MAT 251 ion ratio mass spectrometer (IRMS) at the University of Iceland and the results reported in terms of the conventional  $\delta$ -notation in ‰, relative to the VSMOW standard. Replicate analysis of the reference material yielded a standard deviation of 0.7‰.

Dry gas samples were collected for carbon isotope analysis of  $CO_2$  ( $\delta^{13}C_{CO2}$ ), which was performed at Goethe University, using a Flash EA 1112 (ThermoFisher) connected to a MAT 253 gas source mass spectrometer (Fiebig et al., 2004).  $CO_2$  from two bottles (Oztech), each calibrated

against VPDB, was measured along with the samples to correct for potential instrumental fractionations and scale compression. Hydrocarbon distribution ratios  $(C_1/C_{2+})$  were also determined at Goethe University as outlined by Fiebig et al. (2015).

Samples for methane clumped isotopologue determination were collected into 1L pre-evacuated glass bottles containing ~200 mL 50% KOH (Iceland) or 4M NaOH (Nisyros). Methane was extracted and purified from the gas phase of the flask samples by repeated cycles of vacuuming and flushing with a He carrier gas through a cold trap filled with activated charcoal submerged in liquid nitrogen (-196°C), as described by Wang et al. (2015). This traps gases such as CH<sub>4</sub>, N<sub>2</sub>, CO, and traces of CO<sub>2</sub> that remained in the gas phase. The trap was then heated to separate adsorbed gases through a gas chromatography column packed with Carboxen-1000 (MilliporeSigma, St Louis, MI) held at 30°C, and eluted methane was trapped again in a U-trap containing silica gel at liquid nitrogen temperature. The purified CH<sub>4</sub> samples ranged in volume from 0.7 to 10 ml (STP). The relative abundances of methane isotopologues <sup>12</sup>CH<sub>4</sub>, <sup>13</sup>CH<sub>4</sub>, <sup>12</sup>CH<sub>3</sub>D and <sup>13</sup>CH<sub>3</sub>D were measured by a tunable infrared laser direct absorption spectroscopy (TILDAS) instrument as described previously (Ono et al., 2014; Wang et al., 2015). Two samples from Nisyros were measured for <sup>12</sup>CH<sub>2</sub>D<sub>2</sub> (in addition to the four isotopologues mentioned above) by another TILDAS instrument and reported in Gonzalez et al. (2019).

Sample measurements were bracketed by measurements of CH<sub>4</sub> standard gases of known isotopic composition ( $\delta D$  and  $\delta^{13}C$ ) spanning a large  $\delta D$  range (~200 ‰), which were heated to 250°C for at least two weeks with a Pt catalyst to ensure isotopologue equilibrium (Ono et al., 2014). Isotope values are reported using standard delta notation against VPDB and VSMOW for the ratios  $^{13}C/^{12}C$  and D/H, respectively. This isotope scale was calibrated by the measurements of NGS-1 and NGS-3 (Wang et al., 2015). We used the reference  $\delta^{13}C$  values of -29.0 and -72.8‰, and  $\delta D$  of -138 and -176‰, for NGS-1 and NGS-3, respectively (Hut, 1987).

Results in Table 2 show the 95% confidence interval for the spectroscopic measurements, typically 0.2 to 0.4‰ for  $\Delta^{13}$ CH<sub>3</sub>D values. This does not include potential fractionation during sample preparation. One sample from Iceland (15-AS-03) was collected twice and analyzed as duplicates (Table 2), showing good reproducibility between duplicates with compounded errors similar to the 95% confidence intervals of the measurements (Table 2). Most samples (samples with less than 5ml CH<sub>4</sub>) were measured in a recycling mode, in which sample CH<sub>4</sub> is recovered from the TILDAS absorption cell to the cold trap, and reintroduced to TILDAS for 6 to 10 times. This allows repeated comparison of sample against reference gas. Thus, precision is not necessarily a function of sample size.

We define  $\Delta^{13}CH_3D$  as a measure of the abundance of  $^{13}CH_3D$  relative to a stochastic distribution (Ono et al., 2014):

$$\Delta^{13}CH_3D = \ln\left(\frac{^{13}CH_3D}{^{12}CH_3D} \cdot \frac{^{12}CH_4}{^{13}CH_4}\right)$$
 (2)

The relationship between  $\Delta^{13}CH_3D$  values and apparent methane generation temperatures used in this study is based on the formula:

$$\Delta^{13} \text{CH}_3 \text{D(T)} = -0.11006 \left(\frac{1000}{T}\right)^3 + 1.04151 \left(\frac{1000}{T}\right)^2 - 0.55235 \left(\frac{1000}{T}\right) \tag{3}$$

This equation is approximated from the solution based on the fundamental vibrational frequencies as described in Whitehill et al. (2017).

We note here that different theories predict a different temperature dependence for the value of  $\Delta^{13}$ CH<sub>3</sub>D (e.g. Eldridge et al., 2019; Liu and Liu, 2016; Wang et al., 2015; Webb and Miller, 2014). This will affect our results in two ways: one directly affects the reported  $\Delta^{13}CH_3D$  values and the other affects the equilibrium temperatures derived from given Δ<sup>13</sup>CH<sub>3</sub>D values. In this study, we used methane equilibrated at 250°C to anchor the  $\Delta^{13}$ CH<sub>3</sub>D value of our reference gas. We define this value as 1.98% Error! Reference source not found., which is 0.09% lower than the value of 2.07‰ obtained using the recent calibration by Eldridge et al. (2019). Our unknown sample measurements are calibrated against this 250°C heated methane, therefore in general  $\Delta^{13}$ CH<sub>3</sub>D values reported in this study (Table 2) are <0.1% lower than they would be using the equation of Eldridge et al. (2019). Apparent equilibrium temperatures are in turn calculated from these  $\Delta^{13}$ CH<sub>3</sub>D values using Error! Reference source not found. Due to the relative consistency of the temperature dependence of  $\Delta^{13}CH_3D$  applied both to our equilibrium reference  $CH_4$  at 250°C and to the observed range of  $\Delta^{13}$ CH<sub>3</sub>D values, the derived equilibrium temperatures do not differ significantly between different theories. For example, we would report a temperature of 350°C for a  $\Delta^{13}$ CH<sub>3</sub>D measurement of 1.34‰. The same measurement may yield e.g. 1.43‰ if the calibration of Eldridge et al. (2019) is used to anchor our reference gas, which would return a temperature of 346°C. Such a difference is considerably smaller than the 95% confidence intervals reported in Table 2, hence the application of different models is not expected to significantly alter our results.

# 4. Results

The chemical composition of Iceland and Nisyros steam samples are shown in Table 1. Isotopic compositions of H<sub>2</sub>O ( $\delta$ D), CO<sub>2</sub> ( $\delta$ <sup>13</sup>C) and CH<sub>4</sub> ( $\delta$ <sup>13</sup>C and  $\delta$ D), including clumped isotopologue signatures ( $\Delta$ <sup>13</sup>CH<sub>3</sub>D), are reported in Table 2. All samples have  $\delta$ <sup>13</sup>C<sub>CH4</sub> and  $\delta$ D<sub>CH4</sub> values that plot in a field originally proposed as 'abiotic' (Etiope and Sherwood-Lollar, 2013) but also defined as 'volcanic thermogenic' (Reeves and Fiebig, 2020) (Fig. 3, Table 2).

Iceland samples display a large isotopic variability, with  $\delta D_{\text{CH4}}$  ranging from -319.6‰ to -165.0‰ and  $\delta^{13}C_{\text{CH4}}$  ranging from -39.6‰ to -25.0‰. While some of the  $\delta D$  variability can be attributed to the source of water in the hydrothermal system (meteoric vs seawater), pronounced  $\delta^{13}C$  and  $\delta D$  heterogeneities are observed at the local scale, especially in the Krafla and Námafjall geothermal areas. In contrast, all Nisyros samples yield homogeneous  $\delta D_{\text{CH4}}$  and  $\delta^{13}C_{\text{CH4}}$  values, with  $\delta D_{\text{CH4}}$  ranging from -124.4‰ to -135.4‰ and  $\delta^{13}C_{\text{CH4}}$  from -21.8‰ to -23.4‰. The majority of samples have  $\Delta^{13}CH_3D$  values ranging from 0.82‰ to 1.77‰, corresponding to an apparent equilibrium temperature range from  $278^{+56}_{-44}$  to  $490^{+163}_{-99}$  °C, aligned with previous measurements of CH<sub>4</sub> isotopologues in geothermal systems (Fig. 4).

Additionally, measurements of  $\Delta^{13}\text{CH}_3\text{D}$  and  $\Delta^{12}\text{CH}_2\text{D}_2$  of two Nisyros samples (samples K7 and S4-hi) were made on a separate TILDAS instrument and reported by Gonzalez et al. (2019). The reported  $\Delta^{13}\text{CH}_3\text{D}$  values of 0.97  $\pm$  0.10‰ and 1.14  $\pm$  0.30‰, corresponding to apparent temperatures of  $447^{+26}_{-23}$  and  $400^{+90}_{-65}$ °C, agree within uncertainty with the values reported in this

study of 1.16  $\pm$  0.21‰ and 1.37  $\pm$  0.30‰ (390 $^{+56}_{-45}$  and 345 $^{+69}_{-52}$ °C, respectively). Low  $\Delta^{12}$ CH<sub>2</sub>D<sub>2</sub> values of 0.2  $\pm$  0.4‰ and -0.01  $\pm$  0.6‰ give apparent equilibrium temperatures of  $860^{+122}_{-91}$  and >769°C.

# 3463475. Discussion

# 5.1. Equilibrium and disequilibrium signals for chemical and isotopologue geothermometers

In order to interpret apparent temperatures derived from the methane isotopologue geothermometer, we compare them with several other chemical and isotope geothermometers, and with reservoir temperatures measured directly downhole, which are available for well samples. We first evaluate geothermometers commonly used for geothermal exploration, such as quartz solute equilibria (Gunnarsson and Arnórsson, 2000) and various gas geothermometers (Arnórsson and Gunnlaugsson, 1985; Arnórsson et al., 1998). We then explore geothermometers that involve methane isotopologues, which include those based on deuterium fractionation between  $H_2O-CH_4$  ( $\alpha_{H2O-CH4}$ ),  $^{13}C$  fractionation between  $CO_2-CH_4$  ( $\alpha_{CO2-CH4}$ ), and internal equilibrium characterized by the relative abundances of  $CH_4$  isotopologues ( $\Delta^{13}CH_3D$ ). These solute, gas and isotope geothermometers are tabulated in Table 3.

# 5.1.1. Measured, solute and gas geothermometry temperatures

Solute geothermometers are often applied to determine hydrothermal reservoir temperatures (Arnórsson et al., 1983b; Giggenbach, 1988; Gunnarsson and Arnórsson, 2000). The solubility of quartz (Table 3) is a widely applied geothermometer, which generally agrees with measured downhole temperatures (Table 4 and Fig. 5A). Hence, geothermal wells provide the advantage of sampling fluid directly from the hydrothermal reservoir where the temperatures are known with a reasonable degree of confidence, to which other geothermometers can be compared in order to gain insight on the evolution of the hydrothermal systems.

The majority of our samples, however, come from fumaroles, where measured or quartz temperatures are unavailable. Therefore, we must turn to gas compositions (Table 1) to estimate reservoir temperatures. The use of gas geothermometry relies on the concentrations of gas species within a separated vapor phase, which depends on the extent of fluid-rock interactions in the geothermal system and saturation of the fluid with various secondary minerals (e.g. calcite, epidote, pyrite), as well as on the vapor/liquid partition coefficients of the various gas species. Such geothermometers can be estimated by considering theoretical equilibria of various reactions (e.g. Chiodini and Marini, 1998; Giggenbach, 1980), or constructed empirically by calibrating measured concentrations with measured reservoir temperatures, e.g. from geothermal wells (e.g. Arnórsson et al., 1998; Arnórsson and Gunnlaugsson, 1985; D'Amore and Panichi, 1980). Here we combine three different gas geothermometers based on concentrations of CO<sub>2</sub>, H<sub>2</sub>S, and CO<sub>2</sub>/N<sub>2</sub> ratios (Table 3), which were empirically calibrated by Arnórsson and Gunnlaugsson (1985) and updated by Arnórsson et al. (1998). Figure 5B shows the comparison of gas temperatures to the measured reservoir temperatures for well samples. While the agreement is not as striking as that for quartz geothermometry (Fig. 5A), the temperatures are

generally within 50°C of each other. We also note that the two samples that are furthest from the measured temperatures, Krafla-16 and Námafjall-13, have excess enthalpy (see Table 4), thus the vapor phase composition is more likely to have been modified by boiling or phase segregation, which could have important effects on calculated gas temperatures (e.g. Arnórsson et al., 2007; Chiodini and Marini, 1998). This issue should be less critical for fumaroles, which emit steam at ~100°C and atmospheric pressure. Therefore, gas temperatures calculated for fumarole samples are taken to represent relatively well the temperatures of boiling in the hydrothermal reservoir at depth.

