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**Citation:** Akam, Sajjad A, Lyons, Timothy W, Coffin, Richard B, McGee, David, Naehr, Thomas H et al. 2021. "Carbon-sulfur signals of methane versus crude oil diagenetic decomposition and U-Th age relationships for authigenic carbonates from asphalt seeps, southern Gulf of Mexico." Chemical Geology, 581.

As Published: 10.1016/J.CHEMGE0.2021.120395

Publisher: Elsevier BV

Persistent URL: https://hdl.handle.net/1721.1/148113

**Version:** Original manuscript: author's manuscript prior to formal peer review

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PII:	80009-2541(21)00338-7
DOI:	https://doi.org/10.1016/j.chemgeo.2021.120395
Reference:	CHEMGE 120395
To appear in:	Chemical Geology
Received date:	16 August 2020
Revised date:	9 June 2021
Accepted date:	11 June 2021

Please cite this article as: S.A. Akam, T.W. Lyons, R.B. Coffin, et al., Carbon-sulfur signals of methane versus crude oil diagenetic decomposition and U-Th age relationships for authigenic carbonates from asphalt seeps, southern Gulf of Mexico, *Chemical Geology* (2021), https://doi.org/10.1016/j.chemgeo.2021.120395

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# Carbon-sulfur signals of methane versus crude oil diagenetic decomposition and U-Th age relationships for authigenic carbonates from asphalt seeps, southern Gulf of Mexico.

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## A 'stract:

Offshore hydrocarbon accumulations in u.e Gulf of Mexico (GoM) are often accompanied by natural seepage of oil and gas from subsurface reservoirs into shallow sediments and the water column. This study investigated the temporal patterns and carbon-sulfur (C-S) coupling associated with authigenic carbonate samples recovered from surface sediments of a crude oil seepage site in southern GoW (Chapopote asphalt volcano, Bay of Campeche) using radioactive U-Th dates, and stable C, O, and S isotopes. The results were compared with data from multiple seep sites in the northern GoM where methane seepage is dominant along with non-methane hydrocarbons (ethane , propane, crude oil, etc.). U-Th age-dating of Chapopote asphalt seepage has been ongoing for thousands of years. The results are also consistent with previous studies from the northern GoM that hypothesize that seeps along the GoM continental slope were active during the last deglaciation.  $\delta^{13}C_{CaCO3}$  and  $\delta^{18}O_{CaCO3}$  values from authigenic carbonates at Chapopote indicated a mixed contribution of methane and non-methane hydrocarbons to the dissolved inorganic carbon (DIC) pool, consistent with previous results. Comparison of  $\delta^{13}C_{CaCO3}$  vs.  $\delta^{34}S_{CRS}$  (CRS = chromium reducible sulfur) from carbonate samples showed

noticeable differences at the Chapopote seep site (average  $\delta^{13}C_{CaCO3}$ -25‰ VDPB,  $\delta^{34}S_{CRS}$ -27‰ VCDT) relative to the methane seep-dominated samples from the northern GoM (average  $\delta^{13}C_{CaCO3} < -40\%$  VDPB,  $\delta^{34}S_{CRS} > 0\%$  VCDT). Our results point toward distinguishable differences in the paired  $\delta^{13}C_{DIC}$  and  $\delta^{34}S_{sulfide}$  signatures produced via the diagenetic processes of sulfate-driven anaerobic oxidation of methane versus non-methane hydrocarbons. These results potentially provide an important proxy for identification of such diagenetic processes within the sedimentary records.

Keywords: Hydrocarbon seeps, authigenic carbonates, U-Th dating, Gulf of Mexico, microbial sulfate reduction, anaerobic hydrocarbon oxidation, Carbon-Sulfate cycling

#### 1. Introduction

Tectonic and depositional conditions in the Gulf of Mexico (GoM) provide an ideal setting for the formation of oil and gas as well as its  $u_{\rm I}$  ward migration from subsurface reservoirs to the seafloor and water column (Sassen et al., 1998). Natural hydrocarbon seeps may account for ~47% of all crude oil entering the marine environment (Kvenvolden and Cooper, 2003), with an estimated 160 to 690 thousand barrels of oil entering the Gulf of Mexico annually (Kvenvolden and Harbaugh, 1983; Mitchell et al., 1999; National Research Council, 2003; MacDonald et al., 2015). These seepage sites have a significant impact on geology and biology of the seabed facilitated by microbial<sup>1</sup>y-meutated biogeochemical processes (Judd and Hovland, 2009; Hovland et al., 2012; Cu<sup>ct</sup>m et al., 2015; Chakraborty et al., 2020).

GoM seeps are characterized by significant quantities of authigenic minerals formed by the activities of chemosymbiotic microbial communities (Roberts and Feng, 2013). These processes are tightly linked to biogeochemical turnover and the combination of downward-diffusing seawater and upward advection of hydrocarbon-rich pore fluids (Roberts and Aharon, 1994; Sassen et al., 2004; Roberts and Feng, 2013; Suess, 2018). Furthermore, chemosymbiotic benthic biota are supported by microbially-driven anaerobic oxidation of methane (AOM), which involves a microbial consortium of anaerobic methanotrophic archaea (ANME) and sulfate-reducing bacteria (SRB). AOM anaerobically oxidizes methane while reducing sulfate to sulfide

at the sulfate-methane transition zone (SMTZ), with the following net reaction (Boetius et al., 2000):

$$CH_4 + SO_4^{2-} \rightarrow HCO_3^- + HS^- + H_2O.$$
(1)

