Geochemistry of a continental site of serpentinization, the Tablelands Ophiolite, Gros Morne National Park: A Mars analogue

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The presence of aqueously altered, olivine-rich rocks along with carbonate on Mars suggest that serpentinization may have occurred in the past and may be occurring presently in the subsurface, and possibly contributing methane (\(\text{CH}_4\)) to the martian atmosphere. Serpentinization, the hydration of olivine in ultramafic rocks, yields ultra-basic fluids (\(\text{pH} \geq 10\)) with unique chemistry (i.e. \(\text{Ca}^{2+}–\text{OH}^-\) waters) and hydrogen gas, which can support abiotic production of hydrocarbons (i.e. Fischer–Tropsch Type synthesis) and subsurface chemosynthetic metabolisms. Mars analogue sites of present-day serpentinization can be used to determine what geochemical measurements are required for determining the source methane at sites of serpentinization on Earth and possibly on Mars. The Tablelands Ophiolite is a continental site of present-day serpentinization and a Mars analogue due to the presence of altered olivine-rich ultramafic rocks with both carbonate and serpentine signatures. This study describes the geochemical indicators of present-day serpentinization and a Mars analogue due to the presence of altered olivine-rich ultramafic rocks at sites of serpentinization on Earth and possibly on Mars. The Tablelands Ophiolite is an important terrestrial analog for Mars.

1. Introduction

The detection of methane (\(\text{CH}_4\)) and its short lifetime in the martian atmosphere (Formisano et al., 2004; Krasnopolsky et al., 2004; Mumma et al., 2009), has led to much discussion regarding the source of methane on Mars. A possible source of methane has been attributed to serpentinization, a fluid–rock reaction common in ultramafic rocks that has been hypothesized to occur on Mars (Boston et al., 1992; Formisano et al., 2004; Atreya et al., 2007; Oze and Sharma, 2007). This reaction produces hydrogen (\(\text{H}_2\)) and the reducing conditions necessary for abiogenic hydrocarbon synthesis through the hydration of ultramafic rock, while also producing conditions for the production of methane through chemolithoautotrophic pathways. On Earth, serpentinization has been observed in ultramafic rocks on the sea floor at deep sea hydrothermal vents such as the Lost City hydrothermal field (Kelley et al., 2005), and on continents in ophiolites (Barnes et al., 1978; Abrajano et al., 1990; Blank et al., 2009; Etope et al., 2011). Ophiolites are sections of the ocean crust and upper mantle that have been obducted onto continental crust that can yield ground waters with unique chemistry that are ultra-basic (\(\text{pH} > 10\)), and/or \(\text{Ca}^{2+}–\text{OH}^-\) rich (Barnes et al., 1967). The Tablelands Ophiolite (also referred to in literature as the Bay of Islands Ophiolite) in Gros Morne National Park, Newfoundland, Canada is a continental site exhibiting present-day serpentinization. Olivine-rich ultramafic rocks, prerequisites for serpentinization, have been found on Mars (Hoefen et al., 2003; Hamilton and Christensen, 2005; Quesnel et al., 2009; Ehlmann et al., 2010), and also occur at the Tablelands massif, thus making the Tablelands Ophiolite an important terrestrial analog for Mars.

During serpentinization, ultramafic rocks composed of mostly olivine \([\text{Mg}_x\text{Fe}_{3-x}\text{SiO}_4]\) and pyroxene [orthopyroxene (\(\text{Mg},\text{Fe}\))–\(\text{SiO}_4\) and clinopyroxenes \(\text{Ca}^{2+}\text{(Mg,Fe)}\text{Si}_2\text{O}_6\)] undergo hydration producing serpentine (Eq. (1)). In a half reaction, iron hydroxide, which is produced via hydration of the iron-endmember of olivine, undergoes subsequent oxidation of \(\text{Fe}^{2+}\) and produces magnetite and hydrogen gas (Eq. (2)). The hydration of olivine and clinopyroxenes also releases \(\text{OH}^-\) ions resulting in highly alkaline fluids, which can emerge from fractures in the ultramafic rocks (Coleman and Keith,
Under basic conditions and in the presence of inorganic carbon, \( \text{Ca}^{2+} \) and \( \text{Mg}^{2+} \) form calcium and magnesium carbonate respectively (Eqs. (3) and (4)).

\[
\begin{align*}
3\text{Fe}_2\text{Si}_2\text{O}_5 + 5\text{Mg}_2\text{SiO}_4 + 9\text{H}_2\text{O} & \rightarrow 3\text{Mg}_3\text{Si}_2\text{O}_5(\text{OH})_4 + \text{Mg}(\text{OH})_2 + 2\text{Fe}(\text{OH})_3 \\
\text{fayalite} & \text{jadeite} & \text{serpentine} & \text{brucite} & \text{hydroxide}
\end{align*}
\]

(1)

\[
\begin{align*}
3\text{Fe}(\text{OH})_2 & \rightarrow \text{Fe}_2\text{O}_3 + 2\text{H}_2\text{O} + \text{H}_2 \\
\text{magnetite}
\end{align*}
\]

(2)

\[
\begin{align*}
\text{Ca}^{2+} + \text{CO}_2 & \rightarrow \text{CaCO}_3 \\
\text{calcium carbonate}
\end{align*}
\]

(3)

\[
\begin{align*}
\text{Mg}^{2+} + 2\text{HCO}_3 & \rightarrow \text{MgCO}_3\text{SiO}_4 + \text{CO}_2 + \text{H}_2\text{O} \\
\text{magnesium carbonate}
\end{align*}
\]

(4)

The production of hydrogen gas results in conditions capable of reducing dissolved inorganic carbon species and forming abiotic hydrocarbons (Sleep et al., 2004). In the following generalized reaction for abiotic hydrocarbon formation (i.e. the Tropsch synthesis) (Schulz, 1999), \( \text{H}_2 \) gas reduces inorganic reaction for abiogenic hydrocarbon formation (i.e. the Fischer carbons may also be present at the Tablelands. Therefore, thermogenic hydrocarbons could produce hydrocarbon gases. Therefore, thermogenic hydrocarbons may be present at the Tablelands.

