



GOPEN ACCESS

Citation: Schimmelmann A, Fernandez-Cortes A, Cuezva S, Streil T, Lennon JT (2018) Radiolysis *via* radioactivity is not responsible for rapid methane oxidation in subterranean air. PLoS ONE 13(11): e0206506. https://doi.org/10.1371/journal.pone.0206506

Editor: Elizabeth Grace Wilbanks, University of California Santa Barbara, UNITED STATES

Received: February 14, 2018
Accepted: October 15, 2018
Published: November 1, 2018

Copyright: © 2018 Schimmelmann et al. This is an open access article distributed under the terms of the Creative Commons Attribution License, which permits unrestricted use, distribution, and reproduction in any medium, provided the original author and source are credited.

Data Availability Statement: Our data are in Tables and in the Excel file (i.e. Supplemental Information). The methods have their own DOI under protocols.io.

Funding: This study is based upon work at Indiana University supported by the U.S. Department of Energy, Office of Science, Office of Basic Energy Sciences, Chemical Sciences, Geosciences, and Biosciences Division under Award Number DE-SC0006978 awarded to Arndt Schimmelmann. Work at the Royal Holloway University of London

RESEARCH ARTICLE

Radiolysis *via* radioactivity is not responsible for rapid methane oxidation in subterranean air

Arndt Schimmelmann $^{1^{\bullet}*}$, Angel Fernandez-Cortes $^{2,3^{\bullet}}$, Soledad Cuezva $^{2^{\bullet}}$, Thomas Streil 4† , Jay T. Lennon 5†

- 1 Department of Earth and Atmospheric Sciences, Indiana University, Bloomington, Indiana, United States of America, 2 Department of Biology and Geology, University of Almeria, Almeria, Spain, 3 Department of Earth Sciences, Royal Holloway, University of London, Egham, Surrey, United Kingdom, 4 SARAD GmbH, Dresden, Germany, 5 Department of Biology, Indiana University, Bloomington, Indiana, United States of America
- These authors contributed equally to this work.
- ‡ These authors also contributed equally to this work.
- * aschimme@indiana.edu

Abstract

Atmospheric methane is rapidly lost when it enters humid subterranean critical and vadose zones (e.g., air in soils and caves). Because methane is a source of carbon and energy, it can be consumed by methanotrophic methane-oxidizing bacteria. As an additional subterranean sink, it has been hypothesized that methane is oxidized by natural radioactivityinduced radiolysis that produces energetic ions and radicals, which then trigger abiotic oxidation and consumption of methane within a few hours. Using controlled laboratory experiments, we tested whether radiolysis could rapidly oxidize methane in sealed air with different relative humidities while being exposed to elevated levels of radiation (more than 535 kBg m⁻³) from radon isotopes ²²²Rn and ²²⁰Rn (i.e., thoron). We found no evidence that radiolysis contributed to methane oxidation. In contrast, we observed the rapid loss of methane when moist soil was added to the same apparatus in the absence of elevated radon abundance. Together, our findings are consistent with the view that methane oxidizing bacteria are responsible for the widespread observations of methane depletion in subterranean environments. Further studies are needed on the ability of microbes to consume trace amounts of methane in poorly ventilated caves, even though the trophic and energetic benefits become marginal at very low partial pressures of methane.

Introduction

Energetic radiation generates ions and radicals in fluids *via* radiolysis that can trigger subsequent chemical reactions [1], including the oxidation of organics. Radiolysis has likely affected the evolution of early microbial metabolisms and is crucial for powering the deep microbial biosphere [2, 3]. However, few studies have addressed the quantitative importance of radiolysis for contemporary fluxes in the atmosphere and the critical zone, especially in comparison to processes that compete with biologically mediated transformations.



(RHUL) was supported by the Spanish Ministry of Economy and Competitiveness projects CGL2016-78318-C2-1R and CGL2016-78318-C2-2R AEI/ FEDER/UE awarded to Angel Fernandez-Cortes. Jay T. Lennon was supported by the U.S. National Science Foundation (1442246) and the U.S. Army Research Office (W911NF-14-1- 0411). Jay T. Lennon further acknowledges support from the Indiana University Office of the Vice President of International Affairs. Angel Fernandez-Cortes' work was further supported by project SMACKS (IEF-Marie Curie Actions nº 624204, FP7/2007-2013). The funders had no role in study design, data collection and analysis, decision to publish, or preparation of the manuscript.

Competing interests: T.S. is employed by SARAD GmbH. This does not alter our adherence to PLOS ONE policies on sharing data and materials.

The concentration of methane ($\mathrm{CH_4}$) in the atmosphere has more than doubled since 1850 to ~1.85 ppmv (i.e., parts per million by volume) and now contributes ~15% of anthropogenic forcing of climate change [4]. The Intergovernmental Panel on Climate Change (IPCC) report [5] includes secondary greenhouse warming effects of $\mathrm{CH_4}$ and arrives at 1 W m⁻² for $\mathrm{CH_4}$ relative to 1.7 W m⁻² for $\mathrm{CO_2}$, making $\mathrm{CH_4}$ the second most important anthropogenic climate forcing agent. In the atmosphere, the removal of $\mathrm{CH_4}$ is due primarily to oxidation *via* photochemically generated tropospheric $\mathrm{OH_4}$ radicals ([6], and refs. therein). In spite of intense radiation in the atmosphere from sun and space, the residence time of atmospheric $\mathrm{CH_4}$ is ~12 years. The second largest sink for atmospheric $\mathrm{CH_4}$ is shallow subterranean environments containing aerated soils that are inhabited by $\mathrm{CH_4}$ oxidizing bacteria, MOB [4, 7]. MOB are also found in deeper aerated subterranean environments, such as caves in the vadose zone, although their contribution to global $\mathrm{CH_4}$ cycling has not been quantified or incorporated into earth system models [4].

A growing number of studies have reported that, throughout the world, concentrations of CH_4 are often depleted in the air of caves suggesting that subterranean environments may represent an overlooked sink for atmospheric CH_4 (e.g., [8–13]). Based on ventilation rates and CH_4 pools, it is estimated CH_4 is rapidly consumed in caves on time scales ranging from hours to days [14, 15]. Depletion of CH_4 in caves is often attributed to MOB. However, a study from Spanish caves proposed that rapid CH_4 oxidation may be attributed to non-biological processes *via* radiolysis and ionization of subterranean air by natural radioactivity that could lead to the oxidation of CH_4 at a sufficiently fast rate to account for appreciable consumption of CH_4 [10]. It has been proposed that α -radiation (e.g., from ²²²Rn) can radiolytically ionize, or generate radicals from, atmospheric components (e.g., H_2O) including CH_4 [16, 17]. The study by Haynes and Kebarle [16] determined that α -radiation has a slow effect on pure CH_4 and mixed hydrocarbon gas in the absence of air, making it difficult to extrapolate results to CH_4 in air in the presence of ions and radicals from heteromolecules.

