

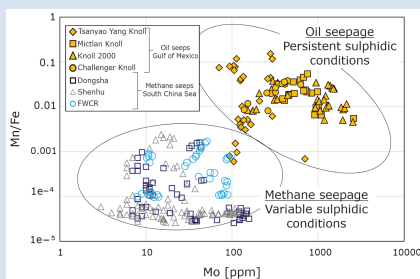
Pyrite-based trace element fingerprints for methane and oil seepage

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<https://doi.org/10.7185/geochemlet.2409>

Abstract



Pyrite forms at marine hydrocarbon seeps as the result of the microbial oxidation of methane, organic matter, and crude oil coupled to sulphate reduction. Redox sensitive and nutrient trace elements in pyrite may hold valuable information on present and past seepage events, the evolution of fluid composition, as well as the presence of heavy hydrocarbon compounds from crude oil. This study uses the trace element compositions of pyrite that formed at methane seeps and crude oil-dominated seeps to constrain element mobilities during the sulphate reduction processes, and examine the degree to which specific trace elements are captured by pyrite. Pyrite forming at oil seeps shows high Mn/Fe ratios and high Mo content compared to pyrite from methane seeps. These patterns suggest either more intense or persistent sulphidic conditions, or an intensified manganese (oxy)hydroxide shuttle process at oil seeps. Copper and Zn are enriched in oil seepage-derived pyrite while Ni and V enrichment is less pronounced, suggesting either a selective uptake of specific elements by pyrite, or varying trace element compositions of organic compounds oxidised *via* microbial reduction.

Received 24 September 2023 | Accepted 30 January 2024 | Published 5 March 2024

Introduction

Marine hydrocarbon seeps support chemosynthesis-based microbial habitats of archaea and sulphate reducing bacteria performing the sulphate driven anaerobic oxidation of methane (SD-AOM). This process triggers the formation of authigenic minerals that record the dynamics and intensity of seepage, redox fluctuations, and the evolution of fluid composition and microbial activity (Feng *et al.*, 2009; Smrzka *et al.*, 2016, 2019a). Pyrite is a widespread mineral that forms during sulphate reduction at seeps, which are hotspots of pyrite authigenesis. Detailed studies on pyrite morphology and its stable isotopic and trace element compositions have recently been conducted in seep settings (Lin *et al.* 2022; Wang *et al.*, 2022).

Pyrites from various environments show different trace element patterns depending on morphology, grain size, formation temperatures, and the composition of parent fluids (Gregory *et al.*, 2015). The trace element inventory of pyrite has been refined into a first order proxy for the deep time evolution of Earth's biosphere (Large *et al.*, 2014). Sedimentary pyrite formation is controlled by the biogeochemical cycles of sulphur, carbon and iron, and constitutes a relevant long term sink for trace elements during early diagenesis (Huerta-Diaz and Morse, 1992). Pyrite scavenges and incorporates trace elements from parent fluids, including many redox sensitive and bio-essential trace metals including Mn, Mo, Ni, Cu, Zn, Cr, As and Se (Huerta-Diaz and Morse, 1992;

Morse and Arakaki, 1993). Pyrite based proxies for fluid composition, sulphate reduction processes and redox conditions are a trending topic in hydrocarbon seep research (Miao *et al.*, 2022, Lin *et al.*, 2022; Wang *et al.*, 2022; Domingos *et al.*, 2023). Hydrocarbon emissions from natural methane and oil seeps represent prominent pathways of carbon transfer from the geosphere to the hydrosphere. Constraining their influence on the marine carbon and sulphur cycles is therefore critical in order to improve the quantification of global methane budgets and to understand the dynamics of – and responses to – natural and anthropogenic oil spills. A pyrite based proxy offers insights into the evolution of fluid composition at seeps, which is critical because fluid composition governs microbial and metazoan ecology (Orcutt *et al.*, 2010). The significance of distinguishing oil-from methane-dominated seep systems has gained traction in recent years in the search for end member system identification (Smrzka *et al.*, 2016; Akam *et al.*, 2021; Krake *et al.*, 2022), and reliable proxies are continuously being explored and refined.