Ultra-basic reducing springs have been identified in the Tablelands Ophiolite (Stevens, 1988). One of the main objectives of this study is to determine if these springs contain \( \text{H}_2 \), \( \text{CH}_4 \) and other hydrocarbons. Highly reducing conditions and the detection of \( \text{H}_2 \) in ground waters discharging from the Tablelands may provide conditions amenable for either abiotic and/or biogenic production of \( \text{CH}_4 \) and subsequently other higher molecular weight hydrocarbons. The process of serpentinization is of particular interest for the exploration of Mars and determining the origin of \( \text{CH}_4 \). Serpentinitization leaves mineralogical evidence of a previous physicochemical environment that would provide conditions for the production of methane and potentially other hydrocarbon gases to the martian atmosphere (Schulte et al., 2006). Near-infrared data from orbiting imaging spectrometer MRO-CRISM have shown the presence of Mg carbonate and serpentine signatures in several outcrops on Mars, suggesting that serpentinization occurred in the past and was active at least \( \gtrsim 3-7 \) Gyr ago (i.e. Noachian) (Ehlmann et al., 2008, 2010). Similarities in the martian crust to terrestrial ophio-

2. Material and methods

2.1. Site description

Highly reducing, ultra-basic springs discharging from serpentinitized peridotite (i.e. harzburgite with some lherzolites) (Suen et al., 1979) rocks were located in the Tablelands Ophiolite in Gros Morne National Park (N49°27′58.9″, W57°57′29.0″) (Fig. 1). The spring fluids are \( \text{Ca}^{2+}-\text{OH}^- \) type waters similar to those described by Barnes and O’Neil (Barnes et al., 1967). Inorganic carbon precipitates in these waters and is found as either carbonate sediment, travertine, or conglomerate cements that form where the springs discharge. Three sampling locations were identified on the northeastern face of the Tablelands and within Winter House Canyon: Wallace Brook (WB), Tablelands East (TLE), and Winter House Canyon (WHC) are shown in Fig. 2. TLE and WB were located on the slope of the Tablelands massif with travertine deposits and discharging \( \text{Ca}^{2+}-\text{OH}^- \) type waters. Located in WHC is a pool of ultra-basic water (labeled WHC2) that is approximately 40 cm
Fig. 1. Geologic map, modified from (Berger et al., 1992), showing the Tablelands massif and approximate sampling locations (star symbols): Wallace Brook (WB), Tablelands East (TLE), Winterhouse Creek (WHC1, 2a, b, c); and Winterhouse Brook (WHB). The Tablelands is located in Gros Morne National Park in Newfoundland, Eastern Canada.

Fig. 2. Images of springs and travertine deposits at sampling locations: (A) Winterhouse Creek 1 (2a, human for scale ~1.70 m); (B) Winterhouse Creek 2 (2b, human for scale ~1.70 m); (C) Tablelands East (2c, human for scale ~1.70 m); and (D) Wallace Brook (2d, spatula for scale ~22 cm).
deep and 126 cm wide, and exposed to the atmosphere at the surface. A survey of the bottom of WHC2 pool using handheld pH and Eh meters showed that there were two distinct locations where the pH was the highest and the Eh was the lowest indicating that there are two ultra-basic, reducing spring discharge points at the bottom of WHC2 (labeled WHC2a and WHC2b). A third sampling location (WHC2c) was selected for this study. At this location freshwater from overland flow trickles into the WHC2 pool. This site was of particular interest for studying the geochemical and biological interactions at a mixing site with large redox and nutrient gradients. The carbonate sediment at the bottom of the WHC2, and the travertine on the side of WHC2 spring were also sampled to determine the source of the inorganic carbon. A small and shallow oval shaped pool (approximately 2 cm deep, 5 cm wide) of ultra-basic spring water seeping from the travertine deposit was located approximately a meter from WHC2. It is identified as WHC1. WHC1 discharges at rate of 1 ml/min, and its surface is continuously exposed to the atmosphere; however, this pool was continuously flushed with ultra-basic groundwater and no freshwater inputs to WHC1 were identified. Winter House Brook (WHB) which flows along the bottom of Winter House Canyon was identified as the freshwater end member.

2.2. Aqueous geochemistry sampling and analysis

Daily field measurements of pH and redox potential ($Eh$) were obtained during sampling trips in July and September, 2009; June, August, and October, 2010; and June, 2011. Redox and pH values of water samples were measured using an ORPTestr 10 m (Eutech Instruments), pH paper and handheld pH meter (IQ Scientific Instruments GLP series IQ180G) respectively. Redox and pH values were measured during sampling to identify any changes in these parameters during or after water sampling, and after rainfall events. Samples were collected at least 48 h following a rainfall event to minimize dilution effects.

Samples for total inorganic carbon (TIC) and dissolved organic carbon (DOC) were collected for both concentration and stable carbon isotope ($\delta^{13}C$) values in pre-combusted 4 mL vials with mercuric chloride ($\text{HgCl}_2$) and phosphoric acid ($\text{H}_3\text{PO}_4$) respectively. Samples for DOC were filtered through a 0.7 µm pre-combusted glass microfiber filter. TIC and DOC concentrations and $\delta^{13}C$ values were determined using an Iol Analytical Aurora 1030 TOC Analyzer equipped with a reduction furnace, water trap, and packed GC column; coupled to a ThermoElectron DeltaV Plus Isotope Ratio Mass Spectrometer (IRMS) system via a Conflo III interface or a Finnigan MAT252 IRMS. The Aurora uses a wet chemical oxidation process to extract carbon as CO$_2$ gas using phosphoric acid for total inorganic carbon (TIC) and Na-per-sulfate for total dissolved organic carbon (DOC). Accuracy and reproducibility for concentration was ±1.5% RSD and ±0.5‰. Accuracy and reproducibility for $\delta^{13}C$ and $\delta^{18}O$ values are reported in delta notation relative to the Vienna Pee Dee Belemnite (PDB) reference standard.

Water samples were collected for hydrogen and oxygen isotope ($\delta^{18}O$, $\delta^{2}H$) in pre-combusted 4 mL vials with no headspace. Oxygen and hydrogen isotope measurements were analyzed at Isotope Tracer Technologies in Waterloo, Ontario on a Picarro Cavity Ring Down Spectroscopy Analyzer (Model L1102-i). Precision on multiple $\delta^{18}O$ and $\delta^{2}H$ measurements was ±0.1‰ and ±0.6‰, respectively. All results for oxygen and hydrogen are reported in delta notation relative to the Vienna Standard Mean Ocean Water (SMOW) reference standard.