Some studies, however, have raised questions about the relative importance of abiotic CH₄ oxidation based on theoretical considerations of kinetics, the inability of α -radiation from metallic uranium and radon to trigger fast oxidation of CH₄ [15, 18]. Laboratory and field experiments implicated MOB with the rapid decline in cave CH₄ concentrations [18], while isotopically uncharacterized radon was unable to remove CH₄ from air in an Australian cave [15]. Studies on radon typically focus on ²²²Rn because its longer half-life of 3.83 days facilitates quantification. No study has yet examined the radiolytic effect on CH₄ oxidation of the relatively more energetic decay of ²²⁰Rn (called thoron, with a half-life 55.6 s), particularly in the air close to cave walls and floors where ²²⁰Rn is relatively more abundant. Also, direct experiments linking the constraints of air humidity and natural radiation from specific radon isotopes to CH₄ oxidation in air are lacking. The current study fills these gaps with detailed independent experiments in two laboratories using energetically distinct radiation levels from isotopes of radon (²²²Rn and ²²⁰Rn) at different humidities and contrasting the results with CH₄-depletion by MOB.

Materials and methods

The authors of this study belong to two teams that had no knowledge of each others' experiments at Indiana University (IU) and Royal Holloway University of London (RHUL). After completion of all experiments, the two groups decided to jointly report their complementary results. Work at IU afforded superior analytical control on radon isotopes and could accurately measure higher dose rates, whereas the more gas-tight experimental setup at RHUL provided more straightforward evidence for the inability of natural radiation levels to rapidly oxidize atmospheric CH_4 at its natural atmospheric abundance.



Details of materials and methods are available from *protocols.io* under http://dx.doi.org/10.17504/protocols.io.s7aehie. We employed two separate, complementary experimental approaches at IU and RHUL. The following two sections offer brief overviews.

Apparatus at IU for active, time-resolved measurements of gas concentrations with circular flow

At IU, we constructed an experimental apparatus to assess the loss of CH_4 in an active (i.e., with pumping of air) and time-resolved manner with or without added radiation from radon isotopes and their progeny (Fig 1). The use of pumping qualifies this method as active and time-resolved in contrast to passive measurements of radon that integrate over time [19]. Approximately 6 L of air was recirculated in the sealed apparatus that included (i) a glass tube with optional thorium carbonate to generate 220 Rn (also called thoron), and (ii) a glass tube containing uranium ore to generate 222 Rn, with an overlying layer of coconut charcoal to limit the escape of co-produced, short-lived 220 Rn. Blank experiments without elevated radiation identified a reproducible loss of CH_4 (likely by diffusion through polymer tubing within the sealed analytical SARAD RTM 2200 instrument) that was subtracted from all other experiments at IU to arrive at net CH_4 losses that are due to other factors, such as radiolysis or microbial methanotrophy.

We conducted a number of experiments at IU to assess the importance of α -radiation intensity, relative humidity, and the presence or absence of soil on CH₄ dynamics. Moisture is critical for the emanation efficiency of radon isotopes from solid sources (i.e., the escape of noble gas radon atoms from the interior of minerals into H₂O-containing pore space *via* recoil subsequent to radioactive decay of parental nuclides; e.g., [20, 21]) and for stabilizing ions and radicals in air. Individual experiments differed in terms of their optional use of elevated humidity, thorium carbonate, and gas flowing through the tube containing uranium ore. The trapped ~6-L volume of air was initially spiked with CH₄ from natural gas to ~70 ppmv and with CO₂ to ~5,000 ppmv (except for experiments with soils) to distinguish it from room air and to increase the analytical precision during the time-series of measurements that lasted over a few days to weeks. Elevated CO₂ concentrations are typical for many cave environments [10].

Most experiments at IU discriminated between α -radiation from radon ^{222}Rn versus thoron ^{220}Rn . Whereas radon ^{222}Rn with a half-life of 3.83 days is relatively homogeneously distributed in cave air (also in our apparatus), the much shorter lived thoron ^{220}Rn with a half-life of only 55.6 s [22] cannot travel far from its parent nuclei residing in minerals [19], thus thoron's highest concentrations in cave air are near cave walls and the floor. The higher α -decay energy of ^{220}Rn (6.3 MeV) relative to ^{222}Rn (5.49 MeV) prompted us to design experiments for separate examinations of the ability of both radon isotopes to trigger the oxidation of CH₄. The more energetic α -decay of thoron ^{220}Rn should ionize air more efficiently than ^{222}Rn . Thoron was generated from thorium carbonate that was optionally loaded into a glass tube attached to the round-bottom flask. In other experiments, ^{222}Rn decay measuring up to 327 kBq m $^{-3}$ was produced *in-situ* in the glass apparatus by uranium ore chips (Fig 1). Escape of co-produced thoron from ore was reduced by using a layer of coconut charcoal in the upper part of the glass tube as a filter [23]. The resulting adsorption of ^{220}Rn on charcoal increased the residence time in the glass tube and let ^{220}Rn decay before it could enter the 5-L glass flask.

We quantified the concentrations of 222 Rn, 220 Rn radiation intensity was either measured *via* 220 Cn radiation



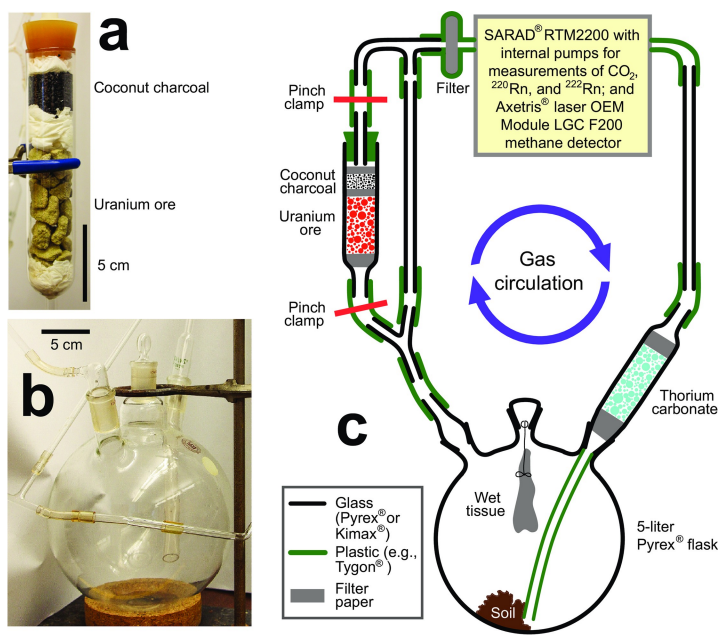


Fig 1. Experimental setup for time-resolved measurements of gas concentrations with circular flow at Indiana University. Approximately 6 L of air was recirculated in a sealed apparatus to assess the loss of CH_4 with or without added radiation from radon isotopes and their radioactive progeny. At the beginning of each experiment, the trapped air was slightly enriched in CH_4 (and CO_2 , except for experiments with soils), followed by hourly measurements of gas concentrations over a few days to weeks. Radon ²²²Rn was generated by uranium ore while charcoal retained ²²⁰Rn (a). The air intake of the 3-neck 5-L glass flask was directed to the bottom of the flask with a plastic insert to facilitate the mixing of air (b); thorium carbonate is not shown. Depicted components of the apparatus (c) are not drawn to scale.

https://doi.org/10.1371/journal.pone.0206506.g001

more powerful membrane pump, or values from flow rates \leq 0.2 L min⁻¹ with the diffusion pump were doubled to adjust for fast ²²⁰Rn decay (see S1 File for detailed control experiments and graphed data). Elevated relative humidity fosters the stabilization of ions in air *via* attachment to clusters of water molecules and may enhance the ability of ions to trigger oxidative degradation of CH₄ (discussed in [10]). Therefore, at IU we recorded humidity in the apparatus along with temperature, air pressure, flow, and battery voltage on an hourly basis. The



accuracy of data from the SARAD RTM 2200 was independently evaluated *via* direct comparison with a newly manufactured and factory-calibrated Thoron Scout instrument (SARAD GmbH, Dresden, Germany; details available in S1 File).