This study presents a first comparison of the trace element compositions of authigenic pyrite derived from methane and oil seeps. Motivated by current efforts to exploit the potential of pyrite based geochemical proxies, we provide new constraints on environmental conditions during seepage and microbial oxidation of heavy hydrocarbons, and establish trace element fingerprints to distinguish methane seeps from oil dominated seeps. These results improve our understanding of trace element

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liberation during microbial sulphate reduction, and their subsequent incorporation into pyrite during early diagenesis, while emphasising the role of pyrite based trace element geochemistry as a main, or complementary, source of information on redox conditions and fluid compositions in modern, and potentially, ancient seepage environments.

Study Sites – Endmembers of Hydrocarbon Seepage

Two localities of methane and heavy hydrocarbon seepage were selected for a comparative study of authigenic pyrite forming in different environments. The Campeche and Sigsbee Knolls in the southern Gulf of Mexico (GoM; Fig. 1a) are two salt provinces that exhibit a set of hummocky seafloor structures related to salt tectonism, enabling the seepage of methane and crude oil, as well as the formation of gas hydrates within the sediments and on the seafloor (Sahling *et al.*, 2016). Crude oil and asphalt in the southern GoM fuel microbial sulphate reduction independent of SD-AOM (Joye *et al.*, 2004), and have shaped a unique environment inhabited by distinct macro- and micro-faunal communities (Orcutt *et al.*, 2010). This study considers the Campeche and Sigsbee Knolls as crude oil dominated end member seepage systems.

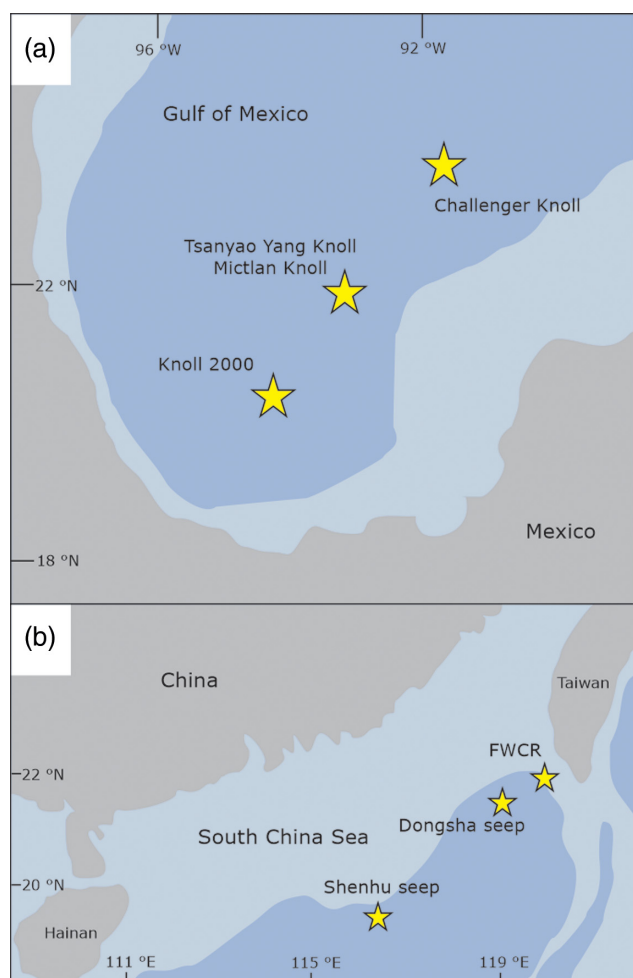


Figure 1 Locations of the investigated sediment cores. (a) Gravity core locations from oil seeps at Campeche and Sigsbee Knolls in the southern Gulf of Mexico. (b) Gravity core locations of methane seeps from the South China Sea.