Water samples were collected for major-ions chemistry and measured on an inductively coupled plasma mass spectrometer (ICP-MS) at Memorial University of Newfoundland using an ELAN DRCII ICP-MS. Samples for ICP-MS were filtered through a 0.45 µm filter and collected in 125 mL Trace-Clean bottles and immediately acidified with 5 N nitric acid. Certified reference materials were used for quality control. The detection limit on conservative ions $\text{Cl}^-$ was <0.01 mg/L and <1.7 mg/L for Br$^-$. The detection limit on $\text{Ca}^{2+}$ and $\text{Mg}^{2+}$ were <167 µg/L and <0.35 µg/L respectively. Total analytical error was ±10% RSD.

2.3. Carbonate mineralogy and isotope analysis

Samples of travertine and carbonate sediment from spring locations were collected for trace element geochemistry, carbon ($\delta^{13}C$) and oxygen ($\delta^{18}O$) isotopes. Samples were frozen upon collection, freeze dried, and ground to a fine powder with a mortar and pestle. X-ray diffraction was employed on powdered carbonates to determine measurements of major minerals present.$\delta^{13}C$ and $\delta^{18}O$ isotopes of carbonates were analyzed using a ThermoElectron Gas Bench II interfaced to an IRMS. Samples were measured into glass vials, placed in a heated block (50 °C), and flushed with helium prior to injection with phosphoric acid. The resultant gases were passed through Nafion dryers and a capillary column prior to entering into an ion source. External calcium carbonate standards CBM, NBS-19, and SPEX were used. Accuracy and reproducibility on $\delta^{18}O$ and $\delta^{13}C$ measurements was ±0.1‰ and ±0.5‰, and reported in delta notation relative to SMOW and PDB standards respectively.

2.3.1. Dissolved gas sampling

Dissolved gases including: $\text{H}_2$, $\text{CH}_4$, and other hydrocarbons including: ethane ($\text{C}_2\text{H}_6$, $\text{C}_2$), propane ($\text{C}_3\text{H}_8$, $\text{C}_3$), butane ($\text{C}_4\text{H}_{10}$, $\text{C}_4$), pentane ($\text{C}_5\text{H}_{12}$, $\text{C}_5$), and hexane ($\text{C}_6\text{H}_{14}$, $\text{C}_6$) were sampled using a modified syringe gas phase equilibration technique by McAuliffe (1971) and Rudd et al. (1974). Twenty milliliters of fluid was withdrawn with a 60 mL sterile syringe and shaken vigorously for 5 min with an equal volume of helium (He). This allowed for partitioning of the dissolved gas in the sample water into the gas phase. The entire gas phase of 2 syringes (40 mL) was injected into a 30 mL serum vial, prefixed with degassed water and sealed with blue butyl stoppers. The dissolved gases in He displaced the water in the serum vial. Samples were collected in triplicates for most sampling periods. The samples that were not sampled in triplicate were at least sampled in duplicate. Samples were fixed with 5 µL-saturated solution of HgCl$_2$ to ensure there was no microbial growth in bottles.

Dissolved gases were sampled for $\delta^{13}C$ analysis by collecting 50 mL of fluid using a 60 mL sterile syringe and injecting samples into a pre-evacuated 125 mL serum vial fixed with HgCl$_2$ and sealed with blue butyl stoppers. Samples were collected in triplicate.

2.3.2. Analysis of dissolved gases

Dissolved gases including $\text{H}_2$, and $\text{CH}_4$ were analyzed for concentration using a portable SRI 8610 Gas Chromatograph (GC) with a Thermal Conductivity Detector (TCD) and a Flame Ionization Detector (FID). The gases were separated using a Carboxen 1000 column packed in stainless steel (15 ft × 1/8 in., 2.1 mm film thickness) with a temperature program: 35 °C, hold 5 min, ramp at 20 °C/min to 225 °C, hold 15 min with a He carrier gas. Hydrocarbons were also analyzed by a GC equipped with a FID on a Porabond-Q column (30 m × 0.25 mm, 0.25 µm film thickness) with a temperature program of 35 °C hold 8 min, 10 °C/min to 150 °C, hold 8 min, ramp 5 °C/min to 210 °C, hold 10 min. Reproducibility on replicate samples was better than 5%.

Stable carbon isotope ratios of $\text{CH}_4$ were measured using an Agilent 6890 GC equipped with the Carboxen 1010 column coupled to a Finnigan MAT252 IRMS via combustion Conflo II Interface (GC-C-IRMS). Methane isotopes were determined using a 5:1 split ratio and a 100 °C isothermal temperature program. Low molecular weight hydrocarbons ($\text{C}_2$, $\text{C}_3$, $\text{C}_4$, $\text{C}_5$, $\text{C}_6$) were determined
3. Results and discussion

3.1. Origin of spring water

The $\delta^{18}O$ and $\delta^{13}C$ of the ultra-basic springs and freshwater including WHB, snowmelt, rainwater, and surface pond water collected from the top of the Tablelands massif plot close to the global meteoric water line (GMWL) (Fig. 3a), indicating that the ultra-basic spring fluids are meteoric in origin. This further suggests that the groundwater discharging from the peridotite rocks has been in recent contact with the atmosphere, excluding connate water or magmatic water sources.

The GMWL is representative of precipitation data collected from different locations globally as determined by (Craig, 1961) and therefore represents the global relationship between $\delta^{18}O$ and $\delta^{13}C$. The $\delta^{18}O$ and $\delta^{13}C$ data from the Tablelands do not plot directly on GMWL, which likely reflects the local conditions that control the $\delta^{18}O$ and $\delta^{13}C$ relationship. A linear line of regression representing the freshwater sources was added to Fig. 3a as a proxy of the local meteoric water line (LMWL) for the Tablelands. $\delta^{18}O$ and $\delta^{13}C$ of spring fluids plot closely to the LMWL indicating a similar $\delta^{18}O$ and $\delta^{13}C$ relationship as the freshwater. Therefore, changes in local meteoric conditions (e.g. temperature) likely govern the $\delta^{18}O$ and $\delta^{13}C$ of freshwater and ultra-basic springs respectively.