We conducted a number of experiments at IU to test for the effects of radiation and microbial activity on CH_4 dynamics in our experimental apparatus. Multi-day time-series of data were collected in closed-circuit air reflux mode (i) as duplicated blank experiments without added radon or thoron, (ii) with enhanced 220 Rn concentration in dry or moist air, (iii) with enhanced 222 Rn concentration in dry or moist air, and (iv) with jointly enhanced 220 Rn and 222 Rn concentrations in moist air to depict an extreme scenario where cave air had a highly elevated α -radiation level. Furthermore, (v) we tested for CH_4 oxidation after placing moist soils, which we assumed contained methanotrophic bacteria (MOB), into the 5-L glass flask, without elevated radioactivity. Certain impurities in industrially conditioned natural gas may act as MOB inhibitors, for example acetylene and carbon monoxide (p. 335 in [24]). As a precaution, the CH_4 spikes in experiments employing two different soils were derived from gas that was collected from a natural seepage of shale gas in New York State [25]. Natural shale gas is not known to contain acetylene or carbon monoxide.

Gas-tight terrarium experiments at RHUL

Experiments at RHUL at atmospheric CH_4 abundance used a gas-tight glass terrarium (i.e., an aquarium without water holding a volume of 13.45 L; Fig 2) with a hermetically sealing glass lid. Two air-tight gas ports allowed the withdrawal of 1-L air samples into Tedlar bags without changes in atmospheric pressure. Fragments of uraninite-bearing pitchblende served as a source of radioactivity. An AlphaLab Air Ion Counter with an integrated fan was placed in the terrarium to measure the abundance of ions in air in 30-s intervals. The α -radiation was quantified on 1-h intervals with a Canary Pro monitor (Airthings, Oslo, Norway) via α -spectrometry. Gas samples in Tedlar bags were analyzed for CH_4 mole fractions with a Picarro G1301 CRDS (Cavity Ring-Down Spectrometer, Picarro Inc., Santa Clara, California, USA).

The initial RHUL experiment #1 (Fig 2A) assessed the production of negative ions and the abundance of 222 Rn over \sim 6 h (i.e., stage 1) without either pitchblende or a beaker with water in the terrarium that had been flushed initially with laboratory air, and subsequently for \sim 15 h in the presence of pitchblende and a beaker with 130 mL of 38 °C warm water in the terrarium (stage 2).

The subsequent RHUL experiment #2 (Fig 2B) in the same terrarium included monitoring of the CH₄ mole fraction of laboratory air sealed in the terrarium where pitchblende and a beaker with 130 mL of water (initially at 38°C) had been placed to provide for elevated radioactivity and relative humidity. Elevated relative humidity was needed to simulate cave conditions. The AlphaLab Air Ion Counter failed to provide useful data due to static interference with Tedlar bags. The second experiment lasted for 76 h and 50 min and reached a ²²²Rn-based radiation level in excess of 50 kBq m⁻³ after 5 h. Approximate 1-L aliquots of air sampled from the terrarium were analytically compared with aliquots of exterior laboratory air on four occasions.

Results and discussion

Active time-series measurements with circular flow at IU

Our controlled experiments with and without 220 Rn and/or 222 Rn were designed to directly test whether or not radiation can oxidize CH₄ in cave air on ecologically relevant time scales (i.e., hours to days). We relied on comparisons of CH₄ inventories in experiments with (i) high radiation intensity from *in-situ* generated 220 Rn and/or 222 Rn with those from (ii) duplicate





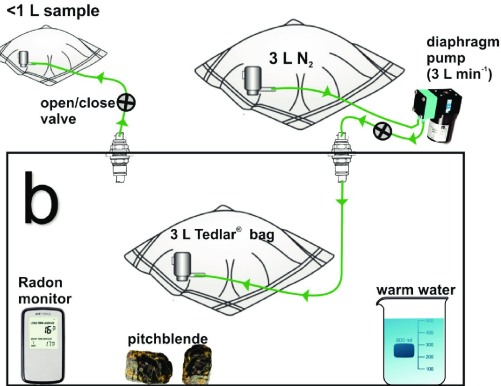


Fig 2. Gas-tight terrarium experiments at Royal Holloway University of London. (a) A hermetically sealed glass terrarium was filled with laboratory air containing atmospheric CH_4 . A radon monitor provided data on 222 Rn abundance, while an AlphaLab Air Ion Counter measured the concentration of negative ions. After \sim 6 h into the RHUL experiment #1, the placement of a beaker filled with deionized, warm water elevated the relative humidity to > 85%. At the same time, two fragments of pitchblende (containing uraninite as a radiation source) were placed into the terrarium



to generate 222 Rn. Tedlar bags in the terrarium are not shown in the photograph. (b) Diagram of the sampling procedure to collect <1-L aliquots of air from the terrarium in RHUL experiment #2. This experiment lasted for 76 h and 50 min and reached a 222 Rn-based radiation level in excess of 50 kBq m $^{-3}$ after 5 h.

https://doi.org/10.1371/journal.pone.0206506.g002

blank experiments with no artificially enhanced radiation to demonstrate the sensitivity of our setup to detect $\mathrm{CH_4}$ -losses. In addition, we conducted (iii) two experiments with moist soils in the absence of added radon isotopes to assess the potential for environmental microorganisms (i.e., MOB) to remove $\mathrm{CH_4}$ as has been demonstrated elsewhere by members of our research team [14, 18]. The comparisons among experiments covered a common range of $\mathrm{CH_4}$ concentration and thus only differed in the lengths of their time windows needed to lower the $\mathrm{CH_4}$ concentration from the upper to the lower threshold (i.e. yellow rectangle in Fig 3A). The 'common window' of $\mathrm{CH_4}$ decline for all 11 experiments maximized the data available for comparison.

Multiple trials in our experimental apparatus revealed that CH₄ dynamics were unaffected by radiation within the precision of measurements. Repeat blank experiments with dry (experiments #1 and #2) or moist air (experiments #3 and #4) without artificially elevated radon or thoron concentrations resulted in reproducible and systematic small losses of both CH₄ and CO₂ over time (Fig 3A and 3B; Table 1A). Although radon isotopes, CH₄ and CO₂ could not diffuse through glass and metal in our apparatus, the SARAD RTM 2200 and its Axetris laser OEM Module LGC F200 methane detector were internally and externally connected to glass and metal components with short segments of various types of clear polymer tubing (Fig 1) that resulted in slow losses via gas diffusion through polymers. The rate of diffusion across a layer of polymer is dependent on the difference in partial pressures between the interior and exterior air, and hence the rates of CH₄ and CO₂ losses via diffusion over time follow curves that asymptotically approach equilibria (Fig 3F). At a CH₄ concentration of ~59 ppmv in the apparatus (i.e., the midpoint of the common CH₄ range; Fig 3A) and outside air with ~1.85 ppmv, the mean CH₄ diffusive loss from air in the apparatus during blank experiments #1 to #4 consistently amounted to 0.39 ppmv h⁻¹ regardless of humidity and small variations in room temperature and air pressure (Table 1A; \$1 File). Such a loss of gas over time could theoretically result from a small internal leak in the system. However, the non-parallel pattern of CO₂ losses in blank experiments (Fig 3B) is inconsistent with a leak and instead argues for varying diffusivity of the polar molecule CO₂ through permeable material at different humidities. The observed degree of CH₄ loss from the system was unavoidable and had to be subtracted from the observed bulk CH4 losses in experiments with enhanced radiation and soils to arrive at any specific losses that are due to radioactivity or presumed microbial methanotrophy.