The South China Sea (SCS) is a large marginal sea in the western Pacific Ocean located at the transect between the Eurasian, Pacific, and Indian plates (Fig. 1b). The northern SCS is a passive continental margin covered by thick successions of Neogene sediments that promote abundant hydrocarbon generation manifested as mud volcanoes and diapirs, gas chimneys, and seafloor seepage sites. Among the numerous seep sites discovered in the SCS over the past decades, the Dongsha and Shenhu seepage provinces are among the best studied sites, which have been previously studied regarding the genesis of authigenic pyrite (Lin *et al.*, 2022). To the northeast of these seepage areas, and related to the convergence of the Eurasian and the Philippine Sea plates, lies the south-western Taiwan accretionary prism, which harbours the Yam Seep area located at the northern crest of Four-Way Closure Ridge (FWCR; Tseng *et al.*, 2023). The Dongsha, Shenhu, and Yam Seep sites are all characterised by seepage of biogenic and thermogenic methane, representing methane dominated end member seepage systems.

Material and Methods

Authigenic pyrite was obtained from a total of five gravity cores, which were sampled over the course of four cruises between 2015 and 2018 in the southern GoM and the northern SCS (Fig. 1). Between 20 and 30 grams of sediment were sampled from the cores at intervals of 20 cm, freeze dried over 24 hrs, and subsequently powdered by hand using an agate pestle and mortar. Aliquots of unpowdered sediment were then sieved with deionised water through a 0.063 mm sieve, and pyrite aggregates were hand picked under a binocular microscope from the coarse fraction. Pyrite grains were mounted onto epoxy discs, polished to a smooth surface and coated with carbon for scanning electron microscopy and electron probe microanalysis (EMPA). Major element content in pyrite was determined using a Cameca SX-100 electron microprobe. Major and trace element composition of pyrite was determined *via* laser ablation inductively coupled plasma mass spectrometry (LA-ICP-MS). All element data and additional information are given in the Supplementary Information.

Results

Manganese, Mo, Cu, and Zn contents in pyrite show different distribution patterns for the two investigated types of hydrocarbon seepage. Pyrite from oil dominated seeps is characterised by higher content of Mn and Mo than in methane seepage-derived pyrite. The Mn enrichment is expressed as Mn/Fe ratios shown in Figure 2, which is two to three orders of magnitude higher in oil seepage-derived pyrite. Molybdenum content is higher by one order of magnitude in oil seep pyrite (Fig. 2). Methane seep pyrite from the four sites shows variable Mn/Fe ratios. The contents of Cu and Zn are higher in oil seep pyrite than in methane seep pyrite, and the distribution of these elements allows for a distinction between the two seepage environments (Fig. 3a). This distinction is less clear for Ni and V (Fig. 3b).

Discussion

A combined Fe-Mn-Mo fingerprint. Microbial sulphate reduction at seeps is coupled to the oxidation of methane, sedimentary organic matter, and high molecular weight hydrocarbons that make up crude oil (Joye *et al.*, 2004; Smrzka *et al.*, 2019a). The microbial oxidation rates of these compounds are governed by the microbial consortium capable of using the particular electron donors, which affects sulphate reduction rates and thus the

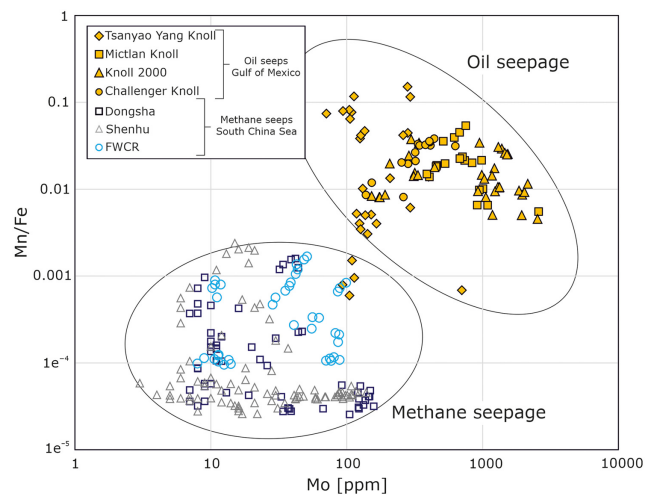


Figure 2 Mn/Fe ratios and Mo content in methane seepage- and oil seepage-derived authigenic pyrite. Data from Dongsha and Shenhu seepage areas are from Lin *et al.* (2022).