Fractionation of $\delta^{18}O$ and $\delta^{18}H$ is a function of temperature, with a greater fractionation associated with decreasing temperatures, resulting in a depletion of $\delta^{18}O$ and $\delta^{18}H$. Samples were collected over different seasons with varying temperatures. A seasonal trend related to temperature was observed in both spring fluids (WHC2b) and freshwater with an isotopic depletion during colder seasons and enrichment during warmer seasons (WHB) (Fig. 3b). The most isotopically depleted $\delta^{18}O$ and $\delta^{18}H$ were observed in early June (winter–spring season), when there was snow cover on the Tablelands and colder surface temperatures. The most enriched $\delta^{18}O$ and $\delta^{18}H$ were observed in September (summer–fall season) when there was no snow cover and warmer surface temperatures.

3.2. Mixing of freshwater and ultra-basic fluids at springs

In order to better understand changes in geochemical parameters in the springs, the contributions of freshwater and ultra-basic fluids were quantified at sites where mixing of the two end members was suspected. Fig. 4 represents a conservative 2-component mixing model between the freshwater end member (represented by WHB) and the ultra-basic reducing end member (represented by WHC1). In this model bromine (Br$^-$) and chlorine (Cl$^-$) are assumed to be conservative tracers of these two end members. The data from WHC2a, WHC2b, and WHC2c are well represented by the model and therefore suggest that the Cl$^-$ or Br$^-$ concentrations...
can be used to calculate the fraction of ultra-basic water ($f_{\text{UB}}$) contributing to the mixing sites (WHC2a, WHC2b, WHC2c). Sampling sites that are farther away from WHB and WHC (i.e. TLE and WB), are not as well described by the mixing model. This could be due to differences in the geochemistry of the end members for the TLE and WB springs. Likewise, TLE and WB, which are located on a steep slope of the Tablelands massif, received water inputs in the form of surface runoff, which could contribute to large differences observed in aqueous geochemistry seasonally, and account for larger standard deviations (Table 1).

Average measurements of $pH$, redox, and dissolved gas composition of fluids from the Tablelands can be seen in Table 1. Standard deviations reflect seasonal and annual variability in the geochemical parameters. The highest $pH$ measurements were found at WHC1 (12.2), WHC2a (12.3), and WHC2b (12.3). Redox measurements remained consistently low at WHC2a and WHC2b (~552 mV) at the bottom of the WHC pool where a distinct oxidized zone was observed. Higher redox values (>415 mV) were measured for the more oxidizing WHB freshwater end member. The redox value of the groundwater spring discharging at WHC1 increased during sampling. This occurred because of the slow recharge rate of the seep and exposure to the atmosphere, which allowed for atmospheric $O_2$ to partition into spring water and increase the redox value while sampling. Redox measurements where higher at mixing spring locations with oxygen-containing freshwater inputs (i.e. WHC2c, TLE, and WB). Differences in redox measurements reflect different amounts of mixing of freshwater and ultra-basic water at different times of sampling.

### Table 1

<table>
<thead>
<tr>
<th>A comparison of average ($\pm$std, 1σ) aqueous geochemical parameters of spring waters.</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Freshwater endmember</strong></td>
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<tr>
<td>WHB</td>
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<tr>
<td><strong>$pH$</strong></td>
</tr>
<tr>
<td><strong>$Eh$ (mv)</strong></td>
</tr>
<tr>
<td><strong>Cl$^-$ (mg/L)</strong></td>
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<tr>
<td><strong>Br$^-$ (mg/L)</strong></td>
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<tr>
<td><strong>Mg$^{2+}$ (mg/L)</strong></td>
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<td><strong>Ca$^{2+}$/Mg$^{2+}$</strong></td>
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<td><strong>TIC (mg/L)</strong></td>
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<td><strong>$\delta^{13}$C%</strong></td>
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<tr>
<td><strong>DOC (mg/L)</strong></td>
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<td><strong>DOC $\delta^{13}$C%</strong></td>
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$dL = \text{less than detection limit.}$

3.3. Geochronological evidence of present-day serpentinization

Fluids found discharging from serpentinized peridotites at the Tablelands Ophiolite are geochemically distinct from freshwater as evidenced by high $pH$, negative redox potentials, and the presence of $H_2$.

Dissolved $H_2$ concentrations were the highest (1.04 and 1.18 mg/L) at the most reducing sites with the least amount of mixing with freshwater and minimum exposure to the atmosphere (WHC2a and WHC2b). Lesser amounts of dissolved hydrogen (0.57–0.36 mg/L) were measured at sites with greater mixing between the ultra-basic reducing groundwater and the surface freshwater (WHC2c, TLE) and at slow discharging sites where the ultra-basic groundwater was directly exposed to the atmosphere for at least 10 min before sampling could occur (WHC1). During this time frame, dissolved hydrogen gas may have been released to the atmosphere. Dissolved hydrogen gas measurements were below detection at WB.

Likewise, major trace elements $Mg^{2+}$ and $Ca^{2+}$, derived from the hydration of olivine and clinopyroxenes respectively, are often found in fluids discharging from serpentinized ultramafic rocks giving evidence for active serpentinization. High $Ca^{2+}/Mg^{2+}$ are found in the ultra-basic springs at continental sites of serpentinization such as Cazadero in California (Barnes et al., 1967). Through serpentinization, soluble Mg$^{2+}$ concentrations become accommodated into serpentine and soluble Ca$^{2+}$ gets released into the water. Therefore, higher ratios of $Ca^{2+}$ to $Mg^{2+}$ are observed. Similarly, high $Ca^{2+}/Mg^{2+}$ ratios were found in the ultra-basic springs at the Tablelands (see Table 1).