There was a comparable loss of CH_4 in recirculating air for all experiments without soil, regardless of the absence or presence of radiation from 220 Rn, 222 Rn, or both 220 Rn and 222 Rn, in dry or moist air (Fig 3A). The time needed to cross the 'common window' of CH_4 decline from 67.2 to 50.9 ppmv was not shorter when radiation from 220 Rn and/or 222 Rn was added (Table 1A, 1B). The slopes of lines representing CH_4 decline within the common window in Fig 3A were not higher for experiments with elevated radiation (mean \sim 0.38 ppmv h⁻¹) than for blank experiments without added radon isotopes (mean \sim 0.39 ppmv h⁻¹; Table 1A and 1B). The mean levels of added radiation from 220 Rn, and especially the cumulative radiation in experiment #9 from simultaneously added 220 Rn and 222 Rn, ranged between \sim 50 and 535 kBq m⁻³ after doubling of experimental 220 Rn values that were measured at \leq 0.2 L min⁻¹ (Table 1B) and thus always exceeded the radiation levels reported in cave air [26], including the air in all Spanish caves where abiotically driven CH_4 oxidation due to radiolysis has been



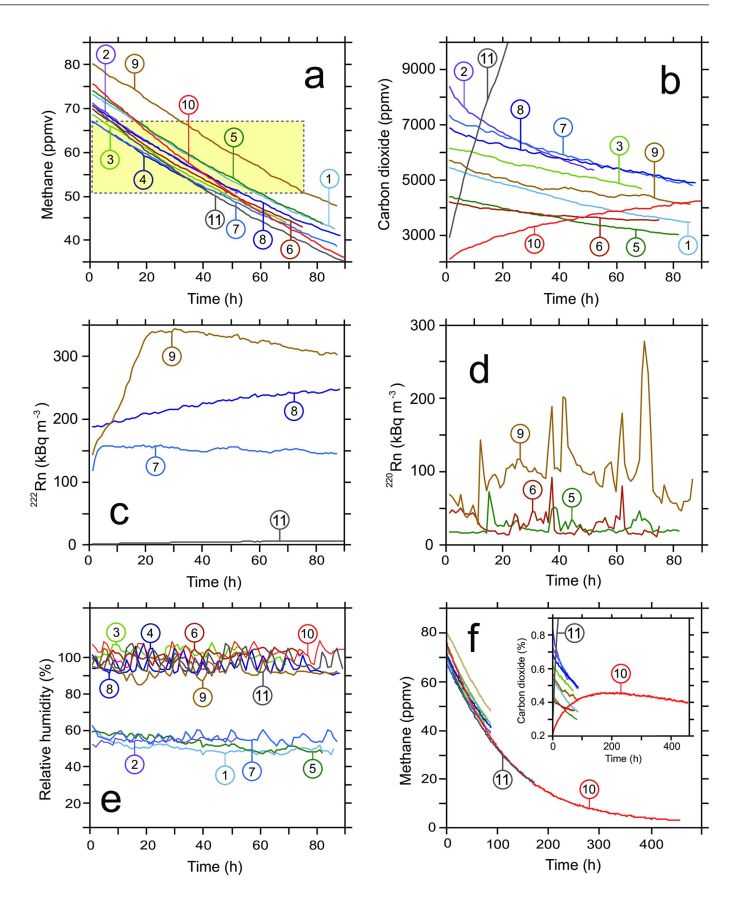




Fig 3. Time-series experiments at Indiana University. Results from time-series experiments to test for abiotic (radiation) and biotic (CH₄ oxidizing bacteria) factors on CH₄ dynamics. (a) The decline of CH₄ concentrations in experiments #1 to #9 (without soil) followed similar trajectories depending on original concentrations, despite major differences in radiation intensity (see the yellow rectangle that identifies a window of CH₄ concentrations that is common to all experiments). Declining CH₄ concentrations are independent of the intensity of α-radiation. Blank experiments #1 to #4 without elevated radiation identify a reproducible loss of CH₄ by diffusion that was subtracted from all other experiments to arrive at net losses that are due to other factors, such as radiolysis or microbial methanotrophy. (b) The decline of CO₂ concentrations in a range of experiments without soil followed similar patterns. In addition to loss due to diffusion through plastic, it was likely influenced by adsorption, solution in water, or possible chemical uptake. In addition, CO₂ was generated from moist soils in experiments #10 and #11. (c) Radon ²²²Rn and (d) thoron ²²⁰Rn concentrations partially depended on relative humidity; soil no. 2 in experiment #11 generated low levels of ²²²Rn over time presumably due to traces of uranium in minerals; ²²⁰Rn concentrations are original data from low flow rates at <0.2 L min⁻¹ when laminar flow conditions in the 5-L glass flask caused heterogeneity and occasional spikes. (e) Noise in relative humidity data partially derived from the automatic battery recharge cycle that influenced the internal temperature of the SARAD RTM 2200 and the algorithm to calculate humidity. (f) Experiments #10 and #11 with moist soils without added ²²²Rn or ²²⁰Rn resulted in a long-term exponential decline of CH₄ concentrations while CO₂ was generated biologically.

https://doi.org/10.1371/journal.pone.0206506.g003

reported [10]. For example, the average rate of CH_4 consumption in Spanish Altamira Cave air of -0.03 ppmv h⁻¹ occurred at a maximum ²²²Rn radiation level of ~6 kBq m⁻³, which is roughly one to three orders of magnitude less than the radiation in any of our experiments

Table 1. Overview on individual experiments performed at Indiana University to constrain the consumption of methane over time.

Experiment #	Overall duration (h)	Common window (h)	Mean values across common window of CH ₄ decrease from 67.2 to 50.9 ppmv							
			²²⁰ Rn (kBq m ⁻³)	222Rn (kBq m ⁻³)	CH ₄ loss (ppmv h ⁻¹)	CO ₂ loss (ppmv h ⁻¹)	Corrected temperature (°C)	Relative humidity (%)	Pressure (kPa)	
(A) Duplicated blank exper	riments without a	dded radon o	r thoron at	low or hig	h relative hun	iidity				
#1, no added radiation, dry	86	42.0	<0.01*	<0.01	0.39 ± 0.06	25.8 ± 14.1	23.2 ± 0.9	49.7 ± 1.5	97.2 ± 0.4	
#2, no added radiation, dry	52	42.7	<0.01*	<0.01	0.39 ± 0.07	47.6 ± 21.3	22.8 ± 0.4	54.7 ± 0.8	98.0 ± 0.1	
#3, no added radiation, wet	69	42.5	<0.2*	~0.4	0.39 ± 0.08	22.9 ± 23.0	23.4 ± 0.7	103 ± 2.7**	98.1 ± 0.2	
#4, no added radiation, wet	43	42.3	<0.3*	<0.2	0.39 ± 0.07	n.d.	25.4 ± 0.7	95.3 ± 4.1	99.1 ± 0.1	
(B) Experiments with enha	nced ²²⁰ Rn and/o	or ²²² Rn conce	ntrations a	t low or hiş	gh relative hui	nidity				
#5, ²²⁰ Rn added, dry	82	42.4	~50*	< 0.01	0.38 ± 0.07	18.4 ± 14.2	22.9 ± 0.6	53.3 ± 2.7	98.6 ± 0.2	
#6 , ²²⁰ Rn added, wet	75	41.8	143	< 0.01	0.39 ± 0.07	10.6 ± 15.0	22.8 ± 0.7	100.4 ± 2.1	98.9 ± 0.1	
#7, ²²² Rn added, dry	87	43.8	< 0.01*	153	0.37 ± 0.07	34.7 ± 35.3	24.3 ± 0.8	56.3 ± 2.2	98.8 ± 0.2	
#8 , ²²² Rn added, wet	88	44.5	< 0.04*	216	0.37 ± 0.09	22.0 ± 29.9	25.1 ± 0.9	95.5 ± 4.1	98.4 ± 0.3	
#9 , ²²² Rn and ²²⁰ Rn added, wet	87	43.5	208	327	0.37 ± 0.11	10.3 ± 27.9	25.8 ± 1.1	92.8 ± 3.4	98.4 ± 0.2	
(C) Experiments with mois	t soils without ad	ded radon or	thoron							
#10 , no added radiation, 45 g soil #1,	456	33.8	<0.03*	<0.2	0.49 ± 0.09	CO ₂ was generated	24.3 ± 0.8	100.6 ± 3.9	98.6 ± 0.2	
#11, no added radiation, 112 g soil #2	172	34.7	~0.1*	~4	0.47 ± 0.11	CO ₂ was generated	24.7 ± 0.9	97.9 ± 4.3	98.8 ± 0.1	