In general, Iceland samples have reservoir temperatures between 250°C and 300°C, which are systematically lower than  $\Delta^{13}\text{CH}_3\text{D}$ -based temperatures (Fig. 5C, Table 4). This effect is most pronounced for Krafla and Námafjall wells, while the fumarole sample from Krysuvík shows closest agreement between  $\Delta^{13}\text{CH}_3\text{D}$  and reservoir temperature. In contrast, gas concentrations from Nisyros samples point to reservoir temperatures around 350°C, which is in good agreement with previously-determined temperatures obtained with other independent geothermometers (Brombach et al., 2003) or based on metamorphic assemblages (Marini et al., 1993). Importantly, these are also in agreement with  $\Delta^{13}\text{CH}_3\text{D}$ -based temperatures (average ±1 s.d.) of  $355^{+50}_{-41}$  °C.

We also compare the temperatures and depths of two-phase well fluids to the pressure-temperature curve of boiling water (Fig. 6). Discharge temperatures and pressures measured at the well head reflect the cooling of ascending fluids through adiabatic boiling and thus usually fall on the boiling point curve. In contrast, temperatures measured at depth for the different wells are lower than those expected for boiling at hydrostatic pressure, reflecting the depressurization of fluids when they are intersected by geothermal wells (e.g. Arnórsson et al., 2007).

# 5.1.2. Deuterium fractionation between H<sub>2</sub>O and CH<sub>4</sub>

While comparison between measured reservoir temperatures, solute and gas geothermometry indicates close to mineral-fluid equilibration at subcritical temperatures (Fig. 5A, B), hydrogen isotope fractionation between H<sub>2</sub>O and CH<sub>4</sub> ( $\alpha_{H2O-CH4}$ ) indicates different degrees of disequilibrium and equilibrium (Fig. 7A, B). The variability is considered to reflect the sluggish kinetics of isotopic exchange between H<sub>2</sub>O and CH<sub>4</sub> (Table 3) due to the chemical stability of methane C-H bonds. Samples exhibit two linear trends of increasing  $\delta D_{CH4}$  values with  $\delta D_{H2O}$  around -90‰ and -10‰ (Fig. 7A), reflecting Icelandic meteoric water and seawater sources, respectively, where maximum  $\delta D_{CH4}$  values associated with each trend are indicative of isotopic equilibrium with the source water in the hydrothermal reservoirs. For example, local seawater is associated with higher  $\delta D_{CH4}$  for Nisyros and Reykjanes and low  $\delta D$  meteoric water is associated with lower  $\delta D_{CH4}$  for other Icelandic sites. Alternatively, the source water could exchange hydrogen isotopes with dissolved or particulate organic matter prior to thermogenic cracking, which would also cause subsequently generated CH<sub>4</sub> to have  $\delta D$  signatures partly derived from local H<sub>2</sub>O.

Steam vents in Nisyros island yielded relatively homogeneous  $\delta D_{CH4}$  values: CH<sub>4</sub> from three sites, Kaminakia, Stefanos and Polybotes Mikros, yielded an average  $\delta D_{CH4}$  of -125.5 ± 1.1% (± 1 s.d.), whereas Phlegeton yielded -135.4% (Table 2). There is a large variability in  $\delta D_{H2O}$ 

values, but this is mostly due to three outliers, with K7 having a very depleted  $\delta D_{H2O}$  value of 56.5%, and the two vents from Alexandros having  $\delta D_{H2O}$  of -4.5% and -4.8%. The depletion observed for K7 is likely due to removal of heavy water through partial condensation occurring during the upflow of steam, which is more likely to occur at the lower vent temperatures measured at Kaminakia (Table 4) (Brombach et al., 2003; Fiebig et al., 2004). The other samples have similar  $\delta D_{H2O}$  values with an average (±1 s.d.) of -11.0 ± 1.6%, which is indistinguishable from the parental hydrothermal liquid, previously suggested to be a mixture between seawater and "andesitic" water with  $\delta D_{H2O}$  of -11 ± 5% (Brombach et al., 2003; Fiebig et al., 2004). Therefore, overall  $\alpha_{CH4-H2O}$  values at Nisyros reflect apparent isotopic equilibrium around 300–350°C (Fig. 7A, B), consistent with gas geothermometry and previous determinations of the hydrothermal reservoir temperature (Brombach et al., 2003; Fiebig et al., 2004; Fiebig et al., 2013).

Methane from Kerlingarfjöll steam vents in Iceland also has  $\delta D_{CH4}$  values that can be interpreted as being in D-equilibrium with H<sub>2</sub>O at reservoir temperatures of ~250–300°C, if we infer values for  $\delta D_{H2O}$  between -80 and -90‰, corresponding to local precipitation (e.g. Stefánsson et al., 2017b) (Table 2 and Fig. 7A, B; "Ker inferred"). The geothermal well sample from Hædarendi also appears to be in H<sub>2</sub>O-CH<sub>4</sub> isotopic equilibrium around ~200°C (Fig. 7A), which is bracketed by the measured reservoir temperature of 170°C and gas temperature of 214°C (Table 4).

In contrast,  $CH_4$  sampled from geothermal wells in Krafla and Námafjall shows D-disequilibrium in the  $H_2O$ - $CH_4$  system, with  $\delta D_{CH4}$  values ranging from -320 to -243‰, which are depleted by approximately 30 to 100‰ compared to the value expected for equilibrium at reservoir temperatures of ~250–300°C (Fig. 7A). There is an array of samples from the most D-depleted (Krafla well 32) to near-equilibrium (Krafla well 16) values (Fig. 7A). Isotopic disequilibrium in the  $H_2O$ - $CH_4$  pair is also observed for the Reykjanes well, with an apparent temperature of <200°C, compared to a reservoir temperature of 290°C (Fig. 7A, Table 4).

We note that the accuracy of the  $\alpha_{H2O-CH4}$  thermometer is highly uncertain because this fractionation factor is derived by the combination of three fractionation factors,  $\alpha_{H2O(g)-H2(g)}$  (e.g. Suess, 1949),  $\alpha_{H2O(I)-H2O(g)}$  (Horita and Wesolowski, 1994) and  $\alpha_{CH4(g)-H2(g)}$  (Horibe and Craig, 1995). The compounded error (1σ) of these fractionation factors is shown as grey shading in Fig. 7B, and illustrates that for a given  $\alpha$  value the possible temperature range (within the  $1\sigma$  uncertainty) spans ~100°C, notably in the 250–374°C range, where the temperature sensitivity of  $\alpha_{H20-CH4}$  is greatly reduced. Since α<sub>H2O-CH4</sub> is simply a small difference between two larger numbers (α<sub>H2O-H2</sub> and  $\alpha_{CH4-H2}$ ), its uncertainty is large relative to its small variations in magnitude. For example,  $\alpha_{H2O-}$ <sub>CH4</sub> ranges from 1.154 at 200°C to 1.123 at 400°C (Horibe and Craig, 1995), while  $\alpha_{H2O-H2}$  ranges from 1.977 to 1.469 and  $\alpha_{CH4-H2}$  from 1.719 to 1.304 over the same temperature interval. It must also be noted that the equilibrium line, based on the calibration by Horibe and Craig (1995), relies on the  $\alpha_{H2O(g)-H2(g)}$  fractionation factor derived by Suess (1949), but that use of other  $\alpha_{H2O(g)-H2(g)}$ relationships (e.g. Bardo and Wolfsberg, 1976; Cerrai et al., 1954) would yield different curves (dashed lines in Fig. 7B). Furthermore, another source of uncertainty arises from the calibration of the  $\delta D_{CH4}$  scale against the VSMOW reference scale as this requires measurement of  $\delta D$  values of reference H<sub>2</sub>O and CH<sub>4</sub> at high precision. However, these differences are small compared to

the measured  $H_2O$ – $CH_4$  fractionations of ~100–300% (Fig. 7A) and do not change the nature of our interpretations.

# 5.1.3. Clumped isotopologue distributions

In comparison to the wide range of  $\delta D_{CH4}$  values,  $\Delta^{13}CH_3D$  signals of methane are largely homogeneous (0.82  $\pm$  0.34 to 1.77  $\pm$  0.34‰), and correspond to apparent equilibrium temperatures ranging from 278 to 490°C (Fig. 4). If  $\Delta^{13}CH_3D$  signals reflect methane formation or secondary equilibration temperature, the inferred temperatures would significantly exceed the thermogenic gas window of ~150 to 220°C typical of sedimentary basins (e.g. Quigley and Mackenzie, 1988; Stolper et al., 2017).

Among measured samples, well samples from Krafla and Námafjall yielded lower  $\Delta^{13}\text{CH}_3\text{D}$  values with an average (±1 s.d.)  $T_{\Delta^{13}CH_3D}=438^{+55}_{-45}^{\circ}\text{C}$ , compared to the rest of the samples, which have generally higher  $\Delta^{13}\text{CH}_3D$  values with an average (±1 s.d.)  $T_{\Delta^{13}CH_3D}=349^{+44}_{-37}^{\circ}\text{C}$  (Table 2). While the four well samples from the Krafla and Námafjall area have a narrow  $\Delta^{13}\text{CH}_3D$  range pointing to the highest apparent clumped temperatures, they exhibit a large range in  $\delta D_{\text{CH}4}$  that indicates pronounced departures from isotopic equilibrium against H<sub>2</sub>O (Fig. 7B). We suggest that sample Krafla well 32, which has the lowest  $\delta D_{\text{CH}4}$ ,  $\delta^{13}\text{C}_{\text{CH}4}$  and  $\Delta^{13}\text{CH}_3D$  values, represents a kinetic end-member, and that the array of data from Krafla and Námafjall reflects different degrees of equilibration from initial kinetic methane (Fig. 5C and Fig. 7A, B). Methane from geothermal wells at Reykjanes and Hædarendi also has  $\Delta^{13}\text{CH}_3D$  values which do not correspond to measured reservoir temperature (Fig. 5C) and the  $1000\ln\alpha_{\text{H}2O\text{-CH}4}$  geothermometer (Fig. 7B). In contrast, samples from Nisyros and Kerlingarfjöll have  $\Delta^{13}\text{CH}_3D$  values which generally agree with the apparent equilibrium temperatures obtained from  $\alpha_{\text{H}2O\text{-CH}4}$  (Fig. 7B), suggesting that they represent equilibrium end-member methane. We examine the rates of deuterium exchange and isotopic equilibration to explain  $\delta D_{\text{CH}4}$  and  $\Delta^{13}\text{CH}_3D$  variability in section 5.3.1.

# 5.1.4. <sup>13</sup>C fractionation between CO<sub>2</sub> and CH<sub>4</sub>

Similar trends of equilibrium and disequilibrium methane are observed in  $^{13}\text{C}_{\text{cractionation}}$  fractionation between CO<sub>2</sub>–CH<sub>4</sub>, where the most negative  $\delta^{13}\text{C}_{\text{CH4}}$  values observed at Krafla, Námafjall and Reykjanes are associated with disequilibrium (i.e. kinetic) signals, while CH<sub>4</sub> from Nisyros and Kerlingarfjöll appears to show equilibrium signals relative to  $\Delta^{13}\text{CH}_3\text{D}$ , with apparent isotopic temperatures from 300 to 350°C (Fig. 7C, D). Variations in the fractionation factor  $\alpha_{\text{CO2-CH4}}$  are primarily due to variations in  $\delta^{13}\text{C}_{\text{CH4}}$ , since  $\delta^{13}\text{C}_{\text{CO2}}$  is relatively constant for all samples from Iceland ( $\delta^{13}\text{C}_{\text{CO2}}$  = -2.4 to -3.8‰) and Nisyros ( $\delta^{13}\text{C}_{\text{CO2}}$  = -0.8 to -1.9‰). While the CO<sub>2</sub>-CH<sub>4</sub> apparent isotopic temperatures (Fig. 7C) at Krafla and Námafjall are not so far from reservoir temperatures between ~250-290°C (e.g. Fig. 5, Table 4), we qualify them as being in disequilibrium relative to the  $\Delta^{13}\text{CH}_3\text{D}$  geothermometer which indicates temperatures of  $438^{+55}_{-45}$ °C. In contrast, the sample from Hædarendi, which is a high CO<sub>2</sub>-producing well (Table 1), shows simultaneous apparent equilibrium between the  $1000\text{In}\alpha_{\text{CO2-CH4}}$  and  $\Delta^{13}\text{CH}_3\text{D}$  geothermometers at a temperature of ~300°C, which is however significantly hotter than the hydrothermal reservoir temperature of 200°C.