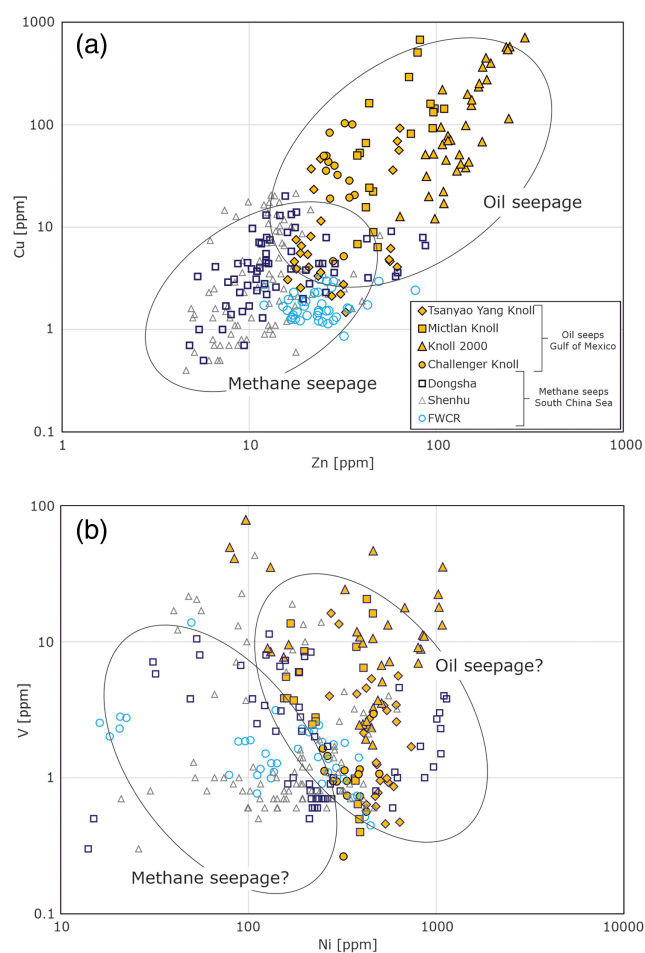


Figure 3 Trace element content in methane seepage- and oil seepage-derived authigenic pyrite. (a) Zn and Cu content, (b) Ni and V content; data from Dongsha and Shenhu seepage areas are from Lin *et al.* (2022).

amount of sulphide released to sedimentary pore water (Smrzka *et al.*, 2019a). The concentration of dissolved sulphide species in pore fluids in turn influences the solubility and mobility of redox sensitive trace elements, particularly Mo and to a lesser extent

Mn. In addition, the oxidation of sedimentary organic matter and crude oil will inevitably lead to the partial or complete breakdown of organic compounds, which are themselves carriers of trace elements (Smrzka *et al.*, 2020).

Manganese and Mo contents in pyrite allow us to distinguish between the two seepage environments (Fig. 2). The Mo content in oil seep pyrite is up to an order of magnitude higher than in methane seep pyrite and sedimentary pyrite reported from black shales (Gregory *et al.*, 2015). Molybdenum is incorporated into pyrite under sulphidic conditions, after stabilising as thiomolybdate, and may also be fixed by organic matter (Morse and Luther, 1999; Gregory *et al.*, 2015). Manganese does not usually reside in pyrite in high concentrations, yet may form sulphides adsorbed to mackinawite; a precursor mineral of pyrite during early diagenesis (Gregory *et al.*, 2015). Manganese is usually incorporated into Ca-rich rhodochrosite or high-Mg calcite under sulphidic conditions during early diagenesis (Suess, 1978), but may incorporate into pyrite at high Mn²⁺ concentrations (Morse and Arakaki, 1993; Morse and Luther, 1999). Manganese incorporation into pyrite may also proceed *via* the uptake of precursor manganese sulphide phases, which are stable at high Mn/Fe ratios and high sulphide levels (Shikazono *et al.*, 1994). A recent study indicates that Mn is distributed randomly within pyrite and held within microcrystals that formed early during diagenesis, suggesting that the partitioning behaviour of dissolved Mn into authigenic minerals is not as straightforward as previously thought (Atienza *et al.*, 2023). Although Mn can be enriched in pyrite formed by SD-AOM (Lin *et al.*, 2022), the Mn/Fe ratios in pyrite from oil seeps are orders of magnitude higher than in pyrite from methane seeps (Fig. 2). High levels of dissolved sulphide favour the incorporation of Mn and Mo into pyrite (Wang *et al.*, 2022), suggesting that the microbial oxidation of crude oil in sediments may enable persistent and highly sulphidic conditions in ambient pore waters controlled by the extent and rate of microbial metabolism. Microbial crude oil oxidation coupled to sulphate reduction enables carbonate precipitation and sulphide production, despite its generally slower microbial oxidation rate compared to SD-AOM (Joye *et al.*, 2004; Smrzka *et al.*, 2019a). Components of crude oils contain variable contents of organic sulphur residing in compounds including thiols, sulphides, and thiophenes (Tissot and Welte, 1984). The high sulphur content in oils (>2 wt. %) from the southern GoM and the increase of dissolved hydrogen sulphide gas emitted at oil seeps (Smrzka *et al.*, 2019a) suggest that the microbial mineralisation of crude oil may act as an additional source of sulphur to pore waters.