Analytical data are mean values with standard deviations for the time window when CH_4 concentrations declined from 67.2 to 50.9 ppmv in each experiment. This window represents the maximum range of methane concentrations that is common to all experiments. The length of time needed to deplete methane from the upper to the lower threshold across the common window (i.e., yellow rectangle in Fig 3A) was interpolated from hourly spaced data.

https://doi.org/10.1371/journal.pone.0206506.t001

^{*} 220 Rn radiation values were doubled when measured at flow rates \leq 0.2 L min⁻¹ to adjust for fast 220 Rn decay, instead of measurements without doubling of values in 10-min intervals (n \geq 10) at a flow rate of 1 L min⁻¹. See justification in S1 File.

^{**} High values of relative humidity are affected by analytical errors in excess of standard deviation.

n.d. = not determined.



with added radon isotopes (#5 through #9). Thus, in terms of radiation intensity, our experiments represent an extreme test of the radiolysis hypothesis. Only the air in shafts of underground uranium mines has been observed to reach even higher radiation levels of one million or more Bq m⁻³ [27].

The consistent pattern of CH₄ decline in our experiments without soils can be better appreciated in light of the observed CO₂ dynamics (Fig 3B). CO₂ is more polar than CH₄, can be more easily adsorbed on surfaces, and is more water-soluble and reactive than CH₄. Therefore, it is possible that changes in room temperature (21.1 to 27.5°C) and atmospheric pressure (96.7 to 99.3 kPa) may have affected adsorption and solubility of CO₂ during our experiments. Moreover, after one week of measurements with a wet paper tissue in the 4-L glass flask without soil, fungi had discolored the paper tissue and metabolically generated CO₂, thus partially stabilizing the CO₂ partial pressure (experiment #6, Fig 3B), apparently without affecting the CH₄ decline (Fig 3A). The paper tissue had been hung by a thread from the central glass stopcock to maximize surface area and to avoid any anoxic microenvironments that could facilitate biological methanogenesis (Fig 1). Subsequent experiments in moist air without soil replaced the wet paper tissue with added deionized water at the bottom of the 4-L glass flask. Experiments with soils initially generated CO₂ via microbial and fungal remineralization of soil organic matter, followed after several days by a decline due to diffusive loss of CO₂.

In the two experiments with moist soils, we documented a CH_4 loss of of ~0.09 ppmv h⁻¹ within the common window of CH_4 concentration decline (Fig 3A), as determined by subtracting the diffusive CH_4 loss in blank experiments from the bulk CH_4 loss in experiments #10 and #11 with soils (Table 1). It is well established that heterogeneously distributed methanotrophic biofilms in the subsurface [28] are capable of scavenging CH_4 from the atmosphere (e.g., [29, 30]). Soil gas can often reach ²²²Rn radiation levels of many thousand Bq m⁻³, depending on local geology [31, 32]. If radiolysis would indeed be able to trigger fast oxidative decay of CH_4 in soil gas, such an important CH_4 sink in dry soils without abundant methanotrophic activity would likely have been documented. Also, radiolysis would compete with methanotrophs in moist soils for CH_4 and would have been identified as a factor in soil CH_4 studies.

Experiments in gas-tight terrarium at RHUL

The first stage of experiment #1 at RHUL (Figs 2A and 4) established background conditions for the abundance of negative ions (~3800 ions cm⁻³) and the concentration of ²²²Rn (17 to 51 Bq m⁻³) in laboratory air at temperatures from 21.4 to 21.7°C and relative humidities from 26.6 to 29.0%. After the onset of stage 2, the placement of pitchblende and a beaker with 130 mL, 38°C warm water into the sealed terrarium strongly increased the abundance of negative ions in air (up to ~200,000 ions cm⁻³) and the concentration of ²²²Rn (~118 kBq m⁻³). The relative humidity exceeded 85%, and the air temperature intermittently rose by 5°C. The measurement uncertainty of the Canary Pro radon monitor increased with the ²²²Rn radiation level (Fig 4). However, the factory-documented uncertainty at the highest measured radiation level and the steadily increasing abundance of negative ions in air suggested that after a run time of ~17 h, the ²²²Rn-based radiation level exceeded 100 kBq m⁻³ (Fig 4; data shown in S1 File).

Experiment #2 at RHUL (Fig 2B) used the same sealed terrarium with pitchblende and high humidity to monitor and compare the CH_4 mole fractions in the air of both the terrarium and the outside laboratory air over ~77 h. The Canary Pro radon monitor in the terrarium indicated an increase in 222 Rn over time parallel to RHUL experiment #1. After 5 h into RHUL experiment #2, the 222 Rn-based radiation in the terrarium was consistently > 50 kBq m⁻³. Despite high levels of ionization and 222 Rn-based radiation in the terrarium, the CH_4 mole

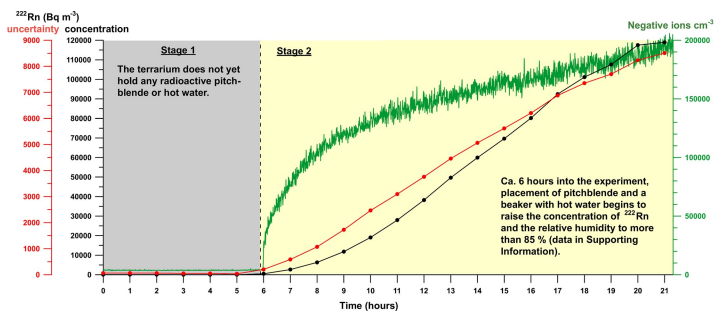


Fig 4. Experiment #1 at Royal Holloway University of London. The experiment proceeded in two stages where the first stage established background conditions in a hermetically closed terrarium at low relative humidity and in the absence of artificially elevated concentrations of 222 Rn. In the second stage, the addition of pitchblende and a beaker with hot water provided a source of 222 Rn and high humidity to simulate conditions characteristic of cave environments. Although the final 222 Rn concentration exceeded 100 kBq m⁻³ and was thus higher than in most caves, the elevated radiation and ionization of air in the terrarium was unable to lower the atmospheric abundance of CH₄ over 77 h in the subsequent RHUL experiment #2 (Table 2).

https://doi.org/10.1371/journal.pone.0206506.g004

fraction of 1.9941 \pm 0.0036 ppm in terrarium air after being sealed for ~77 h was indistinguishable from the starting value of 1.9971 \pm 0.0122 ppm within the uncertainty of measurements (Table 2).