The relatively narrow range in  $\delta^{13}C_{CH4}$  values of Krafla and Námafjall well samples, from -39.6‰ to -35.8‰, compared to their large  $\delta D_{CH4}$  range from -320‰ to -243‰, is consistent with pyrolysis of terrestrial xylite, according to the fractionation factors derived by Berner et al. (1995). However, the two Krafla samples are farthest from each other in  $\delta D_{CH4}$  space but closest to each other in  $\delta^{13}C_{CH4}$  space (Fig. 7A, C), which could mean that while the CH<sub>4</sub> generation process may lead to disequilibria in both  $\delta^{13}C_{CH4}$  and  $\delta D_{CH4}$ , mechanisms controlling D and  $^{13}C$  isotopic equilibration are different. Alternatively, this observation could reflect slight variations in the isotopic composition of source organics. Since the mechanism of carbon isotope exchange between CO<sub>2</sub> and CH<sub>4</sub> is not well understood (e.g. McCollom, 2013) and indeed its use as a geothermometer for natural systems has been put into question (Fiebig et al., 2019),  $^{13}C$  fractionations must be interpreted with caution. Therefore, while CH<sub>4</sub> from Nisyros, Kerlingarfjöll and Hædarendi appears to show simultaneous  $\Delta^{13}CH_3D$  and  $\alpha_{CO2\text{-}CH4}$  equilibrium, we must discuss the rates of isotopic exchange (c.f. section 5.3.2) to assess whether this reflects "true" attainment of equilibrium.

# 5.2. Expected isotopologue signals of thermogenic and abiotic methane

In the previous section, we identified equilibrium and kinetic end-member methane, the latter characterized by low  $\delta D_{CH4}$  and  $\Delta^{13}CH_3D$  values. Here, we discuss whether  $\Delta^{13}CH_3D$  values of "kinetic" methane can place constraints on a thermogenic versus abiotic origin. Disequilibrium  $\Delta^{13}$ CH<sub>3</sub>D signals have been measured for methane generated in laboratory experiments via thermal cracking of organic matter (Shuai et al., 2018a). Other experiments that produced methane through abiotic reactions (Young et al., 2017) have also shown strong  $\Delta^{12}CH_2D_2$ disequilibria along with minor  $\Delta^{13}CH_3D$  disequilibria. The high apparent temperatures associated with 'kinetic' methane at Krafla could thus reflect disequilibrium isotopologue signals from either thermogenic or abiotic methane (Fig 1, yellow ovals). On the other hand, the geothermal systems at Krafla and Námafjall are characterized by steep geothermal gradients, due to the active Krafla magma chamber at a depth of just over 2 km. While the  $\Delta^{13}CH_3D$ -based temperature of  $438^{+55}_{-48}$ °C obtained for sampled wells is higher than reservoir temperatures of ~250-300°C, it is indistinguishable from the temperature of ~440°C measured in fluids at a ~2 km depth during drilling of the IDDP-1 well (Elders et al., 2014; Scott et al., 2015; Heřmanská et al., 2019). Hence, methane could be sourced from a deep superheated reservoir in close vicinity to the magmatic heat source, either thermogenically or abiotically (Fig. 1, red ovals). In any case, if the high apparent clumped temperatures are the result of kinetic processes, these effects may be relatively small, since high temperatures are prevailing in this geothermal system.

# 5.2.1. Isotopologue signals for thermogenic methane

Thermogenic methane produced in sedimentary basins peaks at temperatures between 150 and 220°C (e.g. Quigley and Mackenzie, 1988; Tissot and Welte, 1978). Methane from thermogenic natural gas deposits tends to yield clumped temperatures that usually agree with independently constrained gas formation temperatures (Ono et al., 2014; Stolper et al., 2014a; Wang et al., 2015; Stolper et al., 2017; Shuai et al., 2018b). In addition, thermogenic methane

from hydrothermally heated sediments at Guaymas Basin (Wang et al., 2015) and Juan de Fuca Ridge (Douglas et al., 2017) yielded temperatures of 326°C and 305 to 365°C, respectively, consistent with measured vent fluid temperatures. However, non-equilibrium isotopologue signatures are implied in some unconventional oil-associated reservoirs (Douglas et al., 2017), with derived  $T_{\Delta_{18}}$  exceeding 200°C, the estimated maximum temperature for oil stability under geological timescales (e.g. Hunt, 1996).

Shuai et al. (2018a) investigated the clumped isotopologue signals of experimentally produced thermogenic CH<sub>4</sub> by subjecting coal and shale to pyrolysis at temperatures from 380 to 620°C. Both series of experiments were characterized by near-equilibrium signatures during the early stages of thermal cracking, followed by departures from isotopologue equilibrium with increasing temperatures, in which lower-than-expected  $\Delta_{18}$  values were produced. This led the authors to suggest that CH<sub>4</sub> generated by secondary cracking of C<sub>2-5</sub> hydrocarbons can carry clumped isotopologue signatures characterized by pronounced kinetic isotope effects. These experimental results suggest that kerogen cracking produces methane in clumped isotopologue equilibrium, possibly due to local reversibility in hydrogen exchange, whereas cracking of alkyl groups from oil or C<sub>2+</sub> gases irreversibly generates methane with non-equilibrium clumped signatures (Shuai et al., 2018a).

The conditions that may lead to thermogenic methane generation in hydrothermal systems are quite different from the "dry" experimental conditions investigated by Shuai et al. (2018a), and may thus evoke a different clumped pattern. While these pyrolysis experiments used shale and coal as starting organic materials, thermal cracking in Icelandic hydrothermal fluids is thought to occur principally via the breakdown of organic matter transported to depths by external waters, since no organic sediments are present in these systems (Fiebig et al., 2019). In this respect, the starting organic matter would either represent or be derived from DOC and POC. Moreover, the most pronounced kinetic effects observed by Shuai et al. (2018a) occur when the ratio of  $C_{2-5}/C_{1-5}$  decreases between 0.1 and 0.01 (or as  $C_1/C_{2-5}$  increases from ~10<sup>1</sup> to 10<sup>2</sup>), followed by a return to equilibrium as this ratio goes beyond  $10^2$ . The measured  $C_1/C_{2+}$  ratios in our hydrothermal fluids fall within the range 10<sup>2</sup>–10<sup>4</sup> (Table 1), and could therefore be more analogous to the highest thermal maturities in the experiments of Shuai et al. (2018a) associated with equilibrium clumped signatures. Additionally, the non-equilibrium effects shown by the experiments of Shuai et al. (2018a) co-evolve with increases in  $\delta D$  and  $\delta^{13}C$  values, mirroring increases in thermal maturity. In contrast, such correlation is not observed with the  $\Delta^{13}CH_3D$ values measured in Krafla and Námafjall well fluids, which remain uniformly low over ranges of ~80% in  $\delta D$  and ~5% in  $\delta^{13}C$  values similar to those observed by Shuai et al. (2018a) (Fig. 7B, D). Nonetheless, Krafla and Námafjall may be characterized by fast recharge of fluids and input of immature, low-δ<sup>13</sup>C organic matter, which could produce thermogenic CH<sub>4</sub> with non-equilibrium  $\Delta^{13}$ CH<sub>3</sub>D values, though the magnitude of such kinetic process cannot be evaluated.

# 5.2.2 Isotopologue signals for abiotic methane

Abiotic methane can be produced by a variety of processes and under a wide range of temperatures and pressures, as demonstrated in a growing number of laboratory experiments (Fu et al., 2007; McCollom et al., 2010; Taran et al., 2010; McCollom, 2013). It is expected that

abiotic methane formed by different processes and carbon sources should carry different isotopologue signatures. In natural settings, methane of potentially abiotic origin seems to have higher  $\delta^{13}$ C values compared to conventional biogenic and thermogenic methane, but its  $\delta$ D values largely overlap with these conventional sources (Fig. 3; Reeves and Fiebig, 2020).

Compared to the wide range of  $\delta^{13}$ C and  $\delta D$  values of proposed abiotic methane (Fig. 3), methane from non-sedimented seafloor hydrothermal vents tends to carry homogeneous isotope and isotopologue signals, with  $\delta^{13}$ C,  $\delta D$ , and  $\Delta^{13}$ CH<sub>3</sub>D values ranging from -11 to -18‰, -98 to -127‰, and 0.95 to 3.0‰, respectively, with  $\Delta^{13}$ CH<sub>3</sub>D values above 1.85‰ only observed at the Lost City hydrothermal field (Proskurowski et al., 2008; Wang et al., 2018; Labidi et al., 2020). These relatively uniform isotopologue signals suggest a common origin for methane associated with seafloor hydrothermal vents, proposed to be fluid trapped as fluid inclusions in the oceanic lithosphere (Kelley, 1996; Wang et al., 2018; Klein et al., 2019; Labidi et al., 2020). The kinetic end-member CH<sub>4</sub> measured in this study (Krafla-32) is significantly more depleted in  $\delta^{13}$ C and  $\delta D$  than CH<sub>4</sub> from seafloor hydrothermal vents (even when correcting for the ~80‰ shift between Iceland meteoric water and VSMOW; see Fig. 7A), suggesting that the generation processes of methane for Krafla-32 are different from those of abiotic CH<sub>4</sub> from seafloor vents.

Experimental studies of abiotic methane generation have shown that produced hydrocarbons, including CH<sub>4</sub>, are depleted in  $^{13}$ C compared to the initial carbon source, but the magnitude of the fractionation varies (McCollom, 2013 and references therein). For instance, Taran et al. (2010) carried out open-system Fischer-Tropsch-type (FTT) synthesis of CH<sub>4</sub> via the hydrogenation of CO<sub>2</sub> on iron and cobalt catalysts at 350 and 245°C, respectively, and observed  $^{13}$ C fractionation of >40% between CO<sub>2</sub> and CH<sub>4</sub>. Fu et al. (2007) carried out experiments of CO<sub>2</sub> reduction by H<sub>2</sub> in aqueous NaCl fluids using magnetite as a potential catalyst, at 400°C and 500 bars, and observed  $^{51}$ C isotopic fractionations between CO<sub>2</sub> and CH<sub>4</sub> from 4.6 to 27.1%, whereas the value expected for equilibrium at 400°C is 19‰. They also reported relatively constant  $^{50}$ D fractionation between H<sub>2</sub>O–CH<sub>4</sub> of 214 to 229‰ (equilibrium = 132‰), with CH<sub>4</sub> having lower  $^{50}$ D values than expected for equilibrium fractionation. Therefore, bulk isotopic signatures of abiotic CH<sub>4</sub> produced in the laboratory can exhibit significant variability and overlap with our kinetic end-member Krafla-32.

Young et al. (2017) investigated the clumped isotopologue signatures of abiotic methane produced by two different processes: 1) CH<sub>4</sub> generated via the Sabatier reaction (1) by reacting gaseous mixtures of CO<sub>2</sub> and H<sub>2</sub> with a Re catalyst at 70 and 90°C and 2) hydrolysis of silane (Si<sub>5</sub>C<sub>12</sub>H<sub>36</sub>) at temperatures of 300 to 600°C and 100MPa. The product CH<sub>4</sub> in several experiments showed  $\Delta^{13}$ CH<sub>3</sub>D values that agreed with generation temperatures, although some experiments had  $\Delta^{13}$ CH<sub>3</sub>D deficits of up to ~1‰. Comparatively large deficits in  $\Delta^{12}$ CH<sub>2</sub>D<sub>2</sub> values were observed, particularly for lower temperature experiments. Proposed abiotic methane from deep mine fluids in Precambrian shields (e.g. Sherwood Lollar et al., 2006; 1993) was also shown to carry lower-than-equilibrium  $\Delta^{12}$ CH<sub>2</sub>D<sub>2</sub> signatures combined with equilibrium  $\Delta^{13}$ CH<sub>3</sub>D values, and could thus be produced similarly (Young et al., 2017). These observations suggest that  $\Delta^{13}$ CH<sub>3</sub>D signatures may be relatively insensitive to kinetics. The isotopologue systematics of hydrothermal CH<sub>4</sub> generation from CO<sub>2</sub> reduction, however, have not been studied, making it hard to evaluate whether CH<sub>4</sub> measured in this study carries abiotic signals.