Bicarbonate (HCO<sub>3</sub><sup>-</sup>) from the process enters the dissolved inorganic carbon (DIC) pool in shallow sediments. Authigenic carbonates are common early diagenetic precipitates formed at methane and other hydrocarbon seep sites (Aloisi et al., 2002; Naehr et al., 2007). They are formed primarily through bicarbonate production via AOM, while results in supersaturation of carbonate ions with respect to carbonate minerals in porewriter at the SMTZ and induces carbonate precipitation (Baker and Burns, 1985; Ritger et al.,  $1>^{2}$ , Eq. 2):

$$2\text{HCO}_3^- + \text{Ca}^{2+} \rightarrow \text{CaCO}_3 + \text{CO}_2 + \text{H}_2\text{O}.$$
(2)

Carbonate authigenesis at seep sites is a substantial carbon requestration process and provides an excellent geologic archive of diagenetic geochem. All eactions involving both carbon and sulfur at seeps. At present-day settings, for examine, uuligenic carbonate precipitation is considered to be a significant component of the marine earbon sink (Wallmann et al., 2008; Torres et al., 2020), accounting for 10-15% carbonate arcumulation in pelagic and neritic sediments (Sun and Turchyn, 2014; Akam et al., 2020). Further, carbon sequestration due to authigenic carbonate formation may have played an over larger role in geological past, especially during periods of widespread oceanic anoxia (Schrag et al., 2013). The first-order pathways and processes in the cycles of C, S, and other elements are relatively well constrained for modern seep sites. However, much of the chail, magnitude, and variability of these biogeochemical processes as well as their impacts on geological carbon budgets are still unclear. A better understanding of seep-associated C-S coupling is of great importance in recognizing the role of these processes on the ancient Earth, including relationships to Oceanic Anoxic Events and early evolution of oxygen in Earth's atmosphere (Higgins et al., 2009; Bristow and Grotzinger, 2013; Canfield and Kump, 2013; Havig et al., 2017).

Studies of organic biomarkers in sediments from the GoM (Pancost et al., 2005) revealed disparities between measured rates of sulfate reduction versus methane oxidation that suggest other carbon sources in addition to methane (Joye et al., 2004; Bowles et al., 2011). Further, carbon isotope data from authigenic carbonates (Formolo et al., 2004) also suggest that AOM

may not be the only process responsible for an increase in carbonate alkalinity at many seep sites. Rather, it seems likely that the anaerobic oxidation of non-methane hydrocarbons (AONM) coupled with sulfate reduction (Widdel and Rabus, 2001; Kniemeyer et al., 2007; e.g., hexadecane, Eq. 3, propane, Eq. 4) provides a significant source of metabolic energy and bicarbonate at these locations (Formolo et al., 2004; Joye et al., 2004; Naehr et al., 2009; Mansour and Sassen, 2011; Smrzka et al., 2019; Sun et al., 2020).

$$C_{16}H_{34} + 12.25 \text{ SO}_{4}^{2-} \rightarrow 16 \text{ HCO}_{3}^{-} + 12.25 \text{ HS}^{-} + 3.75 \text{ H}^{+} + \text{H}_{2}\text{O} (3)$$

$$4 \text{ C}_{3}H_{8} + 5 \text{ SO}_{4}^{2-} \rightarrow 6 \text{ HCO}_{3}^{-} + 5 \text{ HS}^{-} + 4 \text{ H}_{2}\text{O} - \text{H}^{+}$$
(4)

Even though seeps dominated by heavy hydrocarbons (like cruce oil) are less common than seeps dominated by light hydrocarbons (like methane), their occurrences are well documented globally (e.g., Hornafius et al., 1999; Noble et al., 2009 Va entine et al., 2010; Jones et al., 2014; Körber et al., 2014; Jiang et al., 2018) and suggest an important effect on microbial diversity and associated biogeochemical cycling (Joye et al. 20<sup>14</sup>, Orcutt et al., 2010; Sahling et al., 2016). For example, GoM hydrocarbon seeps hare been shown to supply subsurface-derived microbial populations, biomass, and metabolic potential along with the geofluids to deep ocean (Chakraborty et al., 2020), significaring inspacting the biogeochemistry of shallow sediments (Hovland et al., 2012; Coffin et al., 2013; Rowe, 2017) as well as the overlying water column (D'souza et al., 2016). Oil seeps annost certainly have persisted over geologic time (Wilson et al., 1974) and are relevant to o can chemistry over the same time scales. Characterizing and comparing the impacts o cru le oil and methane seeps in present-day settings will enable better identification of these processes in the sediment record and their biogeochemical implications over geologic time (Peckmann and Thiel, 2004; Campbell, 2006; Bristow and Grotzinger, 2013). Furthermore, hydrocarbon seeps are highly sensitive in response to changes in oceanographic and tectonic conditions (e.g., Aharon et al., 1997; Berndt et al., 2014; Oppo et al., 2020), and our current understanding of their temporal variation is weak. Such uncertainty also leaves a critical gap in our ability to assess the potential response of these seep systems during future climate change.