Synopsis of combined results

The absence of any experimental evidence for accelerated loss of CH_4 in the presence of elevated radiation makes it highly unlikely that radiation from radon isotopes is important in nature where 220 Rn and 222 Rn concentrations are typically much lower. Our data indicate that natural radiation in cave air cannot be responsible for the rapid consumption of CH_4 in air on time-scales of days, even in caves with high relative humidity. The same conclusion had been reached from earlier laboratory experiments [18] and from observations in Australian cave air [15].

Table 2. Results of experiment #2 at RHUL to assess the consumption of CH_4 over time in the presence of elevated 222 Rn concentrations and high relative humidity.

Date	Time	Sample	CH ₄ (ppm)	CH ₄ st. dev. (ppm)	H ₂ 0 (vol. %)	
March 4, 2016	11:00	Laboratory air	1.9760	0.0006	0.85	
March 4, 2016	11:15	Initial terrarium air	1.9971	0.0122	1.75	
March 4, 2016	13:15	Laboratory air	1.9616	0.0005	0.81	
March 4, 2016	13:15	Terrarium air	1.9873	0.0066	1.10	
March 4, 2016	16:00	Laboratory air	1.9643	0.0006	0.85	
March 4, 2016	16:00	Terrarium air	1.9818	0.0054	1.64	
March 7, 2016	16:05	Laboratory air	1.9693	0.0007	0.77	
March 7, 2016	16:05	Final terrarium air	1.9941	0.0036	1.84	

https://doi.org/10.1371/journal.pone.0206506.t002



Many caves experience seasonally different degrees of venting and even reversals of air flow, which results in differences in air temperature and humidity and is difficult to simulate in laboratory experiments. Still, most cave environments at sufficient distances from cave entrances and vent holes are thermally buffered by surrounding rock and therefore do not express the relatively high diurnal and seasonal temperature and humidity variations as outside environments. Our experiments in laboratories were conducted at relatively constant room temperatures similar to many cave environments. Room temperatures in air-conditioned laboratory buildings are similar to actual temperatures in sub-tropical and tropical caves [14]. The use of water and moist soil in many of our experiments simulated the range of humidity in natural cave air. One possible caveat in terms of dissimilarity between our laboratory settings and actual caves may be the fact that our experiments allowed daylight to reach our experimental setups. However, the amount and timing of indirect light (no direct sunshine) was insufficient to let any photoautotrophs (algae) observably grow in our experiments. A necessary difference between air in our experiments at IU and actual cave air was the presence of traces of CH_4 in our experiments. Some CH₄ was needed to test for possible radiolytic destruction of CH₄. In contrast, most natural cave air is depleted in CH₄ relative to outside air. We conclude that the experimental conditions during experiments at IU and RHUL were reasonable approximations to simulate cave conditions. In the open atmosphere, solar radiation is mainly responsible for the generation of OH• radicals ([6], and refs. therein) that are the longest-lived potential radical reactant with CH₄ in air. Subterranean radiolysis by radioactivity involves far more energy than photochemical dissociation of molecules by solar radiation, hence the speciation of resulting ions and radicals is different. A host of highly energetic, short-lived ions and radicals other than OH• is generated in subterranean air. The first abstraction of an H atom from CH₄ requires a far higher activation energy than those of H atoms from methyl CH₃ and methylene CH₂ moieties. We argue that cave environments with elevated radioactivity may host short-lived, yet highly energetic radicals and ions that can supply the needed activation energy for first H-abstraction from CH₄ more efficiently than OH• in the open atmosphere. Thus, the application of kinetic and energetic findings of photochemical CH₄ oxidation in the open atmosphere may not be warranted for subterranean environments.

The α -radiation level in cave air is typically higher than in the open atmosphere because cave air is relatively close to rock and sediment surfaces with minerals harboring radioactive nuclides. The ionization rate in air via^{222} Rn radon decay is larger close to the ground, as reported for a Finnish forest [33], a spa [34], and in houses [35]. The effect is due to (i) strongly elevated radon concentrations in the air in porous, uranium-containing substrates and the rapid dilution of radon above surfaces upon mixing with the open atmosphere, especially during windy conditions. In contrast, cave air far from cave entrances is typically less turbulent and allows for a more even distribution of 222 Rn in cave air. (ii) Short-lived 220 Rn will always exhibit a greater abundance in air close to its parent nuclides in soil, rock, cave walls and floors [19]. Regardless, even exceptionally high combined radiation levels of 220 Rn and 222 Rn provided no evidence for accelerated CH₄ oxidation in our experiments.

A plausible reason for slow radiolytic reaction kinetics is the mismatch between the large number of CH₄ molecules in 1 m³ of atmosphere containing 1.85 ppmv CH₄ at standard conditions (i.e., $\sim 4.55 \cdot 10^{19}$ molecules CH₄) relative to the small number of radon-related nuclear decay events in the same volume of air (e.g., 10 kBq m⁻³ from ²²²Rn resulting from the decay of 10,000 atoms of ²²²Rn per second). The following simplistic numerical example illustrates the lack of feasibility of radiation-induced rapid oxidation of CH₄. If we assume that 1 m³ of atmosphere entering a cave with 10 kBq m⁻³, even if every decay of ²²²Rn leads to the oxidation of one molecule CH₄, it would require a geologic time period of ~ 144 million years to oxidize all CH₄. In reality, the nuclide-specific radiation from the decay of ²²²Rn alone is dwarfed by the



total radiation from radon, thoron, their radioactive progeny, and any other radioactive nuclides present in a given environment [19]. S1 File offers alternative calculations based on the assumptions that either (i) all energy from α -decay is exclusively invested in radiolytic dissociation of CH₄ and results in the oxidation of multiple molecules of CH₄ per decay event, or (ii) that only a fraction of the energy from α -decay is dissociating CH₄ in the overwhelming presence of other molecules and atoms. The calculated time periods needed to degrade 1.85 ppmv CH₄ at a ²²²Rn radiation level of 10 kBq m⁻³ range from 45.1 to 153,000 years, respectively. Even the most optimistic assumptions cannot speed up the radiolytic reaction kinetics to consume atmospheric CH₄ within hours to days.

We can use the most optimistic scenario for consumption of 1.85 ppmv CH₄ during 45.1 years at 10 kBq m⁻³ and calculate a radiation level of ~165 MBq m⁻³ that would be required to perform the same task in 24 h, which would be commensurate with kinetic CH₄ observations in caves. Natural radiation levels of a few MBq m⁻³ have been measured in air where ²²²Rn emanates through geologic faults from underlying uranium minerals [36]. Radiation levels in the range of MBq m⁻³ have been observed in the air of uranium mines [27]. Still, no location is known to offer values close to the required ~165 MBq m⁻³. We conclude that there is no natural cave environment on earth where the α -radiation level is strong enough to rapidly degrade CH₄. The same conclusion was recently described in a study that included arguments based on radiolytic kinetics of ion-induced reactions [15] that complement our calculations using α -decay and activation energy.