In Fig. 5, $Ca^{2+}/Mg^{2+}$ ratios from spring fluids at WHC2a, WHC2b, and WHC2c at certain sampling periods are well described by the 2-component conservative mixing model defined from the freshwater and ultra-basic end members. The highest $Ca^{2+}/Mg^{2+}$ ratios are observed at most ultra-basic springs, similar to those observed in the Cazadero, CA ultra-basic springs. $Ca^{2+}/Mg^{2+}$ ratios were observed at WHC2b ($f_{\text{UB}} = 0.62$) and WHC2c ($f_{\text{UB}} = 0.32$) are not well described by the 2-component model. A possible third end member rich in $Mg^{2+}$—$HCO_3^-$ may be contributing to the spring fluids. As described by Barnes et al. (1967), $Mg^{2+}$—$HCO_3^-$ rich fluids can be found in localities of incomplete or inactive serpentinization resulting in shallow, moderately basic groundwater that is related to the weathering of serpentine and other magnesium-bearing minerals. High Mg$^{2+}$ concentrations (Table 1) were observed at WHC2b ($f_{\text{UB}} = 0.62$) and WHC2c ($f_{\text{UB}} = 0.32$) suggesting possible mixing of $Mg^{2+}$—$HCO_3^-$ fluids resulting in lower $Ca^{2+}/Mg^{2+}$ ratios. Similar concentrations of $Mg^{2+}$ and low $Ca^{2+}/Mg^{2+}$ ratios were observed at TLE and WB (Table 1), suggesting possible incomplete or inactive serpentinization at these localities.

**Fig. 5.** $Ca^{2+}/Mg^{2+}$ plotted versus the fraction of ultra basic water mixing with the conservative mixing line. $Ca^{2+}/Mg^{2+}$ ratios increase with greater fraction of ultra basic water, $Ca^{2+}/Mg^{2+}$ ratios from the Tablelands plot against $Ca^{2+}/Mg^{2+}$ ratios of spring ($Ca^{2+}/Mg^{2+} = 177$) and meteoric water ($Ca^{2+}/Mg^{2+} = 0.14$) from another continental site of present-day serpentinization in Cazadero, California (Barnes et al., 1967). Error bars of $Ca^{2+}/Mg^{2+}$ are ±10% and may appear smaller than the plotted symbol.
Carbonates have been observed to precipitate at the discharge points of ultra-basic springs emanating from altered ophiolites (Barnes and O'Neil, 1971). Carbonates in the form of travertine and sediment were found at ultra-basic springs discharging from serpentinitized peridotite in the Tablelands. X-ray Diffraction (XRD) analyses of travertine and carbonate sediments from the Tablelands suggest the carbonates found at all sites primarily consist of calcite (>90%) with a lesser amount of aragonite (<10%). Note that δ18O bars are smaller than the plotted symbol.

3.4. Carbonate mineralogy and deposition

Carbonates have been observed to precipitate at the discharge points of ultra-basic springs emanating from altered ophiolites (Barnes and O'Neil, 1971). Carbonates in the form of travertine and sediment were found at ultra-basic springs discharging from serpentinitized peridotite in the Tablelands. X-ray Diffraction (XRD) analyses of travertine and carbonate sediments from the Tablelands suggest the carbonates found at all sites primarily consist of calcite (>90%) with a lesser amount of aragonite (<10%). Note that δ18O bars are smaller than the plotted symbol.

The isotopic composition of carbonates can be used to understand the geochemical conditions under which the carbonates were deposited and can be used to interpret the inorganic carbon origin (Clark and Fritz, 1997). Stable carbon and oxygen isotope compositions (δ13C and δ18O) in travertine and carbonate sediment were determined (Fig. 6) and compared to the δ18O of water and δ13C of dissolved TIC (Table 2).

The large range in the δ13C of the carbonates in this study (Fig. 6) has also been observed in other freshwater carbonates at sites of present-day serpentinization (O'Neil and Barnes, 1971; Clark et al., 1992). Carbon isotope fractionation during chemical precipitation of freshwater carbonates is dependent on the rate of carbonate precipitation, temperature of precipitation, and the carbonate species present; therefore, a large range in the δ13C of carbonates can be observed (Turner, 1982). During this process TIC provides carbon for carbonate precipitation. TIC is the sum of dissolved carbon species: CO2 (aq), H2CO3, HCO3−, and CO3− 2. In the ultra-basic springs at the Tablelands (pH > 10), the carbonate ion, CO3− 2, is the predominant inorganic carbon species and can be assumed to make up most of the TIC composition. The enrichment factor between CO2 and CaCO3 is small (i.e. ~0.9‰ at 25 °C) (Mook, 1974). Therefore, it is not unexpected that the δ13C composition of carbonates reflects the δ13C composition of the TIC within roughly 1‰.

Since the δ13C of carbonates is largely influenced by the isotopic composition of the TIC, it is necessary to understand the origin of inorganic carbon that is contributing to the isotopic composition of the TIC pool. Different sources of inorganic carbon (i.e. CO2) for carbonates can exist and contribute to the δ13C of TIC thus providing a wide range of isotopic compositions. Relative contributions of CO2 (g) can be derived from the atmosphere (δ13C ~ −7‰) (Clark and Fritz, 1997); terrestrial organic matter (δ13C ≈ −24‰) (Deines, 1980); marine organic matter (δ13C ≥ −23‰); marine-carbonate (δ13C ~ 0‰) (Anderson and Arthur, 1983) and CO2 from microbial methane oxidation (δ13C ~ −54 to −30‰), assuming a starting δ13C methane of −60‰ (Whiticar, 1998). In addition to the isotopic composition of the initial CO2, the δ13C of carbonates is also influenced by the temperature-controlled fractionation between CaCO3 and CO2(g). The initial δ13C of CO2(g) contributing to the formation of carbonates was determined using the equilibrium calcite–CO2(g) isotope fractionation (εcalcite–CO2(g)) relationship developed by Bottinga (1969) and the measured temperatures of the sampled spring fluids (Table 3). The calculated starting δ13C of CO2(g) ranging from −36.2‰ to −27.1‰. The calculated δ13C of the initial CO2 is more depleted in 13C compared to atmospheric CO2. Calcite precipitation in equilibrium with CO2 (g) is enriched in carbon by −11‰, and aragonite by 12.8‰ (Rubinson, 1969). Additionally, the δ18O of carbonates can be influenced by the temperature-controlled fractionation between CaCO3 and CO2. Using the measured temperatures of the sampled spring fluids, equilibrium calcite–CO2(g) isotope fractionations were calculated using the εcalcite–CO2(g) relationship developed by Bottinga (1969) (Table 2). If the precipitation of the carbonates were simply due to atmospheric CO2 dissolving into the water and precipitating, the δ13C of the carbonates would be expected to be −19.0‰ to −17.4‰. Using calculated εcalcite–CO2(g), starting δ13C of CO2(g) was calculated (Table 2). Equilibrium fractionations ranging from 10.1 to 12.0 yielding starting δ13C of CO2(g) from −36.2‰ to −27.1‰ suggesting, in addition to atmospheric CO2 dissolution, there may be oxidation of terrestrial organic matter and/or methane contributing to the carbonates associated with the ultra-basic springs.