In this study, we examined authigenic carbonates using carbon ( $\delta^{13}C_{CaCO3}$ ), oxygen ( $\delta^{18}O_{CaCO3}$ ), and sulfur isotopes ( $\delta^{34}S$  of chromium reducible sulfur – CRS) along with the carbonate-

associated sulfate (CAS) to examine the geological signatures of methane and crude oil oxidation (e.g., Eq. 3). CRS represents total reduced inorganic sulfur (pyrite S + acid-volatile sulfide S + elemental S), with the exclusion of organic S and sulfate phases (Canfield et al., 1986). This fraction is typically but not always dominated by pyrite in most sediments (Lyons, 1997; Rickard et al., 2017). We focused our attention on the authigenic carbonates recovered from asphalt seep sites in Chapopote Knoll of Campeche Bay, which were suggested to be dominantly derived from crude oil oxidation. The study site is characterized by extensive and serial asphalt flows, oil and gas seeps, and seafloor gas hydrate deposits, along with seep-associated chemosynthetic communities and authigenic carbonate deposits (MacDonald et al., 2004; Brüning et al., 2010; Sahling et al., 2016). These discoveries added a new dimens on o the inventory of seafloor hydrocarbon seep processes (Bohrmann, 2014; Marcon et al. 2018). Given this importance, we compared the results from Chapopote seep carbonates with h ultiple seep sites from the northern Gulf of Mexico, many of which are methane-dominate.' seeps devoid of oil. As part of this study, we determined U-Th dates for the Chapopote seep carbonate to identify the timing and mechanisms of formation.

#### 2. Study Area

The GoM basin is characterized by multiple, and often large hydrocarbon reservoirs commonly overlying salt deposits, which a sealed by overlying continental margin sediments. Differential sedimentary loading an ( a nsity contrast with the overburden induce salt diapirism and consequent fault generation, paving the way for hydrocarbon leakages toward seafloor (Brooks et al., 1990; Sassen et al., 1993; Roberts, 2001; Fisher et al., 2007; Kennicutt, 2017). The southern GoM is a relatively unexplored area with numerous hydrocarbon seeps and is characterized by two distinct active salt provinces: the Campeche and Sigbee Knolls, separated from the Mississippi-Texas-Louisiana salt province in the northern GoM by the Sigsbee Abyssal Plain (Bryant et al., 1991; Fig. 1). These knolls consist of a series of domes and ridges formed by movement of the Jurassic salt deposits underlying the ~5 to 7 km thick continental margin sediments (Salvador, 1991; Ding et al., 2008). Extensive hydrocarbon transport from sediments to the sea-surface in the region is observable via satellite images as sea-surface oil slicks (MacDonald et al., 2004; MacDonald et al., 2015; Suresh, 2015; Römer et al., 2019).

This study employs carbonate samples collected at a water depth of 2902 m from Chapopote Knoll ( $21^{\circ}54'N/93^{\circ}26'W$ ) located on the northern slope of the Campeche Knolls province during the *R/V Sonne* cruise SO174/2 in 2003 (Bohrmann and Schenck, 2004) and *R/V Meteor* cruise M67/2 in 2006 (Bohrmann et al., 2008). The Campeche Knolls province consists of a cluster of elongated knolls and ridges formed via salt tectonics (Garrison and Martin, 1973). The knolls have relief ranging from 450 to 800 m above the seafloor in water depths of 3000 to 3500 m (Ding et al., 2008). A gravity core GeoB 10602 collected about 50 m away from the asphalt flow, which did not contain any oil, asphalt, or CaCO<sub>3</sub> crusts serves as non-seep background sample for Chapopote seep site (Fig. 1C).

Extensive and overlapping lava-like asphalt flows at Chaport's unanate from a central craterlike depression, generating extensive surface deposits of coluined asphalt with distinct surface textures—leading to the term 'asphalt volcano' (M cD ald et al., 2004). Seismic studies revealed that asphalt seeps derive from a large reservoir buried at shallow depths linked to a deep, heavy petroleum source (Ding et al., 2008. L'ng et al., 2010). Asphalt flow is accompanied by oil and gas seeps (Bohrmann, 2008) and station gas hydrate deposits (Klapp et al., 2010), along with seep-associated chemosymbiotic communities and authigenic carbonate deposits (Bohrmann and Schenck, 2004; MacDanal et al., 2004; Naehr et al., 2009; Brüning et al., 2010; Sahling et al., 2016). Seepage system: support prolific microbial activity involving crude oil degradation and sulfate reduction (Cohubotz et al., 2011a; Schubotz et al., 2011b) in otherwise deep sea pelagic sediments with low organic carbon input from surface waters. Surface sediments from Chapopote, re characterized by locally higher total organic carbon contents (0.9%) and very high calon-nitrogen atomic ratios (C/N<sub>a</sub>) compared to the surrounding abyssal sediments (ratios of 50 vs. 7), suggestive of organic carbon sourced from hydrocarbon seepage rather than deposition (Escobar-Briones and García-Villalobos, 2009). Authigenic carbonates collected from Campeche seeps allow us to investigate the role of crude oil oxidation during carbonate authigenesis (Naehr et al., 2009; Smrzka et al., 2016).