Subterranean radiation does not provide a mechanism for a fast-acting sink of atmospheric CH_4 that would extend to arid and hyperarid environments, unlike microbial methanotrophy. Our study does not invalidate the geochemical data from previous studies documenting CH_4 dynamics in subterranean ecosystems [10]. We do not call into question the fundamental importance of radiolysis of H_2O (and other air components) and subsequent redox reactions that are documented in the geologic record (e.g., [37]) or the long-term subterranean radiolytic impact on sedimentary organic matter [1]. However, the exceedingly slow chemical rates of reaction caused by natural rates of radiolysis would likely take years to geologic time periods in cave environments to deplete trace amounts of atmospheric CH_4 in cave air. As long as no alternative mechanisms have been identified, microbial methanotrophy serves as the only known fast-acting sink for subterranean CH_4 in the critical and vadose zones.

Conclusions

Strong radiation from radon isotopes and subsequent radiolysis of air proved unable to rapidly oxidize methane in dry or moist air. In the absence of a feasible alternative methane oxidation mechanism other than microbial methanotrophy, further studies are needed on the ability of microbes to consume trace amounts of methane in poorly ventilated caves, even though the trophic and energetic benefits become marginal at very low partial pressures of methane.

Supporting information

S1 File. An Excel file contains a first sheet "read me" with instructions and an overview on additional sheets offering analytical details and radiolysis calculations.

(XLSX)

Acknowledgments

We thank Dave Mattey and the staff of the Greenhouse Gas Research Group in the Department of Earth Science at RHUL for their support and technical suggestions during



experiments under project SMACKS (IEF-Marie Curie Actions nº 624204, FP7/2007-2013) and projects CGL2016-78318-C2-1R and 2R AEI/FEDER/UE. This study is based upon work at Indiana University supported by the U.S. Department of Energy, Office of Science, Office of Basic Energy Sciences, Chemical Sciences, Geosciences, and Biosciences Division under Award Number DESC0006978. In addition, financial support was provided by the National Science Foundation (1442246) and the US Army Research Office Grant W911NF-14-1-0411. Jay T. Lennon further acknowledges support from the Indiana University Office of the Vice President of International Affairs. Angel Fernandez-Cortes and Soledad Cuezva's work was further supported by the Spanish Ministry of Economy and Competitiveness projects CGL2016-78318-C2-1R and CGL2016-78318-C2-2R AEI/FEDER/UE. We are indebted to Scott Ensminger for guidance to a natural methane seep in New York State, Dương Nguyễn-Thuỳ and Hướng Nguyễn-Văn at Vietnam National University Hanoi for a ²²⁰Rn-emitting mud brick, and Greg Crouch at Indiana University for technical and radiation safety advice. Steffen Kolb, Tibor Kovacs, Leonie Anna Mueck, and two anonymous reviewers provided expert advice to improve the manuscript.

Author Contributions

Conceptualization: Arndt Schimmelmann, Angel Fernandez-Cortes, Thomas Streil.

Data curation: Arndt Schimmelmann, Angel Fernandez-Cortes.

Formal analysis: Arndt Schimmelmann, Angel Fernandez-Cortes, Soledad Cuezva, Jay T. Lennon.

Funding acquisition: Arndt Schimmelmann, Angel Fernandez-Cortes.

Investigation: Arndt Schimmelmann, Angel Fernandez-Cortes, Soledad Cuezva.

Methodology: Arndt Schimmelmann, Angel Fernandez-Cortes, Soledad Cuezva, Jay T. Lennon.

Project administration: Arndt Schimmelmann, Angel Fernandez-Cortes, Soledad Cuezva.

Resources: Arndt Schimmelmann, Angel Fernandez-Cortes, Thomas Streil.

Supervision: Arndt Schimmelmann, Angel Fernandez-Cortes.

Validation: Arndt Schimmelmann, Angel Fernandez-Cortes, Soledad Cuezva.

Visualization: Arndt Schimmelmann, Angel Fernandez-Cortes.

Writing – original draft: Arndt Schimmelmann, Angel Fernandez-Cortes.

Writing – review & editing: Arndt Schimmelmann, Jay T. Lennon.

References

- Greenwood PF, Shan C, Holman AI, Grice K. The composition and radiolysis impact on aromatic hydrocarbons in sedimentary organic matter from the Mulga Rock (Australia) uranium deposit. Org. Geochem. 2018; 123: 103–112. https://doi.org/10.1016/j.orggeochem.2018.06.013
- Colwell FS, D'Hondt S. Nature and extent of the deep biosphere, Rev. Mineral. Geochem. 2013; 75: 547–574. http://dx.doi.org/10.2138/rmg.2013.75.17
- Colman DR, Poudel S, Stamps BW, Boyd ES, Spear JR. The deep, hot biosphere: Twenty-five years of retrospection. PNAS 2017; 114 (27): 6895–6903. https://dx.doi.org/10.1073/pnas.1701266114 PMID: 28674200
- Xu X, Yuan F, Hanson PJ, Wullschleger SD, Thornton PE, Riley WJ, et al. Reviews and syntheses: Four decades of modeling methane cycling in terrestrial ecosystems. Biogeosciences 2016; 13: 3735–3755. https://dx.doi.org/10.5194/bg-13-3735-2016