The  $\Delta^{12}\text{CH}_2\text{D}_2$  values reported for Nisyros steam vents (Gonzalez et al., 2019) and shown in Table 2 are slightly lower than expected for equilibrium with  $\Delta^{13}\text{CH}_3\text{D}$  and point to higher apparent temperatures, thus they could reflect non-equilibrium processes similar to those described by Young et al. (2017). However, as we outline in the next section, the residence times of Nisyros fluids are thought to be long enough for  $\Delta^{13}\text{CH}_3\text{D}$  signatures to equilibrate, and  $\Delta^{12}\text{CH}_2\text{D}_2$  should equilibrate at least twice as quickly (Labidi et al., 2020), albeit with potentially more pronounced initial kinetic signals. Therefore, the  $\Delta^{12}\text{CH}_2\text{D}_2$  deficits reported by Gonzalez et al. (2019) remain difficult to explain. However, we note that these deficits are small (on the order of 1‰) compared to the large kinetic signals (>10‰) exhibited by the experiments of Young et al. (2017) and could reflect uncertainty associated with sample preparation, which was not characterized.

5.3. Closure temperature of CH<sub>4</sub> in geothermal fluids

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5.3.1 Isotopic exchange between H<sub>2</sub>O and CH<sub>4</sub>

In this section, we evaluate if observed isotopologue equilibrium and disequilibrium signals can be explained by different rates of isotope exchange reactions. Upon cooling of fluids, isotope exchange reactions that operate at faster rates are more likely to re-equilibrate at lower temperatures.

Ohmoto and Lasaga (1982) derived the following rate law for isotope exchange between chemical species X and Y:

$$\ln\left(\frac{\alpha^e - \alpha}{\alpha^e - \alpha^0}\right) = \ln(1 - F) = -k([X] + [Y])t \tag{4a}$$

where  $\alpha^e$ ,  $\alpha^0$  and  $\alpha$  are the isotope fractionation factors at equilibrium, at time 0 and time t, respectively; F is the fraction of isotopic exchange (or progress variable); k is the second order rate constant (e.g. units in cm³ mol⁻¹ s⁻¹); [X] and [Y] are concentrations of reactants (such as H<sub>2</sub>O and CH<sub>4</sub>) and t is time. Equation (4a) shows that the rate of isotope exchange depends upon the sum of the two reactants. This concentration dependence is important for CH<sub>4</sub>/H<sub>2</sub> and CO<sub>2</sub>/CH<sub>4</sub>, where the rate of exchange is limited by the collision of two chemical species. However, when considering isotope exchange with H<sub>2</sub>O (H<sub>2</sub>O-CH<sub>4</sub> and H<sub>2</sub>O-H<sub>2</sub>), we can assume that H<sub>2</sub>O is an infinite reservoir, and equation (4a) reduces to:

$$\ln\left(\frac{\alpha^e - \alpha}{\alpha^e - \alpha^0}\right) = -k't \tag{4b}$$

where  $k' \simeq k[H_2O]$  is the pseudo first order rate constant (e.g. in units of yr<sup>-1</sup>). The same rate law was derived by Pester et al. (2018) for D/H exchange between H<sub>2</sub>O and H<sub>2</sub>.

If the methane isotopologue exchange reaction proceeded via internal equilibration (e.g. Table 3), then  $k' = k([^{12}CH_4 + ^{12}CH_3D] + [^{13}CH_4 + ^{13}CH_3D]) \simeq k[CH_4]$ , and the overall isotopologue exchange rate would be proportional to  $[CH_4]$ . That is, a faster exchange rate would be expected when  $CH_4$  concentration is higher. This is understood as the frequency of  $CH_4$  to  $CH_4$  collisions increases as the square of the concentration of  $CH_4$ . This type of exchange has been suggested by recent studies evaluating the kinetics of dual clumped  $CH_4$  isotope bond reordering (Labidi et

al., 2020; Giunta et al., 2021). Methane isotopologue equilibration in water-dominated systems, however, likely proceeds by D/H exchange with water via the two separate reactions:

$$^{12}\text{CH}_3\text{D} + \text{H}_2\text{O} \rightleftharpoons ^{12}\text{CH}_4 + \text{HDO}$$
 (5)

$$^{13}\text{CH}_3\text{D} + \text{H}_2\text{O} \rightleftharpoons ^{13}\text{CH}_4 + \text{HDO}$$
 (6)

In this case,  $k'=k[H_2O]$ , and the isotopologue exchange rate becomes independent of CH<sub>4</sub> concentration.

Based on experiments performed at different temperatures (e.g. Koepp, 1978; Reeves et al., 2012), Wang et al. (2018) derived an empirical rate law to describe this exchange as:

$$\log k' = 15.247 - \frac{9994.7}{T} \tag{7}$$

where k' is in yr<sup>-1</sup>, and T is temperature in Kelvin. This rate law corresponds to a relatively high activation energy of 191 kJ/mol.

We propose that the elementary reactions that characterize D/H isotope exchange for methane involve H-abstraction from  $CH_4$  by H radicals (reaction 8). The product methyl radical quickly reacts with  $H_2O$  to form  $CH_4$  and an OH radical (reaction 9).

$$CH_4 + {}^{\bullet}H \rightarrow {}^{\bullet}CH_3 + H_2 \tag{8}$$

$${}^{\bullet}\text{CH}_3 + \text{H}_2\text{O} \rightarrow \text{CH}_4 + {}^{\bullet}\text{OH} \tag{9}$$

The activation energy for reaction (8) is 88 kJ/mol (Sutherland et al., 2001). The high apparent activation energy of 191 kJ/mol — based on equation (7) — suggests that the overall exchange rate could be limited by radical initiation reactions such as:

$$H_2O \rightarrow H + OH$$
 (10)

$$H_2 \rightarrow 2^{\bullet}H$$
 (11)

These reactions are highly endothermic, with enthalpies of reactions of 505 and 441 kJ/mol at 700K for (10) and (11), respectively (GRI-Mech 3.0 - Smith et al., 2000), which may partly explain the high activation energy of 191 kJ/mol of the D/H exchange reactions (reactions (5) and (6)). The formation of such radicals could be catalyzed by rock surfaces, aided by opening of fractures in tectonically active environments and by the electrical conductivity of water (Kita et al., 1982; Balk et al., 2009). Under geothermal conditions, HS radicals could also be important in abstracting H from CH<sub>4</sub>. This may also have been the case for the experiments of Reeves et al. (2012), which were used to establish the rate law for D/H exchange between CH<sub>4</sub> and H<sub>2</sub>O (Eq. 7), since they were carried out at the pyrite-pyrrhotite-magnetite buffer.

We used equation (7) to model the evolution of methane isotopologue ratios ( $\Delta^{13}\text{CH}_3\text{D}$  and  $\delta\text{D}$ ) with a starting isotopic composition corresponding to our kinetic end-member Krafla well 32 (15-AS-07), with  $\delta\text{D}_{\text{CH4}}$  = -319.6%,  $\delta\text{D}_{\text{H2O}}$  = -82.4 % (1000ln $\alpha_{\text{H2O-CH4}}$  = 300%) and  $\Delta^{13}\text{CH}_3\text{D}$  = 0.82%. We use this sample as an end-member since it has the lowest  $\delta\text{D}$ ,  $\delta^{13}\text{C}$  and  $\Delta^{13}\text{CH}_3\text{D}$  signatures among measured samples in this study. We also note that this  $\delta\text{D}_{\text{CH4}}$  value is at the lower end of geothermal samples reported by Fiebig et al. (2019) (see Fig. 7A) and is therefore our best estimate for the kinetic end-member. However, as outlined below, the nature of our interpretations does not change if we use a different end-member, e.g. with lower  $\delta\text{D}_{\text{CH4}}$  and  $\Delta^{13}\text{CH}_3\text{D}$ .

Isotopic fractionation according to reactions (5) and (6) can be described by:

 $^{12}\alpha_{\text{H2O-CH4}} = \frac{[^{12}\text{CH}_4] \cdot [\text{HDO}]}{[^{12}\text{CH}_3\text{D}] \cdot [\text{H}_2\text{O}]}$ (12)

$$^{13}\alpha_{\text{H2O-CH4}} = \frac{[^{13}\text{CH}_4] \cdot [\text{HDO}]}{[^{13}\text{CH}_3\text{D}] \cdot [\text{H}_2\text{O}]}$$
(13)

- 723 Similarly, the CH<sub>4</sub> clumped isotopologue distribution (see Table 3 for the isotope exchange
- 724 reaction) can be written as:

$$Q = \frac{[^{13}\text{CH}_3\text{D}] \cdot [^{12}\text{CH}_4]}{[^{12}\text{CH}_3\text{D}] \cdot [^{13}\text{CH}_4]}$$
(14)

725 Hence, it follows that:

$$Q = {}^{12}\alpha/{}^{13}\alpha \tag{15}$$

We assume in our calculations that the equilibrium  $\alpha_{\text{H2O-CH4}}$  relationship derived by Horibe and Craig (1995) (Table 3) corresponds to  $^{12}\alpha_{eq}$  values since the abundance of  $^{12}\text{CH}_4$  is two orders of magnitude higher than  $^{13}\text{CH}_4$ . Then,  $^{13}\alpha_{eq}$  is calculated from  $^{12}\alpha_{eq}$  and  $K_{eq}$ , the value of Q representing clumped isotopologue equilibrium, using equations (2) and (3). Therefore, we model the evolution of  $^{12}\alpha$ ,  $^{13}\alpha$  and Q, at different temperatures following equations (4b), (7) and (15), and starting from the end-member composition (Krafla-32), as well as from an arbitrary end-member with more pronounced disequilibrium (1000ln $\alpha_{\text{H2O-CH4}}$  = 400% and  $\Delta^{13}\text{CH}_3\text{D}$  = 0%). The resulting calculations are reported in Table S1 and illustrated in Figures 7B and 8A, B. Equilibration paths are shown for temperatures of 450°C, 350°C and 250°C. This allows us to explore the equilibration timescales of methane isotopologues across a range of hydrothermal conditions. It is worth noting that the above equations imply that equilibration of  $\delta D_{\text{CH4}}$  and  $\Delta^{13}\text{CH}_3\text{D}$  occur through the same mechanisms and thus have the same timescales. Therefore, equilibration paths shown in Figure 7B are nearly straight lines. Equations 4b and 7 also imply that the timescale of equilibration will be the same for a given temperature regardless of the initial extent of disequilibrium (Fig. 8A, B).

Equilibration at 350°C, for example in the geothermal reservoir at Nisyros, is expected to be achieved within 10-100 years, which is consistent with the observation that CH<sub>4</sub> at Nisyros is in isotopologue equilibrium, considering that the last hydrothermal eruption, which may have introduced disequilibrium in the deep permeable zone, occurred more than 100 years ago (Marini et al., 1993). Figures 8A, B also show that at 250°C, there should be virtually no isotopic

exchange occurring at timescales relevant for this study (<1000 years), thus any methane formed or entrained into the subcritical hydrothermal reservoirs of all of the Icelandic geothermal systems studied here, which have reservoir temperatures of 250–300°C (Table 4), should retain its initial isotopologue ratios. Therefore, apparent H<sub>2</sub>O-CH<sub>4</sub> isotopic equilibrium at the reservoir temperature of ~200°C in the Hædarendi well is unlikely to reflect true attainment of equilibrium. Radium isotopes have been used to estimate fluid residence times of <5 years in Reykjanes wells (Kadko et al., 2007), which would not be sufficient for H<sub>2</sub>O-CH<sub>4</sub> isotopic equilibrium to be attained at the reservoir temperature of 290°C (Fig. 8A, B). This can explain the observation that  $\Delta^{13}$ CH<sub>3</sub>D and  $\alpha_{\text{H2O-CH4}}$  are not in mutual equilibrium for the Reykjanes well (Fig. 7B). The apparent  $\Delta^{13}$ CH<sub>3</sub>D-based temperature for this well is  $379^{+37}_{-32}^{\circ}$ °C, which could represent a kinetic signal from methane formation in the hydrothermal reservoir, but could also imply methane formation in deeper, hotter parts of the geothermal system which have been characterized by deep drilling (e.g. Bali et al., 2020).