For comparison, we also analyzed samples from relatively well-studied sites of northern continental slope of GoM Green Canyon Block 415 (GC 415, 27°33.48N/90°58.86W, water depths 950m; 27°32.61N/90°59.54W, water depth 1045m) and Bush Hill (GC 185, 27°46.97N/91°30.47W, water depths 547m). These sites were also among the expedition targets

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of R/V Sonne cruise 174 (Bohrmann and Schenck, 2004; Fig. 1). Similar to Campeche Knolls, these sites host widespread hydrocarbon seeps due to fractures in sedimentary strata induced by salt tectonics (Roberts and Aharon, 1994; Sassen et al., 2004; Feng et al., 2009). Our preliminary analysis along with previous reports revealed a noticeable contribution of crude oil oxidation to the DIC pool at GC 185 and methane oxidation to DIC pool at GC 415 (Sassen et al., 1994; Akam et al., 2019). Further, we used published literature reports of C-S isotope values for seep carbonates collected from multiple sites in the northern GoM (Atwater Valley - AT340, Green Canyon – GC 180, GC 232, GC 234, GC 852 Garden Banks – GB 260, GB 382, GB 427, GB 647, and Mississippi Canyon MC 118, Fig. 1) (Formolo and Lyons, 2013; Feng et al., 2016; Sun et al., 2020). Among these sites, GC 232 serves as a representation of a crude oil-dominant site (Sun et al., 2020), and other sites represent either methane-dc min int settings or a mix of methane and crude oil seepage. Near-surface sediments (0-1° ch below seafloor) collected from approximately 2.6 km north of a brine pool at GC233 a. 4 away from any visible hydrocarbon seeps serve as a background site for the northern JoM (Formolo and Lyons, 2013). Table 1 summarizes the data sources for this study We iso emphasize that all the samples analyzed in this study and retrieved from literature for pmparison are from surface/near-surface sediments of the seafloor, thus placing our emphasis on the earliest diagenetic processes.

Data Source	Site	Approximate Location	Water depth (m)	Seep type
	Chi poj ote	21°54.0N,	2002	Crude oil
	ínoll	93°26.40W	2902	dominant
This study	Busn Hill GC	27°46.97N,	F 4 7	Crude oil
	185	91°30.47W	547	dominant
	CC 415	27°32.61N,	1045	Methane and
This study	GC 415	90°59.54 W	1045	crude oil
	CC 415	27°33.48N,		Methane and
	GC 415	90°58.86W	950	crude oil
	Chapopote	21°54.01N,	2004	Nessee
	Background	93°26.24W	2884	No seep
Feng et al.	47240	27°38.8N,	2210	Methane
(2016) and Sun	AI 340	88°21.9W	2210	dominant

Table 1: Loca	ation, water	depth, and	leep .ype o	of the samples	studied.
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et al. (2020)	CD2CD	27°42.5N,	502	Methane
	GB200	91°58.5W	503	dominant
	CD427	27°35.50N,	050	Methane and
	GB427	92°25.94W	803	crude oil
	CDC47	27°20.50N,	050	Methane and
	GD047	92°24.18W	950	crude oil
	CC140/194	27°49.16N,	207	Methane and
	GC140/184	91°31.95W	287	crude oil
	CC190	27°28.2N,	252	Methane and
	GC180	91°18.0W	233	crude oil
	ccasa	27°44.50N,	E 27	Crude oil
	GC232	91°19.28W	5.7	dominant
	66229	27°44.45N,	729	Methane
	GC238	91°03.05W	128	dominant
	66053	27°6.36N,	1622	Methane
	66852	91°9.97' <i>N</i>	1055	dominant
	NAC110	27°∩6.3N,	005	Methane and
	IVIC118	9. °0′J.8W	665	crude oil
	CC 224	27°44.79N,	FFO	Methane and
Formolo and	GC 234	91°13.33W	550	crude oil
Lyons (2013)	Northern GoM	?'/°43.42N,		No coop
	Background	91°19.09W	716	No seep



Figure 1: A) Map of the Gulf of Mexico (GoM) showing locations of the study site at Chapopote Knoll and other sites considered within this study within the northern GoM (GC 415 and GC 185). Significant asphalt seepage occurs at Chapopote Knoll whereas the northern GoM sites show a mixture of methane and oil seepage. The green dots indicate locations from published literature where we used those C-S isotopic data for comparison with that from Chapopote Knoll. B) A magnified view of northern GoM sites for an overview of bathymetry. C) A magnified view of Chapopote knoll with 25 m bathymetric contours modified from Naehr et al. (2009). This study uses seep carbonates collected from TVG-6 site (red dot) and background sediment

samples collected from GeoB 10602 (dark gray dot). Light gray shading represents an approximate extent of asphalt flow.

#### 3. Methods

Carbonate samples were collected from surface sediments via video-guided grab sampling. Authigenic carbonate crusts and shell material from chemosynthetic clams were separated from grab samples. Specific samples were defined based on the dive number and location. Fifteen carbonate crusts were analyzed, which included ten crusts from Chap opote (different pieces from three large grab samples, sample ID TVG6 22 -30), four it egu'ar tabular crusts from GC 185 (sample ID GC 185N1-3, N10), and a tabular concretion nom GC 415 (sample ID GC 415T). These materials were split into multiple subsamples based on observations from hand specimens. Carbonate content of the samples were determined by acid-leaching/weight-loss procedure and are reported as weight percent CaCO<sub>3</sub>. Bulk 1, ineralogy was determined by X-ray diffraction (XRD) according to Naehr et al. (2000) a. T MUCC. One gram of crushed sample was mixed with 0.25 gm internal corundum stan.<sup>1</sup>ard ( $\alpha$ -Al<sub>2</sub>O<sub>3</sub>) to prepare randomly-oriented powdered slides. Scans were run from 20°- 60 using a Rigaku Ultima III X-ray diffractometer at TAMUCC at a scan speed of  $0.01^{\circ}$  20's. Relative proportion of magnesium content in carbonate was determined using the shift is spacing of the reflection (104) (Greinert et al., 2001). Calcite with  $MgCO_3 < 4\%$  is referred to us low-Mg calcite, and 4-30% were considered high-Mg calcite (Flügel, 2004). Polished thin sections 50 x 75 mm in size, partially stained with alizarin red and potassium ferrocyanide. v ere used for textural and compositional analysis via standard optical microscopy. Stable isotopes of carbon and oxygen from authigenic carbonates were determined using a ThermoScientific Gasbench Device coupled to a Delta V Advantage Isotope Ratio Mass Spectrometer (IRMS) via a ConFlo IV inlet at the University of California, Riverside. Sample powders were microdrilled from polished slab surfaces for bulk micritic carbonate subsamples (microdrilled samples for each crust were named as A-D in Table 2). Carbon dioxide for  $\delta^{13}C_{CaCO3}$  and  $\delta^{18}O_{CaCO3}$  analyses was produced by reaction of samples with 103% orthophosphoric acid. Precision of  $\delta^{13}C_{CaCO3}$  and  $\delta^{18}O_{CaCO3}$  measurements is 0.2%. The  $\delta^{13}C_{CaCO3}$ and  $\delta^{18}O_{CaCO3}$  are reported with reference to Vienna Pee Dee Belemnite (VPDB) standard.