### Table 2

<table>
<thead>
<tr>
<th>Site</th>
<th>Measured δ13C(TIC) (‰)</th>
<th>Measured δ13CCaCO3 (‰)</th>
<th>Measured δ18O(TIC) (‰)</th>
<th>Measured δ18O(H2O) (‰)</th>
<th>Tm (°C)</th>
<th>Calculated εcalcite–CO2(g)</th>
<th>Calculated δ13CCO2(g) (‰)</th>
<th>Calculated δ18OCaCO3 (‰)</th>
</tr>
</thead>
<tbody>
<tr>
<td>WHC1</td>
<td>−30.6 to −28.2</td>
<td>−25.8 to −19.6</td>
<td>10.9−14.8</td>
<td>−9.5 (±0.1)</td>
<td>21.3−24.3</td>
<td>10.1−10.4</td>
<td>−36.2 to −35.9</td>
<td>19.0−19.7</td>
</tr>
<tr>
<td>WHC2a</td>
<td>−19.8 to −13.0</td>
<td>−16.7 to −11.6</td>
<td>16.1−22.2</td>
<td>−11.0 (±0.2)</td>
<td>10.5−15.5</td>
<td>11.1−11.7</td>
<td>−28.4 to −27.8</td>
<td>19.5−20.7</td>
</tr>
<tr>
<td>WHC2b</td>
<td>−19.2 to −12.5</td>
<td>−15.2 to −12.6</td>
<td>17.9−21.0</td>
<td>−10.8 (±0.3)</td>
<td>10.4−16.2</td>
<td>11.0−11.7</td>
<td>−26.9 to −26.2</td>
<td>19.5−20.9</td>
</tr>
<tr>
<td>WHC2c</td>
<td>−13.3 to −16.5</td>
<td>−15.4 to −8.8</td>
<td>15.7−21.9</td>
<td>−10.5 (±0.5)</td>
<td>11.3−17.2</td>
<td>10.9−11.6</td>
<td>−27.0 to −26.3</td>
<td>19.6−21.0</td>
</tr>
<tr>
<td>WB</td>
<td>−20.7 to −16.5</td>
<td>−16.7 to −9.9</td>
<td>14.6−19.2</td>
<td>−11.6 (±0.1)</td>
<td>10.0−19.9</td>
<td>10.6−11.8</td>
<td>−28.5 to −27.3</td>
<td>17.9−20.2</td>
</tr>
<tr>
<td>TLE</td>
<td>−15.5 to −9.9</td>
<td>−15.2 to −6.2</td>
<td>14.7−19.8</td>
<td>−12.3 (±0.1)</td>
<td>8.3−9.1</td>
<td>11.9−12.0</td>
<td>−27.2 to −27.1</td>
<td>19.7−19.9</td>
</tr>
</tbody>
</table>

Tm = Temperature measured, δ13C reported versus PDB and δ18O reported versus SMOW.

- Calculated equilibrium carbon enrichment between calcite and CO2(g) using fractionation equation: 1000ln x = 1.435 × 106T2 − 6.13 (Bottinga, 1969).
- Calculated equilibrium carbon–oxygen isotope composition using fractionation equation: 1000ln x = 2.78 × 106T2 − 2.89 (O'Neil et al., 1969; corrected in Friedman and O'Neil (1977)).
$\delta^{18}$O of carbonates are influenced by the temperature-controlled fractionation between CaCO$_3$ and water, and the $\delta^{18}$O of the water. The observed $\delta^{18}$O$_{CaCO_3}$ for travertine and carbonate sediment ranged from 10.9‰ to 21.9‰ (Fig. 6). Using the measured isotopic compositions and temperatures of the sampled spring fluids, and calcite-water oxygen isotope fractionations (O’Neil et al., 1969), corrected in Friedman and O’Neil (1977), theoretical calcite $\delta^{18}$O$_{CaCO_3}$ VSMOW values were calculated (Table 2). This isotopic fractionation assumes that the carbonates were precipitated in equilibrium with the water. The calculated equilibrium $\delta^{18}$O$_{CaCO_3}$ values ranged from 17.9‰ to 21.0‰. The $\delta^{18}$O$_{CaCO_3}$ data collected from WHC2b and WB fall within the calculated equilibrium $\delta^{18}$O range based on the water isotopes, and temperature at the time of sampling. Data from TLE, WHC2a, and WHC2c only partially fall within the calculated equilibrium $\delta^{18}$O range, where as data from WHC1 does not fall within the calculated equilibrium range. For the most part, $\delta^{18}$O$_{CaCO_3}$ that are not well described by the equilibrium $\delta^{18}$O range are more depleted in 18O. A similar isotopic depletion in 18O has been measured in travertine deposits and carbonate sediments at ultra-basic reducing springs at continental sites such as Cazadero, CA and the Oman ophiolite (O’Neil and Barnes, 1971; Clark et al., 1992). O’Neil and Barnes (1971) attributed this depletion in 18O to the rapid precipitation of CO$_2$ in an alkaline solution where by two-thirds of the oxygen in the solid carbonate comes from the CO$_2$ and therefore isotopically depleted oxygen of CaCO$_3$ would have been expected.

3.5. Source of CH$_4$ and low molecular weight hydrocarbons in spring fluids

Dissolved hydrocarbon gases including: methane, ethane, propane, n-butane, n-pentane, and n-hexane were measured in ultra-basic reducing springs. Highest hydrocarbon concentrations were at the most reducing sites WHC2a and WHC2b (Table 3). Lesser amounts of dissolved hydrocarbon gases were measured at sites with greater mixing between the ultra-basic reducing groundwater and the surface freshwater (WHC2c) and at slow discharge rates (WHC1). In the latter case hydrocarbons may have been lost to the atmosphere before they could be sampled. In order to obtain a fresh spring water sample, all water was removed from WHC1 and the small depression in the travertine was allowed to fill again before the water was sampled. It was noted that the $E_a$ became more positive while the pool was filling up, suggesting that the geochemistry of the water was changing very quickly in the presence of air. Therefore, hydrocarbons, inorganic carbon, and dissolved organic matter may have been partially or completely lost before sampling. At sites with the greatest amount of mixing, dissolved hydrocarbon gases were below detection (TLE and WB).