- IPCC. Climate Change 2013: The Physical Science Basis, Figure SPM.5 of the Summary for Policy-makers; 2013. http://www.ipcc.ch/pdf/assessment-report/ar5/wg1/WG1AR5_SPM_FINAL.pdf (accessed on September 4th, 2018)
- Prather MJ, Holmes CD. Overexplaining or underexplaining methane's role in climate change. PNAS 2017; 114 (21): 5324–5326. https://dx.doi.org/10.1073/pnas.1704884114 PMID: 28507153
- Edwards CR, Onstott TC, Miller JM, Wiggins JB, Wang W, Lee CK, et al. Draft genome sequence of uncultured upland soil cluster Gammaproteobacteria gives molecular insights into high-affinity methanotrophy. Genome Announc. 2017; 5: e00047–17. https://doi.org/10.1128/genomeA.00047-17 PMID: 28450499
- Waring CL, Griffith DWT, Wilson S, Hurry S. Cave atmosphere: A guide to calcification and a methane sink. Geochim. Cosmochim. Acta 2009; 73 (13): A1419–A1419. http://apo.ansto.gov.au/dspace/ handle/10238/2461?mode=full (accessed on September 4th, 2018)
- Mattey DP, Fisher R, Atkinson TC, Latin JP, Durrell R, Ainsworth M, et al. Methane in underground air in Gibraltar karst. Earth Planet. Sci. Lett. 2013; 374: 71–80. http://dx.doi.org/10.1016/j.epsl.2013.05. 011
- Fernandez-Cortes A, Cuezva S, Alvarez-Gallego M, Garcia-Anton E, Pla C, Benavente D, et al. Subterranean atmospheres may act as daily methane sinks. Nat. Commun. 2015; 6: 7003. https://dx.doi.org/10.1038/ncomms8003 PMID: 25912519
- McDonough LK, Iverach CP, Beckmann S, Manefield M, Rau GC, Baker A, et al. Spatial variability of cave-air carbon dioxide and methane concentrations and isotopic compositions in a semi-arid karst environment. Environ. Earth Sci. 2016: 75: 700. http://dx.doi.org/10.1007/s12665-016-5497-5
- Webster KD, Mirza A, Deli JM, Sauer PE, Schimmelmann A. Consumption of atmospheric methane in a limestone cave in Indiana, USA. Chem. Geol. 2016; 443: 1–9. https://dx.doi.org/10.1016/j.chemgeo. 2016.09.020
- Webster KD, Drobniak A, Etiope G, Mastalerz M, Sauer PE, Schimmelmann A. Subterranean karst environments as a global sink for atmospheric methane. Earth Planet. Sci. Lett. 2018; 485: 9–18. https://doi.org/10.1016/j.epsl.2017.12.025
- Nguyễn-Thuỳ D, Schimmelmann A, Nguyễn-Văn H, Drobniak A, Lennon JT, Tạ PH, et al. Subterranean microbial oxidation of atmospheric methane in cavernous tropical karst. Chem. Geol. 2017; 466: 229– 238. https://doi.org/10.1016/j.chemgeo.2017.06.014
- Waring CL, Hankin SI, Griffith DWT, Kertesz MA, Kobylski V, Wilson NL, et al. Seasonal total methane depletion in limestone caves. Sci. Rep. 2017; 7: 8314. https://dx.doi.org/10.1038/s41598-017-07769-6 PMID: 28814720
- Haynes RM, Kebarle P. Mass-spectrometric study of ions at near-atmospheric pressure. III. The alpha radiolysis of methane. J. Chem. Phys. 1966; 45 (11): 3899–3906. https://dx.doi.org/10.1063/1. 1727435
- Rasul G, Prakash GKS, Olah GA. Comparison of structures and energies of CH52+• with CH4+• and their possible role in superacidic methane activation. PNAS 1997; 94 (21): 11159–11162. https://dx.doi.org/10.1073/pnas.94.21.11159 PMID: 11038583
- Lennon JT, Nguyễn-Thùy D, Phạm TM, Drobniak A, Tạ PH, et al. Microbial contributions to subterranean methane sinks. Geobiology 2017; 15 (2): 254–258. https://dx.doi.org/10.1111/gbi.12214 PMID: 27671735
- Meisenberg O, Mishra R, Joshi M, Gierl S, Rout R, Guo L, et al. Radon and thoron inhalation doses in dwellings with earthen architecture: Comparison of measurement methods. Sci. Total Environ. 2017; 579: 1855–1862. http://dx.doi.org/10.1016/j.scitotenv.2016.11.170 PMID: 27939079
- Markkanen M, Arvela H. Radon emanation from soils. Radiat. Prot. Dosimetry 1992; 45 (1–4), 269– 272. https://doi.org/10.1093/oxfordjournals.rpd.a081541
- Morawska L, Phillips CR. Dependence of the radon emanation coefficient on radium distribution and internal structure of the material. Geochim. Cosmochim. Acta 1993; 57 (8): 1783–1797. https://doi.org/10.1016/0016-7037(93)90113-B
- Hursh JB. Thoron half-life. J. Inorg. Nucl. Chem. 1966; 28 (12): 2771–2776. https://doi.org/10.1016/
 0022-1902(66)80001-5
- Wang J, Meisenberg O, Chen Y, Karg E, Tschiersch J. Mitigation of radon and thoron decay products by filtration. Sci. Total Environ. 2011; 409 (19): 3613–3619. https://doi.org/10.1016/j.scitotenv.2011.06. 030 PMID: 21767867
- Oremland RS, Capone DG. Use of "specific" inhibitors in biogeochemistry and microbial ecology. In: Marshall KC, editor. Advances in Microbial Ecology 10 (ch. 8), Plenum Publishing Corporation; 1988. p. 335. https://dx.doi.org/10.1007/978-1-4684-5409-3_8



- Farhan UI Haque M, Crombie AT, Ensminger SA, Baciu C, Murrell JC. Facultative methanotrophs are abundant at terrestrial natural gas seeps. Microbiome 2018; 6: 118. https://doi.org/10.1186/s40168-018-0500-x PMID: 29954460
- Cigna AA. Radon in caves. Int. J. Speleol. 2005; 34 (1–2): 1–18. http://scholarcommons.usf.edu/cgi/ viewcontent.cgi?article=1203&context=ijs (accessed on September 4th, 2018)
- 27. Wismut GmbH and MECSEK-ÖKO. Annual Report 2010 on the Cooperation between Wismut GmbH and MECSEK-ÖKO. 2011; p. 24–26. https://www.google.com/url?sa=t&rct=j&q=&esrc=s&source=web&cd=3&ved=2ahUKEwjm7rrq5JrdAhWF34MKHScrBTQQFjACegQICBAC&url=http%3A%2F%2Fgov.pecs.hu%2Fdownload%2Findex.php%3Fid%3D309232&usg=AOvVaw0u7lRhg7ZYhYU2lGyshAXv (accessed on September 4th, 2018)
- Yan Z, Liu C, Liu Y, Bailey VL. Multiscale investigation on biofilm distribution and its impact on macroscopic biogeochemical reaction rates. Water Resour. Res. 2017; 53 (11): 8698–8714. https://dx.doi.org/10.1002/2017WR020570
- Mancinelli RL. The regulation of methane oxidation in soil. Annu. Rev. Microbiol. 1995; 49: 581–605. https://doi.org/10.1146/annurev.mi.49.100195.003053 PMID: 8561473
- Conrad R. Microbial ecology of methanogens and methanotrophs. Adv. Agron. 2007; 96: 1–63. https://doi.org/10.1016/S0065-2113(07)96005-8
- Lindmark A, Rosen B. Radon in soil gas—Exhalation tests and in situ measurements. Sci. Total Environ. 1985; 45: 397–404. https://doi.org/10.1016/0048-9697(85)90243-8 PMID: 4081740
- Papastefanou C. Measuring radon in soil gas and groundwaters: a review. Annals Geophys. 2007; 50
 (4): 569–578. http://www.annalsofgeophysics.eu/index.php/annals/article/download/3070/3113
 (accessed on September 4th, 2018)
- Tammet H, H

 forrak U, Laakso L, Kulmala M. Factors of air ion balance in a coniferous forest according to measurements in Hyytiälä, Finland. Atmos. Chem. Phys. 2006; 6: 3377–3390. https://doi.org/10.5194/acp-6-3377-2006
- Cosma C, Suciu I, Jäntschi L, Bolboacă SD. Ion-molecule reactions and chemical composition of emanated from Herculane Spa geothermal sources. Int. J. Mol. Sci. 2008; 9 (6): 1024–1033. https://dx.doi.org/10.3390/ijms9061024 PMID: 19325844
- Kolarz PM, Filipović DM, Marinković BP. Daily variations of indoor air-ion and radon concentrations.
 Appl. Radiat. Isot. 2009; 67 (11): 2062–2067. https://doi.org/10.1016/j.apradiso.2009.07.023 PMID: 19700332
- Seyis C, İnan S, Streil T. Ground and indoor radon measurements in a geothermal area. Acta Geophys. 2010; 58 (5): 939–946. https://link.springer.com/article/10.2478/s11600-010-0012-y
- Lefticariu L, Pratt LA, LaVerne JA, Schimmelmann A. Anoxic pyrite oxidation by water radiolysis products—A potential source of biosustaining energy. Earth Planet. Sci. Lett. 2010; 292 (1–2): 57–67. https://dx.doi.org/10.1016/j.epsl.2010.01.020