U/Th analyses were performed on samples micro-drilled from polished samples targeted for early diagenetic, particularly micritic, cements. Thirty samples were initially screened for  $^{238}$ U/ $^{232}$ Th ratios to determine whether they could be dated by U/Th methods. One milligram of sample was dissolved in 1.5 mL 0.5 M nitric acid, and U, Th, and Ca were measured by quadrupole ICP-MS at the MIT Center for Environmental Health Sciences to determine U concentrations and U/Th ratios. A gravimetric U/Th/Ca solution with similar abundances to the samples was used to calibrate the instrument, and a standard was run after every 15-20 samples to monitor instrument drift. Samples with <sup>238</sup>U/<sup>232</sup>Th mass ratios greater than two were selected for U/Th dating analysis. Radioactive U/Th isotope dating of the carbonate crusts was performed Nu Plasma II-ES MC-ICP-MS at the Department of Earth, Atm spheric and Planetary Sciences, Massachusetts Institute of Technology (MIT). Micro-drillea carbonate powder samples ranging in weight from 10 to 70 mg were used. Samples were dirsolved in 8 N HNO<sub>3</sub> and spiked with a <sup>229</sup>Th/<sup>233</sup>U/<sup>236</sup>U tracer. Uranium and Th were separated following procedures described by Edwards et al (1987) and Bayon et al., (2009). Faw data were corrected with an initial  $^{230}$ Th/ $^{232}$ Th atomic ratio of 4.4 ± 2.2 x 10<sup>-6</sup> ... surving a typical upper continental crust value for  $^{238}$ U/ $^{232}$ Th, and the errors were arbitrarily  $\therefore$  sumed to be 50%. This initial  $^{230}$ Th/ $^{232}$ Th ratio is consistent with values used in some provi studies of seep carbonates from the GoM and other areas (Feng et al., 2010; Chen et al., 20.9), but we note that Aharon et al. (1997) used initial  $^{230}$ Th/ $^{232}$ Th ratios ranging from 8-  $^{22}$  x 10<sup>-6</sup>. Use of a higher initial  $^{230}$ Th/ $^{232}$ Th ratio would result in younger corrected ages; we note the need for future work to better constrain this ratio at our site.  $\delta^{234}U_{initial}$  was calculated based on <sup>230</sup>Th age (T), that is,  $\delta^{234}U_{initial} = \delta^{234}U_{measured} \times e^{\lambda 234*T}$ , and T is the corrected . re. Decay constants for <sup>230</sup>Th and <sup>234</sup>U are from Cheng et al. (2013): the decay constant for  ${}^{238}$ U : 1.55125 x 10<sup>-10</sup> yr<sup>-1</sup> (Jaffey et al., 1971).

Carbonate associated sulfate (CAS) was extracted on micro-drilled samples targeted for micritic and early diagenetic cements using a technique modified from Lyons et al (2004). This method includes removal of initial sulfate salts (before acid addition) with a NaCl solution and treatment for sulfides and organically bound sulfur (e.g., ester bound sulfates) using a 5% sodium hypochlorite rinse. Furthermore, 5% (by weight) SnCl<sub>2</sub> was used to inhibit pyrite oxidation during the 4N HCl treatment. Despite these precautions, our attempt to measure CAS in Chapopote samples failed, yielding many highly negative  $\delta^{34}S_{CAS}$  values rather than the positive values from seawater and as modified via microbial sulfate reduction. We attribute these

problems to extremely low CAS concentrations (as low as 31 ppm) and concomitant vulnerability to any sulfide oxidation during extraction, as well as residual sulfate derived from sulfide during sample collection, storage, or naturally prior to collection. Thus, our CAS data are not valid because of extremely low concentrations and are not reported.

Chromium reducible sulfur (CRS) from bulk micritic carbonate samples was extracted using a modified method of Canfield et al (1986). Homogenized 5 g samples were reacted with 10 ml ethanol, 25 ml of 6M HCl, and 25 ml of a chromium chloride (1 M CrCl<sub>3</sub>.6H<sub>2</sub>0 in 0.5 M HCl) solution. CRS was converted to H<sub>2</sub>S gas and carried via a N<sub>2</sub> ca rier gas to an AgNO<sub>3</sub>-NH<sub>4</sub>OH trap where it was quantitatively converted to silver sulfide ( $\Delta_2 \circ_5$ ). Ag<sub>2</sub>S precipitates were recovered on polycarbonate membrane filters using vacuur. Function before being dried and weighed to determine the CRS weight percent.  $\delta^{34}S_{CRS}$  v due, were measured using a Thermo Scientific Delta V Plus IRMS connected to a Costech 4 )10 ECS via a ConFlo III interface at the University of California, Riverside. Standard deviatior for  $\zeta^{34}S$  analysis was ± 0.23‰ (VCDT).