As a first approach, a Bernard plot (Fig. 7) was used to determine the primary source of CH$_4$ in the ultra-basic reducing springs. The Bernard plot has been used previously to discriminate between microbial and thermogenic methane production (Bernard et al., 1977). Bernard et al. (1977) found that on a plot of $\delta^{13}$C$_{CH4}$ versus CH$_4$/C$_2+$ (where C$_2+$ is the sum of the concentrations of C$_2$, C$_3$, and C$_4$) that microbially produced methane has high CH$_4$/C$_2+$ ratios (>400) and very negative $\delta^{13}$C values (<−50‰) relative to thermogenic methane with lower CH$_4$/C$_2+$ (<100) and more positive $\delta^{13}$C values (>−50‰). Dissolved methane from the ultra-basic springs at the Tablelands plot in the thermogenic field suggesting that the primary source of methane is not microbial in these springs. This observation is consistent with initial microbiological studies of the Tablelands spring fluids, which have found a lack of significant contribution from biological methanogenesis (Brazelton et al., 2012). All methanogens are members of the Archaea, and attempts to detect archaeal 16S ribosomal RNA genes in Tablelands fluids via the polymerase chain reaction have failed. In contrast, bacterial 16S ribosomal RNA genes are readily detected (Brazelton et al., 2012). Furthermore, metagenomic sequences predicted to represent methanogens comprise only 0.2% of the full metagenomic dataset from WHC2b (Brazelton et al., 2012); data are available at http://metagenomics.anl.gov under dataset “WHC2B sff”). These predictions are based on automated sequence similarity algorithms, and none of the predicted sequences actually encode proteins directly involved in the methanogenesis pathway. In short, the available microbiological data indicate that if methanogens are present in WHC2b at all, they are extremely rare.

The measured methane plots within a similar isotopic range (Fig. 7). However, one data point collected from WHC2c in August 2010 had an enriched value ($\delta^{13}$C$_{CH4} = −15.9‰$, where C$_2+$ was below detection limit. This enriched CH$_4$ may be evidence of methane oxidation occurring at WHC2c where there is O$_2$ present. Assuming a carbon isotopic signature of oxidized methane similar to that reported in June 2010 (−25.6‰), and uptake of the isotopically light carbon for metabolism, the left over carbon pool would be enriched in 13C as observed at WHC2c from August 2010. Likewise, the TIC produced via oxidation will be depleted relative to

![Fig. 7. Modified Bernard plot of CH$_4$/C$_2+$ versus 13C of methane from ultra-basic springs. Two distinct fields for microbially and thermogenically produced methane are plotted. Data points for hydrocarbon gases collected at the Tablelands fall within the thermogenic field.](image-url)

Table 3: Gaseous composition of spring waters.

<table>
<thead>
<tr>
<th></th>
<th>TLE</th>
<th>WHC2a</th>
<th>WHC2b</th>
<th>WHC2a</th>
<th>WHC2b</th>
</tr>
</thead>
<tbody>
<tr>
<td>H$_2$ (mg/L)</td>
<td>0.13–0.36</td>
<td>0.07–0.57</td>
<td>0.47–1.04</td>
<td>0.47–1.18</td>
<td>0.06</td>
</tr>
<tr>
<td>CH$_4$ (mg/L)</td>
<td>&lt;d.l.</td>
<td>0.03–0.05</td>
<td>0.04–0.32</td>
<td>0.16–0.38</td>
<td>0.03–0.06</td>
</tr>
<tr>
<td>C$_2$H$_6$ (mg/L)</td>
<td>&lt;d.l.</td>
<td>0.004–0.03</td>
<td>0.01–0.03</td>
<td>0.02–0.04</td>
<td>0.00–0.01</td>
</tr>
<tr>
<td>C$_3$H$_8$ (mg/L)</td>
<td>&lt;d.l.</td>
<td>0.01–0.03</td>
<td>0.01–0.04</td>
<td>0.03–0.05</td>
<td>0.00–0.01</td>
</tr>
<tr>
<td>nC$<em>4$H$</em>{10}$ (mg/L)</td>
<td>&lt;d.l.</td>
<td>0.01</td>
<td>0.01–0.02</td>
<td>0.01–0.02</td>
<td>0.004</td>
</tr>
<tr>
<td>nC$<em>6$H$</em>{14}$ (mg/L)</td>
<td>&lt;d.l.</td>
<td>0.005–0.01</td>
<td>0.01–0.02</td>
<td>0.02–0.04</td>
<td>0.00–0.01</td>
</tr>
<tr>
<td>CH$_4$/C$_2+$</td>
<td>NA</td>
<td>0.68–3.65</td>
<td>2.78–3.78</td>
<td>2.62–3.14</td>
<td>3.08–3.72</td>
</tr>
<tr>
<td>$\delta^{13}$C$_{CH4}$</td>
<td>&lt;d.l.</td>
<td>−15.9 to −26.3</td>
<td>−26.4 to −28.0</td>
<td>−26.4 to −27.7</td>
<td>−27.2 to −28.5</td>
</tr>
</tbody>
</table>

<d.l. = Less than detection limit, NA = not analyzed.
the carbon source, as was observed with a depleted $^{13}C_{\text{DIC}}$ ($-17.3\%$) sampled in August 2010. A decrease in CH$_4$ concentration and increase in TIC concentration was also observed in August 2010 (0.01 $\text{mg/L}$, 2.16 $\text{mg/L}$) as compared to June (0.03 $\text{mg/L}$, 1.73 $\text{mg/L}$) suggesting possible CH$_4$ uptake and CO$_2$ production from methane oxidation. During microbial oxidation the TIC would increase, and become depleted in $^{13}C_{\text{DIC}}$.