At 450°C, which could represent a superheated vapor phase in the vicinity of magmatic intrusions at Krafla and Námafjall (Elders et al., 2014; Heřmanská et al., 2019), equilibration of  $\delta D_{\text{CH4}}$  and  $\Delta^{13}\text{CH}_3\text{D}$  values is expected to occur within days to one month (Fig. 8A, B). Therefore, if methane is generated in such phase, the superheated vapors must rapidly ascend into the overlying ~250-300°C subcritical reservoir (e.g. Figs. 1, 6) to preserve the observed range of disequilibrium  $\delta D_{\text{CH4}}$  values. Further equilibration at reservoir temperatures would likely proceed more slowly than fluid residence times of ~5-50 years (e.g. Stefánsson et al., 2015). Hence, if the range of  $\delta D_{\text{CH4}}$  values for Krafla and Námafjall reflects high-temperature equilibration in a superheated vapor phase, the  $\Delta^{13}\text{CH}_3\text{D}$  value of kinetic end-member Krafla-32 actually closely records the expected high temperature of superheated vapor. This would imply that CH<sub>4</sub> is formed with near-equilibrium  $\Delta^{13}\text{CH}_3\text{D}$  signatures but pronounced  $\delta D_{\text{CH4}}$  deficits. However, it is also possible that CH<sub>4</sub> forms at reservoir temperatures of 250–300°C with both kinetic  $\delta D$  and small  $\Delta^{13}\text{CH}_3\text{D}$  deficits, which would be preserved due to slow kinetics. The range in  $\delta D_{\text{CH4}}$  values of Krafla and Námafjall samples would in this case be explained by varying thermal maturity of degrading organic matter.

# 5.3.2. Rates and mechanisms of carbon isotopic exchange between CH<sub>4</sub> and CO<sub>2</sub>

The rates and mechanisms of chemical and isotopic equilibration between CH<sub>4</sub> and CO<sub>2</sub> (1) are poorly constrained but thought to be much slower than the D/H isotope exchange between CH<sub>4</sub> and H<sub>2</sub>O (Giggenbach, 1997; Cole and Chakraborty, 2001). While overall isotopic exchange can be represented by the <sup>13</sup>C exchange reaction shown in Table 3, Giggenbach (1997) points out that this reaction is unlikely since it requires a complete rearrangement of two C=O bonds and four C-H bonds between reacting CO<sub>2</sub> and CH<sub>4</sub> molecules. By analogy to the mechanism of S-isotope exchange between H<sub>2</sub>S and H<sub>2</sub>SO<sub>4</sub> proposed by Ohmoto and Lasaga (1982), carbon isotope exchange between CH<sub>4</sub> and CO<sub>2</sub> could proceed via two-carbon intermediate species such as acetic acids from a series of reactions such as association (16), intramolecular C-isotope exchange (17), or dissociation reactions (18) (Blake and Jackson, 1969; Mackie and Doolan, 1984; Duan and Page, 1995):

$$^{13}\text{CO}_2 + ^{12}\text{CH}_4 \rightarrow ^{12}\text{CH}_3 + ^{13}\text{COOH}$$
 (16)

$$^{12}\text{CH}_3^{13}\text{COOH} \rightleftharpoons ^{13}\text{CH}_3^{12}\text{COOH}$$
 (17)

$$^{13}\text{CH}_3^{12}\text{COOH} \rightarrow ^{13}\text{CH}_4 + ^{12}\text{CO}_2$$
 (18)

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While an experimental study by Dias et al. (2002) demonstrated that C-isotope exchange between CO<sub>2</sub> and the carboxylic group of acetic acid occurs at a reasonable rate, the rate for intramolecular C-isotope exchange (17) is expected to be slow and likely to be the rate limiting step.

Another potentially important mechanism for the C-isotope system is exchange of C between CH<sub>4</sub> and organic matter. Sackett (1993) reported on a pyrolysis experiment demonstrating that C-isotope exchange between kerogen and CH<sub>4</sub> at 700°C is completed in about 200 hours, corresponding to a half-exchange timescale of ~40h. Using the Arrhenius relationship, we can calculate that at 400°C this would be about half a year. The exchange rate in natural settings would depend upon the reactive surface area of organics and is therefore difficult to estimate, but we expect that this process could be relevant at Nisyros, where the basement rocks include cretaceous limestone (Marini et al., 1993), which could provide kerogen that would exchange C isotopes with methane.

A third mechanism of C-isotope exchange could be via partial chemical equilibration between CO<sub>2</sub> and CH<sub>4</sub> following reaction (1), such as:

$${}^{12}\text{CO}_2 + 4\text{H}_2 \rightleftharpoons {}^{12}\text{CH}_4 + 2\text{H}_2\text{O} \tag{19}$$

$$^{12}\text{CO}_2 + 4\text{H}_2 \rightleftharpoons ^{12}\text{CH}_4 + 2\text{H}_2\text{O}$$
 (19)  
 $^{13}\text{CO}_2 + 4\text{H}_2 \rightleftharpoons ^{13}\text{CH}_4 + 2\text{H}_2\text{O}$  (20)

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(e.g. Fiebig et al., 2004; Giggenbach, 1997). As outlined earlier, the rates for this reaction are poorly known. Early experiments by Berndt et al. (1996) suggested a reduction of 1% CO<sub>2</sub> during serpentinization of olivine at 300°C in a period of about two months, from which the forward rate could be calculated. However, more recent experiments using a more robust <sup>13</sup>C-labelling technique (McCollom, 2016) showed that rates are likely much slower, and that many previous experimental work invoking abiotic reduction of inorganic carbon compounds may have been affected by background concentrations of reduced carbon, potentially from the reacting minerals. The experiments of McCollom (2016) failed to produce any significant <sup>13</sup>C-labelled methane despite having a high thermodynamic drive for reaction (20) (i.e. high  $fH_2$ ), except for one experiment where a separate vapor phase was allowed to form.

In geothermal systems fH2 depends on a ladder of reactions involving hydrolysis and progressive oxidation of Fe<sup>2+</sup> to Fe<sup>3+</sup>, which in turn depend on fluid pH, on the chemical composition of the surrounding rocks (e.g. mafic vs felsic) and on the stability of secondary minerals formed (Stefánsson, 2017). In general, this provides much lower thermodynamic drive for CO<sub>2</sub> reduction than those expected in serpentinization environments. Nevertheless, we use the <sup>13</sup>C-labelled experiment of McCollom (2016) where some CH<sub>4</sub> was produced as an upper estimate for the rates of chemical equilibration between CO<sub>2</sub> and CH<sub>4</sub>. This experiment was

conducted at 300°C and found that 52 µmol/kg of <sup>13</sup>CH<sub>4</sub> was produced over 3885h, with a final  $\Sigma$ CO<sub>2</sub> of 11mmol/kg. From this we derive a pseudo-first-order rate constant  $k_c$  of 0.001 a<sup>-1</sup> (where  $k_c' = \ln\left(\frac{c_0}{c_t}\right)/t$ ), which is about 60 times slower than the rate calculated from the Berndt et al. (1996) experiments.

Giggenbach (1997) suggested that <sup>13</sup>C isotope exchange is slower than chemical exchange by a factor of ~400. We apply the same approach to derive the following pseudo-first-order rate constants of chemical  $(k_c')$  and isotopic exchange  $(k_i')$ :

$$\log k_c' = 4.77 - 4440/T \tag{21}$$

$$\log k_i' = 2.16 - 4440/T \tag{22}$$

where the second term is based on a previously determined activation energy for the Fischer-Tropsch process of ~85 kJ/mol (Huff and Satterfield, 1984; Zimmerman and Bukur, 1990).

These rates are substituted into equation (4b) to compute progressive equilibration of  $\alpha_{\text{CO2-CH4}}$  values. While the reaction order is likely different, for the sake of comparison with H<sub>2</sub>O- $CH_4$  exchange we make the simplifying assumption that  $k_c$  and  $k_i$  are the overall pseudo-first order rate constants and that the equation ln(1-F) = -k't (4b) is still valid. Progressive isotopic equilibration according to equations (22) and (4b) is shown in Fig. 7D and 8C using the same endmember as in the deuterium exchange calculations (Krafla well 32). From this it is clear that rates of <sup>13</sup>C exchange are expected to be much slower than those of D exchange at high temperature, and are reflected by unreasonably long timescales for any <sup>13</sup>C equilibration to occur in the Icelandic systems studied, and in Nisyros provided that chemical and isotopic disequilibrium was likely introduced by the 1873 eruption (Fig. 7D, 8C). Therefore, it appears that if any <sup>13</sup>C isotopic exchange occurred, such as implied by the apparent equilibrium  $\alpha_{CO2-CH4}$  values observed at Nisyros, Kerlingarfjöll and Hædarendi (Fig. 7D), then CO<sub>2</sub> reduction must be catalyzed by either sedimentary organic material or acetic acids as discussed earlier. Alternatively, the apparent <sup>13</sup>C isotopic equilibrium is coincidental in these systems and reflects the <sup>13</sup>C-enrichment trend associated with open-system breakdown of organic matter. In its advanced stage, the latter process could generate methane having an isotopic composition consistent with equilibrium-like  $\delta^{13}C_{CH4}$  values (Fiebig et al., 2019).

# 5.4. Process of CH<sub>4</sub> generation and equilibration

Overall, the kinetic end-member CH<sub>4</sub> isotopologue composition of Krafla-32 is consistent with thermogenic methane produced by thermal cracking of organic material at various possible temperatures (Fig. 8A, B). For example, CH<sub>4</sub> generated in the hydrothermal reservoir at ~250°C would preserve any kinetic fractionations associated with thermal cracking. However, the trend from low  $\delta D_{CH4}$  values towards equilibrium at Krafla is well reproduced by partial isotopologue equilibration (Fig. 7A, B), and this equilibration process appears to be associated with highenthalpy wells (Fig. 9). Such wells have a high vapor/liquid ratio, which could indicate the addition of superheated vapor to the subcritical reservoir (Elders et al., 2014). The input of superheated vapor is supported by the observation of a partially equilibrated acid well with high enthalpy (Fig. 9), which is thought to represent condensed superheated vapor (Heřmanská et al., 2019). Therefore, low  $\Delta^{13}$ CH<sub>3</sub>D values observed in Krafla could instead represent a unique window into the fluid processes occurring in the deep roots of geothermal systems.

Fluid flow in a hydrothermal reservoir is largely controlled by host rock permeability and by the thermal structure of the geothermal system, which is in turn determined by their magmatic history (Hayba and Ingebritsen, 1997; Fournier, 1999; Scott et al., 2016). Numerical simulations show that the extent of boiling zones is controlled by both magmatic emplacement depth and host rock permeability: for example, boiling zones are restricted to shallow levels ( $\leq 1$ km) in systems with high permeability ( $\geq 10^{-14}$  m²), whereas they may span the whole depth range from surface to intrusion in systems with intermediate permeability ( $\sim 10^{-15}$  m²), provided that the intrusion depth is < 3 km (Scott et al., 2016). These conditions are optimal for the development of supercritical geothermal resources, as observed in the Krafla geothermal system where the intrusion depth is just over 2 km (Scott et al., 2015). Therefore, the observation of kinetic CH<sub>4</sub> at Krafla and Námafjall with  $\Delta^{13}$ CH<sub>3</sub>D-based temperatures ( $438^{+55}_{-45}^{+55}$ °C) corresponding to the temperatures of superheated vapor sampled by the IDDP-1 well (Elders et al., 2014) could reflect the recent magmatic history of the geothermal system, shallow depth to magmatic bodies and appropriate host rock permeability (Fig. 10A).

Local variations in host-rock permeability, for example due to its temperature dependence near the magmatic intrusion (e.g. Hayba and Ingebritsen, 1997; Fournier, 1999) may control the extent of H<sub>2</sub>O-CH<sub>4</sub> isotopic exchange, since it would affect fluid velocities and thus, residence times. For example, Krafla well 32 (kinetic end-member CH<sub>4</sub>) could sample a zone where host-rock permeability is slightly higher, which would promote faster recharge of meteoric fluids with liquid enthalpy and quenching of kinetic CH<sub>4</sub> isotopologue signals. In contrast, highenthalpy wells may sample zones with lower host-rock permeability, which could provide more favorable conditions for the development of more extensive and longer-lived "pockets" of superheated vapor where CH<sub>4</sub> isotopologue equilibration proceeds over longer durations (Fig. 10A). Hence, the unique thermal structure of the Krafla geothermal system, where conditions leading to supercritical geothermal resource formation are optimized (Scott et al., 2015), could in large part be responsible for the unique CH<sub>4</sub> isotopologue signals observed there (Fig. 7B).