A cluster analysis of the  $\delta^{13}C_{CaCO3}$  and  $\delta^{34}S_{CRS}$  data pairs was performed on data obtained from this study along with the data collected from published literature from additional sites mentioned in Table 1 (Supp. Table 1) via *k*-means unsupervised learning algorithm using Python. This approach allowed an additional tool to pressify the  $\delta^{13}C_{CaCO3}$  and  $\delta^{34}S_{CRS}$  data pairs, along with classification using seep types mentioned in Table 1. K-means divides objects into clusters that show similar characteristics and are dissimilar to the objects belonging to another cluster. The elbow method, which fits the models with a range of values of k (number of clusters), was used to determine the optimum value of k. Visual inspection of spread of the data and the site characteristics were also considered here to choose the optimal k value. Once k is determined, each data point is assigned to a cluster based on minimizing the distance between the points within a cluster. The final field boundaries to classify the  $\delta^{13}C_{CaCO3}$  and  $\delta^{34}S_{CRS}$  data pairs were delineated by combining cluster density and site characteristics (seep type, Table 1).

#### 4. Results

#### 4.1 Petrography

Petrographic observations of carbonate macro- and microfacies from Chapopote seep carbonates were previously reported by Canet et al., (2006) and Naehr et al (2009). Aragonite was the primary phase for Chapopote seep carbonates. The siliciclastic components consisted of detrital quartz, feldspar, and clay minerals. Petrographic observations of the samples showed wide-ranging textural variations (Figure 2). Aragonite cement types included acicular, botryoidal, peloidal, micritic, and sparitic cement. Microsparitic aragonite was the volumetrically dominant phase, commonly occurring as peloidal and clotted textures. Aragonite crystals were often coated with crude oil residues. Macroscopically, the samples consisted of *cl* irregularly-shaped large clasts that were held together by a microsparitic to sparitic angonite components (primarily bivalve and foraminifera shells). Primary pore space was vide spread resulting from incomplete filling of voids between clasts and microsparitic or fibuous cements (Fig. 2B). Secondary pores, potentially caused by carbonate dissolution and *in situ* brecciation, were also observed (Fig. 2A). Framboidal pyrite was pervasive (Fig. 2a)



Figure 2: Petrographic observations on Chapopote seep carbonates. All images are in planepolarized light, and blue color indicates pore spaces. A) Multiple cement types including peloidal (P), Sparitic (S), and micritic (M) cement. Note fracture (arrow) post-dating cement formation. B) Banded and botryoidal aragonite (B) that also exhibits fibrous and needle-like

crystals. C) Sediment (S) cemented by microcrystalline and peloidal aragonite. Yellow box is enlarged in D. D) Zoomed view of yellow box of C. Microsparite (MS) filling of spaces between sediments and cement surrounding the clasts is incomplete, leaving significant pore space. Aragonite cement fills pore spaces, creating a peloidal texture in the left half of the picture. A bivalve shell fragment (black arrow) is visible. E) Peloidal texture suggestive of microbial activity and early diagenetic cementation. A shell fragment (yellow arrow) and oil coatings (white arrows) are visible. Oil coating (white arrows) surrounding the aragonite cement can be seen. F) Pyrite clumps (black arrow) composed of pyrite framboids approximately 15 to 20 µm in size is suggestive of sulfate reduction. Clots of residual hydrocarbox s (white arrow) can be seen around the pore spaces adjacent to cemented structures as w<sup>2</sup>ll (s coating on the aragonite cements.

#### 4.2 U-Th dates

Of the 30 samples screened for U-Th dating, nine Chapopote samples had  $^{238}$ U/ $^{232}$ Th greater than 2, and are thus suitable for age-dating. These samples were analyzed for U/Th isotopic composition and age dated (Table 2). The  $^{238}$ U concentrations for these selected samples ranged from 5230 to 13200 ppb, and  $^{232}$ Th concentrations ranged from 482 to 1220 ppb.  $\delta^{234}$ U initial values averaged 143‰ when corrected for detrital U (based on an initial  $^{230}$ Th/ $^{232}$ Th atomic ratio of 4.4 ± 2.2 x 10<sup>-6</sup> and assuming that  $^{230}$ Th/ $^{238}$ U and  $^{234}$ U/ $^{238}$ U activity ratios in detrital matter are 1 ± 0.1). The initial  $\delta^{234}$ U are soft the carbonate fraction of the Chapopote samples is 147 ± 1‰, consistent with the average seawater composition of 146.8 ± 0.1‰ (Andersen et al., 2010). U-Th dates ranged from 4.6 ± 1.0 ka to 13.5 ± 2.1 ka (BP).  $^{230}$ Th/ $^{230}$ Th large. As a result, uncertainties in the corrected ages are dominated by the uncertainty in the initial  $^{230}$ Th correction.