The high Mn/Fe ratios of oil seep pyrite may also be due to an effective manganese (oxy)hydroxide shuttle process, transferring adsorbed trace elements from seawater to the sediments (Scholz *et al.*, 2013; Smrzka *et al.*, 2020). This shuttle may be intensified by the seepage of crude oil, acting as an additional source of Mn from sedimentary pore waters to bottom waters. Oil seeps at the Campeche Knolls emit oil droplets, oily gas bubbles, and asphalt fragments (Sahling *et al.*, 2016), which could act as an additional transport agent for trace elements to bottom waters. The emitted oil components will be oxidised aerobically in the bottom waters around the locus of seepage, releasing adsorbed or incorporated trace elements from organic compounds. This oil enhanced manganese (oxy)hydroxide shuttle would also explain the observed co-enrichment of Mn and Mo (*cf.* Scholz *et al.*, 2013) and to a lesser extent Cu (Fig. 3a) in oil seep pyrite.

A combined Cu-Zn-Ni-V fingerprint. Copper and Zn are commonly co-enriched in marine organic matter and seafloor sediments, yet they are taken up into pyrite to different degrees. Copper forms strong complexes with organic matter and

precipitates as copper sulphide during sulphate reduction (Morse and Luther, 1999). Copper enrichment in oil seep pyrite is likely derived from two sources, either released by reductive dissolution of manganese (oxy)hydroxides leading to a co-enrichment of Mn and Cu (Figs. 2, 3a) and released directly from crude oil during its microbial mineralisation. The affinity for Zn incorporation into pyrite is lower than for Cu due to ZnS precipitation prior to pyrite formation (Morse and Luther, 1999). Zinc content varies considerably in diagenetic pyrite where it may be present as sphalerite inclusions (Gregory *et al.*, 2015). However, both Zn and Cu are co-enriched in oil seep pyrite, suggesting an overriding effect of microbial crude oil oxidation coupled to sulphate reduction. Nickel and V are micronutrients for phytoplankton growth, and are transported to the seafloor by organic particles and metal (oxy)hydroxides (Smrzka *et al.*, 2019b). While Ni is commonly divalent in marine sediments, V is sensitive toward pH, redox conditions and dissolved sulphide concentrations, and V enrichment in reducing sediments relative to average continental crust (Thomson *et al.*, 1998). Whereas Ni is incorporated into pyrite during early diagenesis, V usually resides in the non-sulphide fraction such as organic matter, silicates and carbonates (Huerta-Diaz and Morse, 1992; Gregory *et al.*, 2015). In contrast to Cu and Zn, the distribution of Ni and V show large overlap without clear enrichments between methane and oil seep pyrite (Fig. 3b).