In contrast, the other systems studied here (Reykjanes, Kerlingarfjöll, Nisyros) have older magmatic histories (~100-1000 years) and deeper heat sources (~3-5 km) (e.g. Ármannsson, 2016; Bachmann et al., 2012; Bali et al., 2020; Caliro et al., 2005) where supercritical fluids may not be as important, or not present at all (Scott et al., 2016). This could in part explain their systematically higher  $\Delta^{13}$ CH<sub>3</sub>D values, whereas the degree of H<sub>2</sub>O-CH<sub>4</sub> isotopic equilibration will be determined by reservoir temperature and fluid residence times. Nisyros can be characterized as an equilibrium CH<sub>4</sub> isotopologue end-member, with all geothermometers pointing to reservoir temperatures of 350°C (e.g. Fig. 5C and 7). Host rock permeability in this system is characterized by two permeable zones separated by an aquiclude (Chiodini et al., 1993; Marini et al., 1993), and thus fluid residence times are longer, probably on the order of 100-1000 years. Figure 8 shows that this is largely sufficient for  $\Delta^{13}$ CH<sub>3</sub>D and  $\delta$ D<sub>CH4</sub> isotopologue equilibrium to be attained, but probably not for CO<sub>2</sub>-CH<sub>4</sub> isotopic equilibrium. A schematic representation is shown in Fig. 10B.

Organic matter in small concentrations brought from the surface may reach the roots of geothermal systems without breaking down significantly. As fluids approach the heat source thermogenic breakdown would become more effective. This is analogous to CH<sub>4</sub> generation during pyrolysis experiments conducted over short timescales, which is commonly optimized at temperatures above 400°C (Littke et al., 1995; Shuai et al., 2018a). The thermal cracking

generates methane with disequilibrium  $\delta D$  and  $\delta^{13}C$  signals, which may be exemplified by samples from the Krafla and Námafjall areas, where these signals are preserved as superheated vapors are rapidly quenched in the overlying reservoirs. The difference in isotopologue signals from other locations could reflect the absence of substantial supercritical resources and/or longer fluid residence times. Alternatively, thermogenic breakdown may occur in shallower regions of the hydrothermal reservoirs, with low  $\Delta^{13}CH_3D$  values mostly reflecting kinetic effects associated with breakdown of organic matter at hydrothermal reservoir temperatures. The lower reservoir temperatures of Icelandic systems (Table 4) would limit the extent of  $H_2O-CH_4$  isotopic exchange and thus favor the preservation of kinetic signals, while the higher temperatures at Nisyros could lead to isotopic and isotopologue equilibration. However, we consider this latter scenario less likely, since it requires the preservation of large  $\delta D_{CH4}$  kinetic signals with invariant  $\Delta^{13}CH_3D$  signals for Krafla and Námafjall samples.

Because isotopologue signals for CH<sub>4</sub> produced from CO<sub>2</sub> reduction in hydrothermal settings are not well characterized, we also cannot exclude the possibility of a contribution of abiotic methane. The low  $\Delta^{13}$ CH<sub>3</sub>D values raise the interesting possibility that methane could be produced abiotically in the supercritical reservoir at Krafla, at temperatures as high as 490°C. Such high temperatures and reactions in a superheated vapor phase could overcome the kinetic barriers associated with CO<sub>2</sub> reduction (McCollom, 2013; Reeves and Fiebig, 2020). It is also interesting to note that locations where CO<sub>2</sub> flux is particularly high, such as Nisyros (Bini et al., 2019) and Hædarendi (Ármannsson, 2016), also have higher concentrations of CH<sub>4</sub> (Table 1) in simultaneous apparent clumped and <sup>13</sup>C equilibrium, suggesting possible high-temperature <sup>13</sup>C isotopic exchange. Our model of CH<sub>4</sub> generation at Krafla and Námafjall (Fig. 10A) suggests a decoupled origin for methane and the bulk of the geothermal fluids. This is reminiscent of the interpretations of CH<sub>4</sub> isotopologues in seafloor vents, where most CH<sub>4</sub> is thought to be produced in fluid inclusions and picked up by actively circulating hydrothermal fluids. There again, low  $\Delta^{13}$ CH<sub>3</sub>D values suggest a deeper origin for CH<sub>4</sub> compared to the hydrothermal environment. In line with abiotic CH<sub>4</sub> production experiments (e.g. McCollom, 2016) and interpretations of the origin of abiotic CH<sub>4</sub> in seafloor hydrothermal vents, the formation of a superheated vapor phase in some geothermal systems could provide a favorable environment for abiotic CH<sub>4</sub> production.

Abiotic synthesis of hydrocarbons through the serpentinization of mantle rocks in marine hydrothermal environments is supported by a growing body of evidence (Proskurowski et al., 2008; Lang et al., 2010; Ménez et al., 2018; Wang et al., 2018; Klein et al., 2019). As mentioned above, the reducing conditions prevailing in these environments may be more conducive to CH<sub>4</sub> production compared to the more oxidizing conditions of subaerial geothermal systems. However, the mechanism may be similar in many regards, following reaction 1 and controlled by fH<sub>2</sub> which is determined by fluid-rock reaction progress and mineral assemblages (Stefánsson, 2017; Leong and Shock, 2020), possibly aided by the presence of a vapor phase (McCollom, 2016). The measured CH<sub>4</sub>/CO<sub>2</sub> ratios in marine systems tend to be higher than those of geothermal systems, which could reflect this difference in redox conditions. Moreover, the apparent clumped temperatures reported here are higher than those reported for marine systems (Fig. 8A), which could either reflect the higher kinetic barriers to overcome in more oxidizing systems, or longer residence times and thus lower temperature equilibration of magmatic fluids in the oceanic crust (or both).

# **Conclusions**

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Methane isotopologue distributions in steam vents and geothermal well fluids from Iceland and Nisyros are characterized by relatively low  $\Delta^{13}$ CH<sub>3</sub>D values (0.82 to 1.77‰), reflecting the generation of methane and subsequent equilibration at high temperatures generally in excess of 300°C. The lowest  $\Delta^{13}CH_3D$  values were measured at exploited geothermal wells in Krafla and Námafjall in Iceland. These low  $\Delta^{13}CH_3D$  values are associated with low  $\delta D_{CH4}$  and  $\delta^{13}C_{CH4}$  values, suggesting kinetic as opposed to equilibrium signals. These kinetic signals are consistent with methane generation by thermal cracking of organic matter, although abiotic generation of methane in superheated vapor at temperatures in excess of 400°C cannot be excluded. Partial re-equilibration of  $\delta D_{CH4}$  values with  $\delta D_{H2O}$  occurs and is best recorded in highenthalpy fluids that may have experienced supercritical conditions for longer times. In contrast, methane samples from natural steam vent fields, including the Kerlingarfjöll in Iceland and the volcanic caldera on Nisyros island, yielded higher Δ<sup>13</sup>CH<sub>3</sub>D values from 1.10 to 1.77‰. The average of  $\Delta^{13}$ CH<sub>3</sub>D values from Nisyros island (1.32±0.21‰), with corresponding apparent equilibrium temperature of  $355^{+50}_{-41}$ °C, are consistent with the measured reservoir temperatures of 350°C, suggesting isotopologue equilibration. From available experimental data, we estimated that isotopologue equilibration for  $\Delta^{13}CH_3D$  and  $\delta D$  between methane and water is achieved in tens of years at 350°C. Such timescale is consistent with the observed kinetic and equilibrium signals in exploited geothermal wells and matured natural geothermal systems. It is unclear whether any <sup>13</sup>CO<sub>2</sub>-<sup>13</sup>CH<sub>4</sub> exchange is occurring, since differences in the thermal maturity of degrading organic matter may also produce  $\delta^{13}C_{CH4}$  signatures overlapping those expected for isotopic equilibrium (Fig. 2), and since calculations suggest extremely slow rates for such exchange (Fig. 7D and 8C). More experimental work simulating the conditions of geothermal environments will be required to better characterize the possible reaction mechanisms of CO<sub>2</sub> reduction to CH<sub>4</sub> and evaluate the potential for abiotic methanogenesis.

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Table 1: Gas chemistry of geothermal well discharges (Iceland) and fumaroles (Iceland and Nisyros, Greece)

Sample Name	Location	H <sub>2</sub> O	CO <sub>2</sub>	H <sub>2</sub> S	H <sub>2</sub>	N <sub>2</sub>	Ar	02	CH <sub>4</sub>	СО	C <sub>1</sub> /C <sub>2+</sub>
Iceland wells											
15-AS-07	Krafla, w32	998030	1317	426	194	21.9	0.35	0.200	1.33	8.77E-03	256
15-AS-08	Krafla, w16	990059	8411	818	683	17.4	0.21	0.081	4.53	6.30E-03	248
15-AS-9	Námafjall, w13	998643	583	421	316	22.6	0.34	0.097	1.93	1.13E-02	742
15-AS-10	Námafjall, w9	997055	1107	837	896	63.0	1.05	0.256	26.9	1.26E-02	596
15-AS-17	Reykjanes, w15	994709	5069	156	11.8	48.7	0.90	0.107	1.25	3.03E-03	132
15-AS-15	Haedarendi, w7	464561	533293	140	15.7	1752.3	26.60	11.304	186	1.42E-02	2343
Iceland fumaroles	_										
15-AS-01*	Krysuvik	994240	5263	363	107	23.0	0.44	0.146	1.41	2.14E-03	316
15-AS-03	Kerlingarfjoll	995899	3239	406	413	37.7	0.54	0.200	2.71	1.70E-03	823
15-AS-05	Kerlingarfjoll	995991	2734	626	613	31.0	0.34	0.191	4.41	1.95E-04	870
15-AS-06	Kerlingarfjoll	996133	2693	554	592	21.8	0.25	0.103	4.59	3.54E-05	1016
15-AS-11	Námafjall	999129	635	132	20.8	60.0	1.32	1.983	20.0	n.m.	566
Nisyros fumaroles											
K7	Kaminakia	921217	69842	6098	871	404	5.6	7.4	1554	0.079	n.m.
K6	Kaminakia	946953	47480	3199	334	175	3.2	2.0	1854	0.053	n.m.
S4-lo	Stefanos	985744	10925	3059	109	53.3	0.78	0.90	108	0.242	n.m.
S4-hi	Stefanos	986871	10091	2828	90	32.0	0.45	0.29	86.8	0.079	n.m.
S15	Stefanos	986042	10431	3290	109	31.7	0.69	1.1	93.9	0.045	n.m.
A14	Phlegeton	982341	13728	3679	144	74.7	1.1	4.9	26.8	0.037	n.m.
A13	Phlegeton	977011	16788	3197	165	2763	40.5	4.9	30.5	0.025	n.m.
PP9S	Polybotes Mikros	989379	8779	1670	104	36.1	0.75	0.66	29.8	n.m.	n.m.
PP9N	Polybotes Mikros	986586	10824	2360	130	51.7	0.76	2.0	44.4	0.014	n.m.

All concentrations in µmol/mol. \*averaged composition of 15-AS-01 and 15-AS-02. C<sub>2+</sub> refer to C<sub>2</sub>-C<sub>6</sub> values of alkanes and alkenes. n.m.: not measured

Table 2: Isotopic composition of H and C in H<sub>2</sub>O-H<sub>2</sub>-CO<sub>2</sub>-CH<sub>4</sub> fluids