The methane measured may be thermogenic in origin. Thermogenic methane could be produced from the thermal alteration of the sedimentary organic matter underlying the Tablelands Ophiolite, and then migrated to the surface through cracks and fissures in the altered peridotite. Sedimentary organic matter most likely exists in the siliciclastic marine sandstones from the Blow-me-Down Brook formation, siliciclastic marine melange from the Croily Cove formation, and carbonate limestone from McKenzie’s formation (see geologic map, Fig. 1). Therefore, a thermogenic origin of the methane must be considered; however, an abiogenic origin must also be considered.

Although an abiogenic field is not found on a Bernard plot, experimental and field studies have shown the CH$_4$/C$_2+$, for abiogenic hydrocarbons (0.02–30) can overlap with the lower range of CH$_3$/C$_2+$, ratios for thermogenic hydrocarbons, especially if mixing with thermogenic hydrocarbons (McCollom and Seewald, 2006; Taran et al., 2007, and references there in; McCollom et al., 2010). CH$_4$/C$_2+$, ratios from the Tablelands springs (0.68–3.78) fall within this range. Therefore, the Bernard plot cannot discriminate between thermogenic and abiogenic sources of CH$_4$ within this range.

To distinguish between thermogenic and abiogenic origins of methane and higher molecular weight hydrocarbons, patterns of $\delta^{13}C$ and $\delta^D$ values of C$_1$–C$_6$ alkanes were considered. Thermogenic hydrocarbons have been shown to have a $^{13}C$ enrichment with increasing carbon number. This isotopic trend is attributed to kinetic isotopic effects where alkyl groups cleave from the source organic matter. The weaker $^{12}C$–$^{12}C$ bonds will break at a faster rate than the heavier $^{12}C$–$^{13}C$ bond (DesMarais et al., 1981). As a result of breaking the weaker $^{12}C$–$^{12}C$ bond, residual alkanes will be more enriched in the $^{13}C$ with increasing molecular mass. In contrast, it has been suggested that ethane can form abiogenically via a polymerization reaction demonstrating an isotopic depletion in $^{13}C$ and enrichment in D with respect to methane (Sherwood Lollar et al., 2002). In this polymerization reaction, the lighter isotopes $^{12}C$–$^{12}C$ will bond at a faster rate than the heavier and lighter isotopes $^{12}C$–$^{13}C$, resulting in a depletion of $^{13}C$ of ethane compared to that of methane. In this study, the $\delta^{13}C$ of alkanes from WHC2a and WHC2b follow a general isotopic depletion of C$_2$–C$_6$ relative to CH$_4$ with increasing carbon number and a clear $^{13}C$ depletion between methane and ethane similar to other putative abiogenic hydrocarbons (Fig. 8). However, it must be noted that this $^{13}C$ depletion between methane and ethane is not consistently observed for abiogenic hydrocarbons (Taran et al., 2007).

Sherwood Lollar et al. (2008) proposed that a saw-tooth pattern, in this case a $^{13}C$ depletion between methane and ethane followed by a $^{13}C$ enrichment between ethane and propane, can be modeled assuming rapid abiogenic polymerized chain growth with negligible carbon isotope fractionation in the formation of C$_2+$ compounds (Fig. 8). The Tablelands C$_2+$ data is not well described by such a model suggesting that the carbon isotopic patterns observed at the Tablelands may be due to fractionation effects associated with secondary alteration processes such as oxidation or diffusion, or mixing of hydrocarbons from thermogenic and/or abiogenic sources.

### 4. Implications and conclusions

Spatial variations in methane concentrations have been observed in the martian atmosphere suggesting the presence of localized source regions (Formisano et al., 2004; Mumma et al., 2009). This suggests that production and/or release of methane to the martian atmosphere may be related to the geology of the planet’s surface or subsurface. Possible source regions include Nili Fossae and NE Syrtis Major where aqueous altered olivine-rich rocks are exposed (Hamilton and Christensen, 2005; Mumma et al., 2009; Ehlmann et al., 2010). At Nili Fossae Mg-carbonates and Mg-rich serpentines have also been found (Ehlmann et al., 2010), and calcium pyroxenes have been detected at Syrtis Major (Rogers and Christensen, 2007). The presence of carbonate and serpentine signatures at localized methane source regions suggests that serpentinization may have occurred in the past on Mars. Given the short residence time of methane, detected methane at source regions would suggest either recent serpentinization where liquid water would need to be present, possibly in the subsurface, or Noachian and trapped as methane clathrates in the subsurface, and/or polar ice being released in the subsurface (Mumma et al., 2009).

The Tablelands Ophiolite is an active site of serpentinization and a Mars analogue for the presence of altered olivine-rich ultramafic rocks with both carbonate and serpentine signatures. Meteoric water reacting with ultramafic peridotites at the Tablelands forms geochemically distinct ultra-basic and reducing springs. These springs have characteristic Ca$^{2+}$ and Mg$^{2+}$ ion ratios. Calcium-rich carbonate sediment and travertine deposits have been found in and around the ultra-basic springs. Isotopic analyses of carbonates suggest that carbonate sediment and travertine derived carbon from CO$_2$ and are precipitated in non-equilibrium with the atmosphere and fluids. Dissolved gases including hydrogen, methane, and low molecular weight alkanes (C$_2$–C$_6$) have been measured in ultra-basic springs. The primary source of methane sampled from the springs is not microbial, but either thermogenic or abiogenic in origin, or a mixture of both. Additional work is needed to distinguish between the two possible sources of hydrocarbons. One possible approach is to examine the stable isotope composition of hydrogen of the hydrocarbon gases to determine if they are abiogenic or thermogenic in origin (Sherwood Lollar et al., 2008). With the current Mars Science Laboratory rover, goals...
to survey gases such as methane in the martian atmosphere, stable isotope and abundance analyses of gases can be applied to help provide information on the source of methane on Mars. Likewise, serpentinization at the Tablelands Ophiolite can be used as a mineralogical indicator to understand previous aqueous environment conditions and as a geochemical indicator to understand the conditions suitable for methane production on Earth and potentially on other ultramafic planetary bodies such as Mars.

Overall, key measurements on serpentinizing systems including aqueous geochemistry, carbonate mineralogy, and evidence of organic molecules can help in both interpreting data from MSL and planning future Mars exploration missions.

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