Table 2: U-Th data from Chapopote samples

Sample	<sup>238</sup> U	<sup>232</sup> Th	$^{230}$ Th/ $^{232}$ T	δ <sup>234</sup> U	<sup>230</sup> Th/ <sup>238</sup>	<sup>230</sup> Th Age	<sup>230</sup> Th	$\delta^{234}$ U
ID	±	±	h	(‰)	U	(ka BP)	Age (ka	initial

	$(2\sigma)^{a}$	$(2\sigma)^{a}$	± (2σ)	$\pm (2\sigma)^{b}$	± (2σ)	$\pm (2\sigma)^{c}$	BP)	(‰) ±
						<b>x</b> ,	$\pm (2\sigma)^{d,e}$	(2 <b>σ</b> ) <sup>1</sup>
	(nnh)	(nnh)	(atomic	(measured	(activity)	(uncorrecte	(corrected	(corrected
	(ppb)	(bbp)	×10 °)	)	(activity)	d)	)	)
	1315	118			0.1074			
TVG_6	$0 \pm$	$3 \pm$	$19.0 \pm$		±	$10.788 \pm$	8.32 ±	
-21a	260	24	0.3	$143 \pm 2$	0.0017	0.19	1.2	$146 \pm 2$
		106			0.1340			
TVG_6	6673	$0 \pm$	$13.4 \pm$	$138 \pm$	土	$13.750 \pm$	$9.40 \pm$	
-23a	± 133	21	0.3	0.3	0.0029	0.22	2.2	$141\pm3$
		110			0.1109			
TVG_6	7253	4 ±	11.6 ±	$138 \pm$	±	$11.260 \pm$	$7.10 \pm$	
-25a	± 145	22	0.3	0.3	0.002s	0.31	2.1	$140\pm2$
		116			0.1(91			
TVG_6	7799	$3 \pm$	$18.0 \pm$	$139 \pm$	±	$17.670 \pm$	$13.50 \pm$	
-26a	±156	23	0.3	0.3	(1),028	0.33	2.1	$143\pm2$
		101			0.1673			
TVG 6	5231	$8 \pm$	$13.7 \pm$	137 +	±	$17.460 \pm$	$12.10 \pm$	
-26c	± 105	20	0.3	0.3	0.0036	0.41	2.8	$141\pm3$
		107		0	0.1235			
TVG 6	7736	$4 \pm$	14.1 +	$142 \pm$	±	$12.630 \pm$	$8.79 \pm$	
-27a	± 155	22	0.5	0.3	0.0026	0.29	1.9	$144 \pm 2$
					0.0659			
TVG 6	6824	422	$1.1.8 \pm$	$143 \pm$	±	$6.507 \pm$	$4.55 \pm$	
-27c	± 482	$\pm 10$	0.3	0.3	0.0014	0.15	0.96	$144 \pm 2$
					0.1612			
TVG 6	5990	993	$15.4 \pm$	$140 \pm$	±	$16.700 \pm$	$12.10 \pm$	
-28a	± 120	$\pm 20$	0.3	0.3	0.0031	0.35	2.3	$144 \pm 2$
	1016	122			0.0864			
TVG 6	$0 \pm$	$1 \pm$	$11.4 \pm$	$142 \pm$	±	$8.630 \pm$	5.32 ±	
-29d	200	24	0.3	0.3	0.0022	0.23	1.67	$144 \pm 2$

Notes:

Decay constants for <sup>230</sup>Th and <sup>234</sup>U are from Cheng et al. (2013); decay constant for <sup>238</sup>U is  $1.55125 \times 10^{-10} \text{ yr}^{-1}$  (Jaffey et al., 1971).

<sup>a</sup>Reported errors for  $^{238}$ U and  $^{232}$ Th concentrations are estimated to be  $\pm 1\%$  due to uncertainties in spike concentration; analytical uncertainties are smaller.

 ${}^{b} \delta^{234} U = ([{}^{234} U/{}^{238} U]_{activity} - 1) \times 1000.$  ${}^{c} [{}^{230} Th/{}^{238} U]_{activity} = 1 - e^{-\lambda 230T} + (\delta^{234} U_{measured}/1000) [\lambda_{230}/(\lambda_{230} - \lambda_{234})] (1 - e^{-(\lambda 230 - \lambda 234)T}), \text{ where } T \text{ is the age.}$ "Uncorrected" indicates that no correction has been made for initial <sup>230</sup>Th.

<sup>d</sup>Ages are corrected for detrital <sup>230</sup>Th assuming an initial <sup>230</sup>Th/<sup>232</sup>Th of (4.4  $\pm$  2.2) x 10<sup>-6</sup>.

<sup>e</sup>B.P. stands for "Before Present" where the "Present" is defined as Janu. "71, 1950 C.E.  $^{f}\delta^{234}U_{initial}$  corrected was calculated based on  $^{230}$ Th age (T), i.e.,  $\delta^{234}U_{ir} = \delta^{234}U_{measured} \times e^{\lambda 234*T}$ , and T is corrected age.

#### 4.3 C-O and S isotopes

Our measured values of  $\delta^{13}C_{CaCO3}$ ,  $\delta^{18}O_{CaCO3}$ , and  $\delta^{34}S_{C_4}$ , are shown in Table 2.  $\delta^{13}C_{CaCO3}$  and  $\delta^{18}O_{CaCO3}$  values from Chapopote samples average 1 -25‰ and +4.5‰, respectively. Samples from the northern GoM, GC 185 and GC 4<sup>15</sup>, related average  $\delta^{13}C_{CaCO3}$  values of -19.8‰ and 4.4‰ and  $\delta^{18}O_{CaCO3}$  values of -34.6‰ and +4.8‰, respectively (Fig. 3A; Table 2).  $\delta^{34}S_{CRS}$ values from Chapopote carbonates were noticeably low, ranging from -14% to -38.7%, with an average value -27.4‰. Samples from CC 415 and GC 185 averaged -4.8‰ and +18.9‰, respectively. Data for additional cites compiled from literature is summarized in supplementary Table 1. Overall, the  $\delta^{13}C_{CaCO_2}$ ,  $\delta^{18}O_{CaCO_3}$ , and  $\delta^{34}S_{CRS}$  data showed noticeable spread within and between the sites (Fig. 3B-D).