The release of trace elements during organic matter oxidation represents a source of trace elements during early diagenesis (Smrzka *et al.*, 2020), reflected by the trace element composition of pyrite (Miao *et al.*, 2022; Chen *et al.*, 2023). The present data set expands this distinction regarding the presence of oil compounds and their microbial mineralisation in marine sediments. Crude oils contain trace elements including Ni, V, Cu, and Zn, which are present as metalloporphyrin complexes derived from bacterial and plant pigments, as metal centres of microbial enzymatic cofactors, and in large organic matrices or other metal-binding functional groups (Duyck *et al.*, 2007). Only Cu and Zn are systematically enriched in oil seep pyrite relative to its methane seep counterpart. These distribution patterns may be controlled by differences in the incorporation behaviour of the respective element into the pyrite structure (Gregory *et al.*, 2015), the trace element composition of the oxidised organic compounds, the varying rates of microbial oil degradation, and the composition of the host sediment (Figure S-2) during pyrite formation.

Nickel and V reside in crude oils primarily as petroporphyrin complexes that are primarily part of the asphaltene fraction (Duyck *et al.*, 2007). Petroleum hydrocarbons from southern GoM seeps are biodegraded, weathered (Schubotz *et al.*, 2011) and of low thermal maturity with Zn, Cu and Ni content between 10 and 100 ppm, and V content of up to 400 ppm (Smrzka *et al.*, 2019a). The lack of V enrichment in oil seep pyrite is most likely due to its low affinity towards pyrite incorporation, which is lower than for other trace metals (Gregory *et al.*, 2015). Although a slight Ni enrichment in oil seep pyrite is present, it cannot be used as an unambiguous indicator for microbial crude oil mineralisation. Manganese (oxy)hydroxides are also effective transport agents for Ni and V to sediments (Smrzka *et al.*, 2019b), yet contrary to Mn and Mo, they are not distinctly enriched in oil seep pyrite. Also, the degree to which Ni and V are released from organic matter during microbial remineralisation may be smaller than for Cu and Zn. For instance, although all four elements reside in metalloporphyrins of organic matter and oils, biodegradation of nickel porphyrins is slower than for copper and vanadyl porphyrins (Sadowski *et al.*, 2007). Microbial mineralisation of crude oil, which occurs both anaerobically in the sediments and aerobically in bottom waters, may

selectively release trace elements from the oils depending on the complexity and size of their respective organic molecules.

Synthesis

Manganese and Mo are enriched in authigenic pyrite derived from oil seeps compared to pyrite derived from methane seeps. This co-enrichment may be due to a persistent sulphidic environment that is seldom disrupted by pulses of oxygenation due to shifting redox boundaries in the sediment; conditions maintained by the oxidation of crude oil. These conditions would allow for an effective incorporation of Mo into pyrite, while also providing conditions that facilitate Mn uptake by pyrite. The presence of a Mn (oxy)hydroxide shuttle process at oil seeps would also explain the co-enrichment of Mn and Mo in oil seep pyrite. The enrichment of Cu, Zn, and Ni at oil seeps also argues for persistent sulphidic conditions and an additional source of these elements to pore water, both being triggered by the microbial oxidation of crude oil. The presence of an enhanced particulate shuttle process driven by Mn (oxy)hydroxides at oil seeps as invoked above is currently unknown, and represents a yet poorly constrained source for trace elements to bottom waters above seepage environments. Distinguishing the exact sources of trace elements in the two seepage systems that may include crude oil, as well as the dissolution of carbonates and clay minerals in the host sediments, will shed more light on trace element dynamics during early diagenesis. Comparing these results to other seepage sites from more diverse environments will increase the potential of pyrite based trace element proxies.

Acknowledgements

This research was funded by the “Independent Projects for Postdocs” grant scheme awarded to DS by the Central Research Development Fund of the University of Bremen. We thank Anne Hübner, Stefan Sopke, Janice Malnati, and Andreas Klügel (University of Bremen) for sample preparation and assistance in EMPA analyses.

Editor: Juan Liu

Data Availability Statement

Data will be made available on the Pangaea database (www.pangaea.de).

Additional Information

Supplementary Information accompanies this letter at <https://www.geochemicalperspectivesletters.org/article2409>.



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Cite this letter as: Smrzka, D., Lin, Z., Monien, P., Chen, T., Bach, W., Peckmann, J., Bohrmann, G. (2024) Pyrite-based trace

element fingerprints for methane and oil seepage. *Geochem. Persp. Let.* 29, 33–37. <https://doi.org/10.7185/geochemlet.2409>

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