Sample Name	Location	δD <sub>H20</sub> (‰)	δD <sub>CH4</sub> (‰)	δ <sup>13</sup> С <sub>СН4</sub> (‰)	δ <sup>13</sup> Cco <sub>2</sub> (‰)	Δ <sup>13</sup> CH <sub>3</sub> D (‰)	95% CI (‰)	T <sub>Δ</sub> 13 <sub>CH3D</sub> (°C)				
Iceland wells												
15-AS-07	Krafla, w32	-82.4	-319.6	-39.6	-3.8	0.82	0.34	$490^{+163}_{-99}$				
15-AS-08	Krafla, w16	-83.1	-242.9	-38.7	-3.5	0.95	0.33	$447^{+129}_{-84}$				
15-AS-9	Namafjall, w13	-94.8	-262.4	-37.4	-3.7	1.22	0.28	$376^{+73}_{-55}$				
15-AS-10	Namafjall, w9	-88.5	-293.7	-35.8	-3.0	0.94	0.18	$450^{+61}_{-49}$				
15-AS-17	Reykjanes, w15	-27.8	-165.0	-31.5	-2.8	1.21	0.15	$379^{+37}_{-32}$				
15-AS-15	Haedarendi, w7	-86.1	-215.7	-27.0	-2.4	1.69	0.23	$290^{+38}_{-32}$				
Iceland fumaroles												
15-AS-1**	Krysuvik		-205.5	-36.4	-3.2	1.55	0.22	$312^{+41}_{-34}$				
15-AS-03*	Kerlingarfjoll†	-82	-201.0	-25.0	-3.2	1.31	0.44	$357^{+116}_{-76}$				
15-AS-03 (1)			-200.8	-24.6		1.09	0.40	$408^{+133}_{-85}$				
15-AS-03 (2)			-201.1	-25.4		1.53	0.20	$316^{+36}_{-31}$				
15-AS-05	Kerlingarfjoll†	-86	-201.4	-27.2	-3.1	1.34	0.34	$351^{+82}_{-60}$				
15-AS-06	Kerlingarfjoll†	-88	-201.0	-27.2	-3.0	1.32	0.14	$355^{+30}_{-27}$				
15-AS-11	Namafjall		-310.9	-35.2	-3.0	1.22	0.48	$376^{+145}_{-88}$	Measure	ments from	Gonzalez et al.	(2019)
Nisyros fumaroles									Δ <sup>13</sup> CH <sub>3</sub> D (‰)	$T_{\Delta^{13}CH_3D}$ (°C)	$\Delta^{12}CH_2D_2$ (%0)	$T_{\Delta^{12}CH_2}$ (°C)
K7	Kaminakia	-56.5	-125.8	-22.5	-1.5	1.16	0.21	$390^{+56}_{-45}$	$0.97 \pm 0.10$	447 +26 -23	$0.2 \pm 0.4$	860 +112
К6	Kaminakia	-9.1	-126.6	-22.4	-1.9	1.10	0.40	405+133				
S4-lo	Stefanos	-10.7	-125.1	-23.3	-1.4	1.24	0.20	$372^{+48}_{-40}$				
S4-hi	Stefanos	-11.3	-125.0	-23.3	-1.6	1.37	0.30	345+69	$1.14 \pm 0.30$	$400^{+90}_{-65}$	$-0.01 \pm 0.6$	>769
S15	Stefanos	-13.8	-125.1	-23.4	-0.9	1.20	0.24	381+63				
A14	Phlegeton	-4.5	-135.4	-23.4	-0.9	1.77	0.34	278 +56				
A13 PP9S	Phlegeton Polybotes	-4.8	-135.4	-23.3	-1.3	1.14	0.37	395 +113				
PP9N	Mikros Polybotes	-10.9	-124.4	-21.8	-0.8	1.40	0.19	339 +39 -33				
	Mikros	-10.4	-124.5	-22.0	-1.0	1.50	0.67	$321^{+166}_{-93}$				

\*duplicate sample:  $\delta 1^3 C_{CH4}$ - $\delta D_{CH4}$ - $\Delta 1^3 CH_3 D$  values represent averages of the two measurements shown below, and  $2\sigma$  values are compounded errors. \*\*composite sample (15-AS-01+15-AS-02).  $\dagger \delta D_{H2O}$  values of Kerlingarfjoll samples were not measured, but were approximated as a likely  $\delta D_{H2O}$  range for local meteoric water (Stefansson et al. 2017), consistent with a narrow range of  $\delta D$  values in Iceland's dilute hydrothermal systems (e.g. see Fig. 7A). Assigned values are italicized.

Table 3: Chemical and isotopic geothermometers discussed in this study

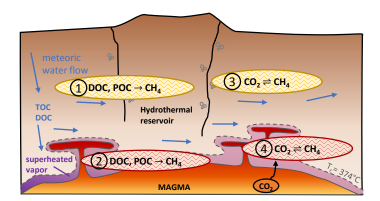
Reactions	Temperature dependence	Source				
$SiO_{2,s} + 2H_2O \rightleftharpoons H_4SiO_4^{\circ}$	$\log K_{qtz} - 34.188 + 197.47 * T^{-1} - 5.851 * 10^{-6} * T^{2} + 12.245 * \log T$					
T <sub>CO2</sub> (empirical)*	$T_{CO2} = 121.8 + 72.012Q - 11.068Q^2 + 4.724Q^3$					
T <sub>H2S</sub> (empirical)*	$T_{H2S} = 177.6 + 66.152Q + 4.811Q^2$					
T <sub>CO2/N2</sub> (empirical)*	$T_{CO2/N2} = 173.2 + 48.751Q + 7.599Q^2 + 1.739Q^3$	(b,c)				
$CH_4 + HDO \rightleftharpoons CH_3D + H_2O$	$\alpha_{H_2O-CH_4} = 1.0997 + 8.456 * \left(\frac{10^3}{T^2}\right) + 0.9611 * \left(\frac{10^9}{T^4}\right) - 27.82 * \left(\frac{10^{12}}{T^6}\right)$	(f)				
$^{12}\text{CH}_3\text{D} + ^{13}\text{CH}_4 \rightleftharpoons ^{13}\text{CH}_3\text{D} + ^{12}\text{CH}_4$	$\Delta^{13}\text{CH}_3\text{D}(T) = -0.11006 \left(\frac{10^3}{T}\right)^3 + 1.04151 \left(\frac{10^3}{T}\right)^2 - 0.55235 \left(\frac{10^3}{T}\right)$	(g)				
$^{12}\text{CO}_2 + ^{13}\text{CH}_4 \rightleftharpoons ^{13}\text{CO}_2 + ^{12}\text{CH}_4$	$1000 \ln \alpha_{CO_2 - CH_4}$ $= 0.16 + 11.754 * \left(\frac{10^6}{T^2}\right) - 2.3655 * \left(\frac{10^9}{T^3}\right) + 0.2054 * \left(\frac{10^{12}}{T^4}\right)$	(d,e)				

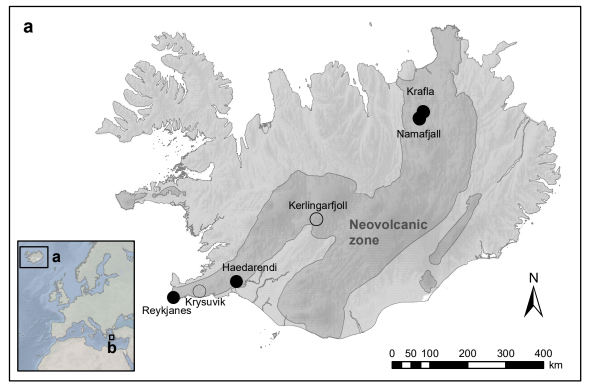
<sup>(</sup>a) Gunnarsson and Arnórsson (2000); (b) Arnórsson et al. (1998), updated from (c) Arnórsson and Gunnlaugsson (1985); (d) Horita (2001); (e) Richet et al. (1977); (f) Horibe and Craig (1995); Whitehill et al., (2017).\*Q is the logarithm of the concentration (in mmol/kg) or of the concentration ratio.

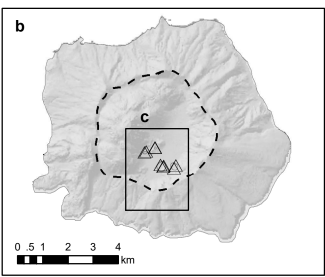
Table 4: Temperature-depth characteristics of boreholes, and inferred temperatures from gas geothermometry for fumaroles

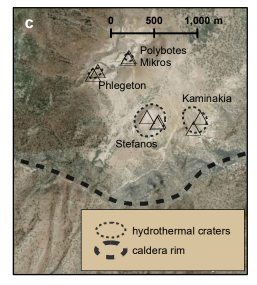
Sample Name	Location	Borehole depth (km)	T <sub>vent</sub>	Tres, measured	T <sub>qtz</sub> (°C)	T <sub>gas</sub> (°C)	T <sub>Δ</sub> 13 <sub>CH3D</sub> (°C)	Discharge enthalpy (kJ/kg)
Iceland wells								
15-AS-07	Krafla, w32	1.9		250	249	273	$490^{+163}_{-99}$	1253
15-AS-08	Krafla, w16	2.0		270	264	339	$447^{+129}_{-84}$	2767
15-AS-9	Námafjall, w13	2.2		290	290	254	$376^{+73}_{-55}$	1544
15-AS-10	Námafjall, w9	1.3		235	236	264	$450^{+61}_{-49}$	1083
15-AS-17	Reykjanes, w15	2.5		290	281	287	$379^{+37}_{-32}$	
15-AS-15	Haedarendi, w7	0.9		170		222*	$290^{+38}_{-32}$	
Iceland fumaroles								
15-AS-1	Krýsuvík					309	$312^{+41}_{-34}$	
15-AS-03	Kerlingarfjöll					289	$357^{+116}_{-76}$	
15-AS-05	Kerlingarfjöll					292	$351^{+82}_{-60}$	
15-AS-06	Kerlingarfjöll					295	$355^{+30}_{-27}$	
15-AS-11	Námafjall					232	$376^{+145}_{-88}$	
Nisyros								
K7	Kaminakia		97			356**	$390^{+56}_{-45}$	
К6	Kaminakia		97			355**	$405^{+133}_{-85}$	
S4-lo	Stefanos		102			345	$372^{+48}_{-40}$	
S4-hi	Stefanos		101			349	$345^{+69}_{-52}$	
S15	Stefanos		100			353	$381^{+63}_{-49}$	
A14	Phlegeton		102			349	$278^{+56}_{-44}$	
A13	Phlegeton		102			355†	$395^{+113}_{-76}$	
PP9S	Polybotes Mikros		100			336	339 +39 -33	
PP9N	Polybotes Mikros		99			341	$321^{+166}_{-93}$	

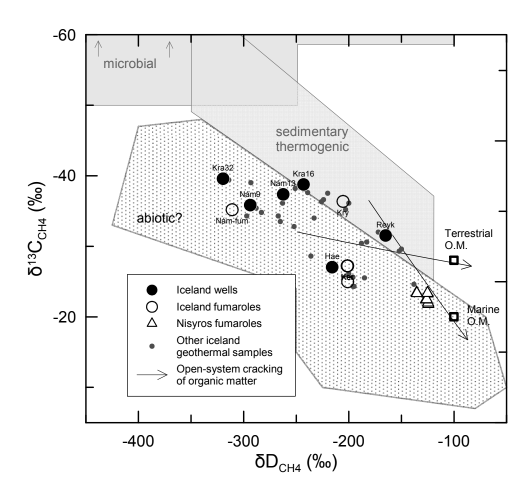
T<sub>gas</sub> calculated as the average of the CO<sub>2</sub>, H<sub>2</sub>S and CO<sub>2</sub>/N<sub>2</sub> geothermomers (Arnórsson et al., 1998), with following exceptions: \*T<sub>gas</sub> from H<sub>2</sub>S thermometer only. \*\*CO<sub>2</sub>-only geothermometer excluded. †CO<sub>2</sub>/N<sub>2</sub> geothermometer excluded

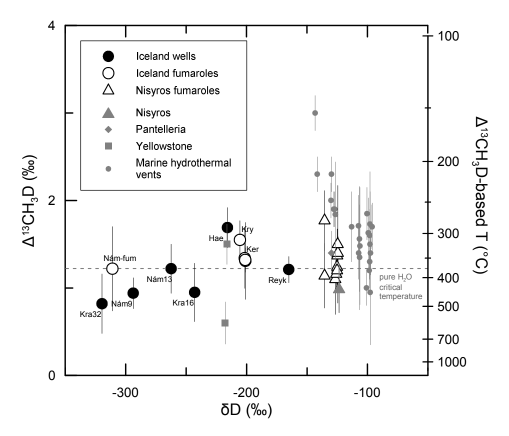


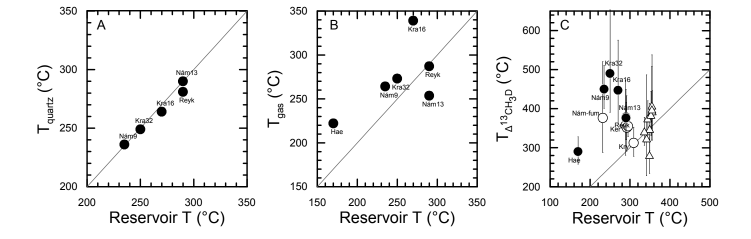


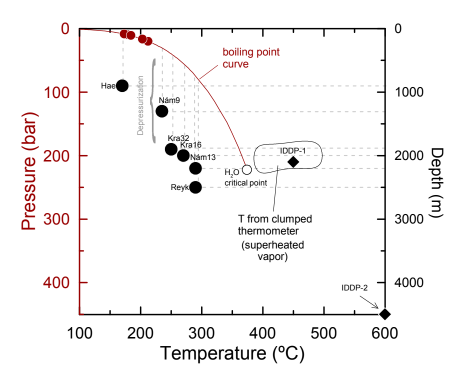


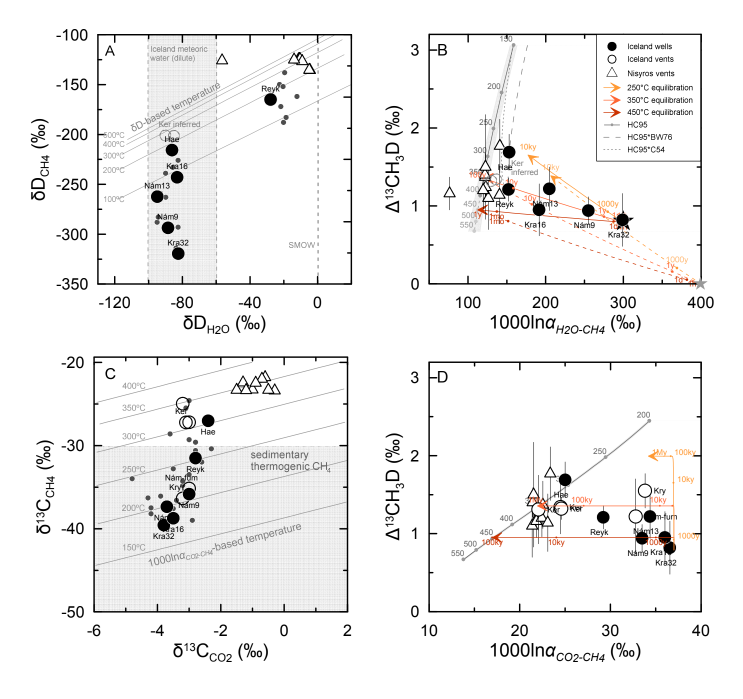


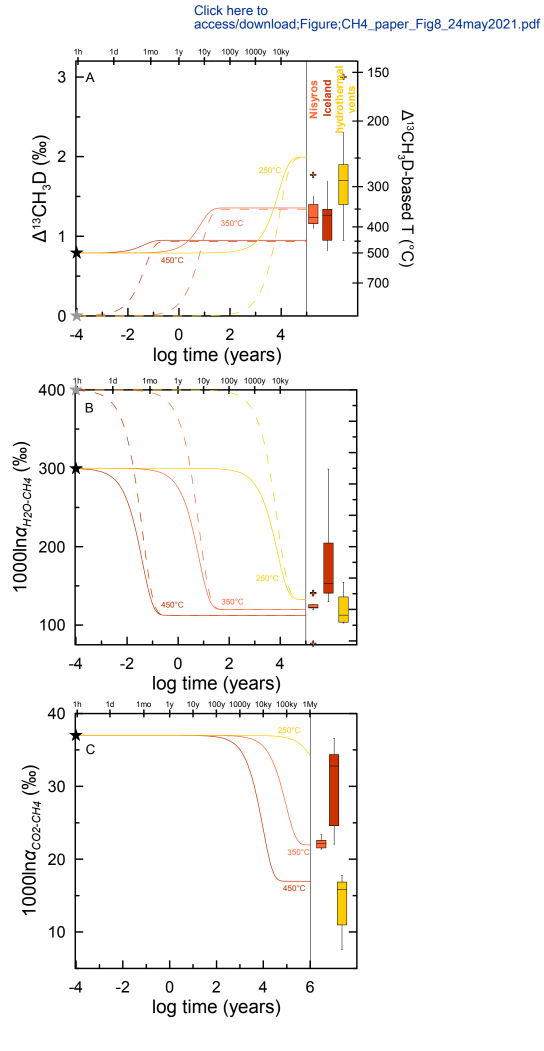


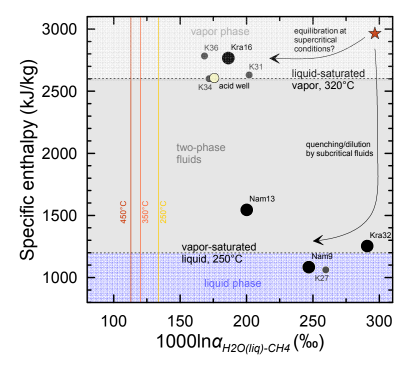


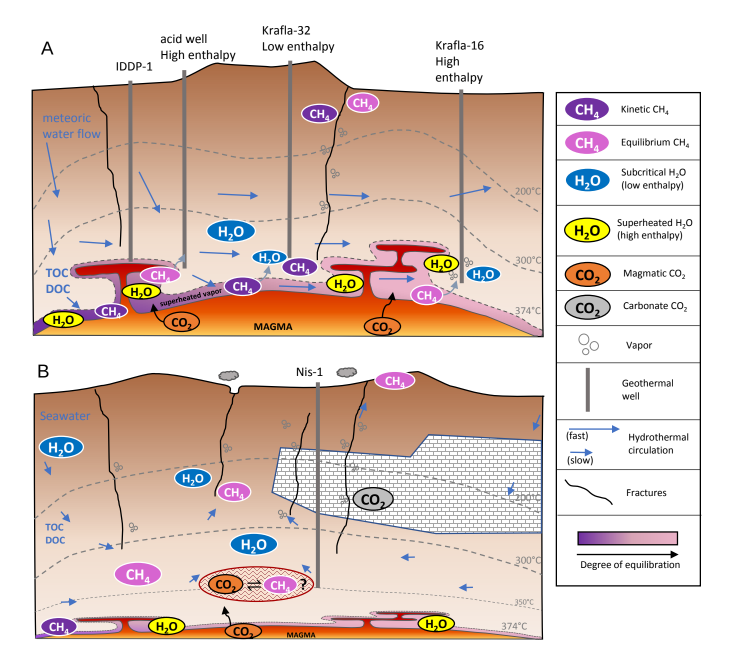












# **Figure captions**

Figure 1: Schematic representation of a terrestrial geothermal system and possible origins for low  $\Delta^{13}\text{CH}_3\text{D}$  signals observed to date. 1) methane is produced by the breakdown of organic matter (DOC, POC) contained in circulating groundwater or adsorbed on host rocks in the subcritical hydrothermal reservoir, and preserves low and kinetic  $\Delta^{13}\text{CH}_3\text{D}$  signals; 2) OM is transported close to the magmatic heat source and break down to CH<sub>4</sub> with equilibrium  $\Delta^{13}\text{CH}_3\text{D}$  signals; 3) CH<sub>4</sub> is produced abiotically in the subcritical reservoir via CO<sub>2</sub> reduction, controlled by the redox conditions of the system, preserving kinetic  $\Delta^{13}\text{CH}_3\text{D}$  signals; 4) CH<sub>4</sub> is produced abiotically in a superheated vapor phase via reduction of magmatic CO<sub>2</sub> and preserves equilibrium  $\Delta^{13}\text{CH}_3\text{D}$  signals.

**Figure 2**: Maps of Iceland (A) and Nisyros (B) showing sample locations and relevant geological features. Filled circles are geothermal well samples, and open circles and triangles are steam vents.

**Figure 3**:  $\delta^{13}C_{CH4}$  vs  $\delta D_{CH4}$  values of methane analyzed in this study and Fiebig et al. (2019) [1]. Also shown are typical ranges for microbial and sedimentary thermogenic methane, and the proposed range for abiotic methane from (Etiope and Sherwood Lollar, 2013). Grey dots ("Other Iceland samples") and the modeled trends for methane produced by thermal cracking from terrestrial and marine organic matter are from Fiebig et al. (2019). These trends do not consider potential secondary exchange of  $CH_4$  with water, which would flatten the corresponding pyrolysis slopes (Fiebig et al., 2019).

**Figure 4**:  $\Delta^{13}$ CH<sub>3</sub>D vs.  $\delta D_{CH4}$  plot showing the constancy of low  $\Delta^{13}$ CH<sub>3</sub>D values for all geothermal samples, including existing literature data from Nisyros, Pantelleria and Yellowstone (grey symbols, Douglas et al. (2017))\* and from marine hydrothermal vents (Douglas et al., 2017; Wang et al., 2018 and Labidi et al., 2020). Dashed line representing the critical temperature of pure water shown for reference. \*Data from Douglas et al. (2017) are  $\Delta_{18}$  values (*see* text in Results section).

**Figure 5:** (A) Comparison of measured hydrothermal reservoir temperatures with temperatures derived by the quartz geothermometer for well samples. 1:1 line shown in grey; (B) Comparison of temperatures derived by gas geothermometry with measured reservoir temperatures for well samples. (C) Apparent equilibrium  $\Delta^{13}$ CH<sub>3</sub>D-based temperatures compared with measured reservoir temperatures (wells) and calculated reservoir temperatures from gas geothermometry (vents). Symbols as in Figs. 1-3.

**Figure 6:** Pressure-temperature diagram showing the measured reservoir temperatures and borehole depths of sampled wells (black dots). The Pressure and Depth axes are approximately equivalent assuming hydrostatic pressure, hence samples follow the boiling point curve of water (red solid line) but plot left of the line, indicating depressurization of the hydrothermal reservoir upon intersection by wells. The sampling temperature and pressure (red circles) are along the

water vapor saturation curve. Maximum depths and temperatures of wells IDDP-1 (Krafla) and IDDP-2 (Reykjanes) also shown for reference; note that IDDP-1 stopped at 2.1 km when it encountered rhyolite magma (Elders et al., 2014).

**Figure 7**: Isotope and isotopologue systematics of geothermal samples, showing measured isotope values against those expected for equilibrium at different temperatures. (A)  $\delta D_{CH4}$  vs  $\delta D_{H2O}$ ; (B)  $\Delta^{13}CH_3D$  vs  $1000ln\alpha_{H2O-CH4}$ ; (C)  $\delta^{13}C_{CH4}$  vs  $\delta^{13}C_{CO2}$ ; (D)  $\Delta^{13}CH_3D$  vs  $1000ln\alpha_{CO2-CH4}$ ; dark grey dots in (A, C) are other Iceland samples—as in Fig. 3—reported in Fiebig et al. (2019). Colored arrows in (B, D) indicate the expected evolution of CH<sub>4</sub> isotopologues equilibration at different temperatures (shown in Fig. 7), starting from kinetic end-member Krafla-32 (black star). Dashed lines in Fig. 7B shows expected equilibration path from a more fractionated kinetic end-member (grey star), with  $\Delta^{13}CH_3D = 0\%$  and  $1000ln\alpha_{H2O-CH4} = 400\%$ . Equilibrium temperatures in (A) and (B) are from the calibration of Horibe and Craig (1995) (HC95); uncertainty (1σ) shown as grey envelope in (B). Dashed lines are shown when  $\alpha_{H2O(I)-H2}$  factors of Bardo and Wolfsberg (1976) (HC95\*BW76) and Cerrai (1954) (HC95\*C54) are used with  $\alpha_{CH4-H2}$  calibration of Horibe and Craig (1995). Equilibrium temperatures in (C) and (D) are from Horita (2001). Note the slower rates required for <sup>13</sup>C equilibration (D).

**Figure 8**: Evolution of CH<sub>4</sub> isotopologue signatures (A), deuterium fractionation (B) and <sup>13</sup>C fractionation (C) at various temperatures, assuming the starting CH<sub>4</sub> (black star) has the composition of kinetic end-member Krafla-32, or a more fractionated kinetic end-member (grey star and dashed lines) as in Fig. 7. Equilibration paths are compared to the ranges of measured values in Nisyros, Iceland and marine hydrothermal vents (box and whiskers plots).

**Figure 9:** Fluid enthalpy in Krafla and Námafjall well samples vs.  $1000 ln\alpha_{H2O-CH4}$ , showing that wells with high discharge enthalpy (Kra16, K31, K34, and K36) are closer to deuterium isotope equilibrium between  $H_2O$  and  $CH_4$ , shown for different temperatures as vertical colored lines. Black circles: samples measured in this study.  $1000 ln\alpha_{H2O-CH4}$  values of grey dots and yellow circle calculated with data from Fiebig et al. (2019). Acid well (yellow circle) shown for reference, as such wells are thought to be composed of condensed superheated vapor (Heřmanská et al., 2019). Specific enthalpy values are shown as horizontal dashed lines for liquid-saturated vapor at  $320^{\circ}C$ , based on fluid ascent paths for Krafla from Scott et al. (2015), and for vapor-saturated (boiling) liquid at  $250^{\circ}C$ , corresponding to the lower bound of measured reservoir temperatures.

**Figure 10:** Schematic diagrams illustrating the possible sources of CH<sub>4</sub> and processes of generation and equilibration. An idealized representation of the Krafla geothermal system is depicted in (A), where fast fluid circulation favors the preservation of kinetic isotopologue signals. The direction of fluid circulation (e.g. Pope et al. 2016) is consistent with longer equilibration timescales for CH<sub>4</sub> emanating from the area surrounding well 16. An idealized representation of the Nisyros is shown in (B), where fluid circulation is slower and thus equilibration timescales are longer. All CH<sub>4</sub> that reaches the surface has equilibrium isotopologue distributions. (A) and (B) can be viewed as early and late stages in the evolution of geothermal systems, respectively (e.g. Scott et al., 2016).