![](_page_19_Figure_1.jpeg)

Figure 3: A) Average values of  $\delta^{13}C_{CaCO3}$ ,  $\delta^{18}O_{CaC3}$ ,  $\delta^{18}O_{CaC3}$ , and  $\delta^{34}S_{CRS}$  obtained from samples in this study. B-D: Box and whisker plot for  $\delta^{13}C_{LaC03}$  ( $\Gamma$ 'g. 3B),  $\delta^{18}O_{CaCO3}$  (Fig. 3C), and  $\delta^{34}S_{CRS}$  (Fig. 3D) from seven sites (Chapopote, GC 232, CC 415, GC 234, GC 852, Chapopote Background and northern GoM background) that shared 83% of the total data in this study which includes new data and that compiled from literature. Refer to Table 1 for site descriptions. Note that the near-surface (<10 cmbsf) background samples from Chapopote and northern GoM did not yield measurable CRS content.

Table 2:  $\delta^{13}C_{CaCO3}$ ,  $\delta^{18}C_{CaCO3}$ , and  $\delta^{34}S_{CRS}$  values, with chromium reducible sulfur (CRS) and calcium carbonate (CaCO<sub>3</sub>) content from studied samples. Sample materials involve authigenic carbonate crusts, shell material at seep sites and surface sediments of background sites. An extended database, which includes a compilation of published literature data from multiple seep sites in GoM, is provided in Supplementary Table 1.

Location	Wat er dep th (m)	Sample ID	CaC O <sub>3</sub> (wt %)	CRS (wt %)	δ <sup>13</sup> C <sub>Ca</sub> <sub>co3</sub> (‰ VPDB )	δ <sup>18</sup> O <sub>Ca</sub> <sub>CO3</sub> (‰ VPDB)	δ <sup>34</sup> S crs (‰ VCD T)	Mineral ogy
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				0.01				Aragonit
		I VG-6 22A	96%	%	-25.0	4.2	-20.1	e
			000/	0.05				Aragonit
		TVG-6 22B	90%	%	-24.8	3.8	-26.2	e
			070/	0.06				Aragonit
		TVG-6 22C	97%	%	-24.5	4.6	-19	e
			010/	0.00				Aragonit
		TVG-6 22D	91%	%	-24.1	4.4	-18.8	e
			0.40/	0.12				Aragonit
		TVG-6 23A	84%	%	-27.6	5.7	-28	e
			0.00/	0.17				Aragonit
		TVG-6 23B	88%	%	-26.2	3.8	-23.7	e
			700/					Aragonit
		IVG-023C	/0%	trace	-26.7	3.7	-22.0	e
			070/	0.36				Aragonit
		I VG-0 24B	8/%	%	-25.J	5.9	-27.3	e
		TUC ( )4C	950/	0.17				Aragonit
		I VG-6 24C	85%	%	-26.3	4.4	-28.5	e
		TVG-6 24D-	700/	0.(0				Aragonit
	2902	SHELL	/0%	355	-4.1	3.3	-27.5	e
			0,%	<b>U.21</b>				Aragonit
Chapopote		1 VG-0 25A		%	-24.2	4.1	-27.6	e
Knoll		TVG-6 25P	71.74	0.53				Aragonit
$21^{\circ}54.0$ IN,			70.0	%	-25.7	5.1	-24.4	e
95 20.40W		TVG 62 A	000/	0.08				Aragonit
		1 VG-0 20P.	99%	%	-26.5	4.3	-21.7	e
		TVC 6 26P	<b>Q10</b> /	0.26				Aragonit
			81%	%	-25.0	5.3	-21.6	e
		CVG-626C	83%	0.16				Aragonit
			0370	%	-25.6	4.3	-28.5	e
		TVG-6 27A	87%	0.08				Aragonit
		1V0-027A	0770	%	-26.0	4.0	-34.8	e
		TVG-6 27B	75%	0.38				Aragonit
		1100210	1570	%	-28.5	4.5	-33.4	e
		TVG-6 27C	87%	0.14				Aragonit
		1100270	0770	%	-23.3	4.3	-33.4	e
		TVG-6 28A	64%	0.06				Aragonit
		110 0 2011	0170	%	-26.3	4.0	-31.3	e
		TVG-6 28B	89%	0.02				Aragonit
		1,00200	0770	%	-25.8	4.7	-38.4	e
		TVG-6 28C	71%	0.02				Aragonit
		1,00200	/1/0	%	-25.2	4.4	-26.5	e
		TVG-6 28D-	96%	0.07				Aragonit
		SHELL	2070	%	-23.0	4.3	-33.1	e
		TVG-6 29B	96%	0.21	-25	4.6	-27.4	Aragonit

				%				e
		TVG 6 20C	0004	0.48				Aragonit
		1 0 -0 290	9970	%	-27.4	4.9	-38.7	e
		TVG-6 29F	91%	0.01				Aragonit
		110020	2170	%	-26.9	3.9	-34.9	e
		TVG-6 30A	88%	0.47	26.0	56	276	Aragonit
				% 1.04	-20.8	3.0	-27.0	Aragonit
		TVG-6 30B	85%	1.04	-29.4	51	-14 1	e
		GeoB BGA		trace	0	0.09	-	NA
Chapopote		GeoB BGB		trace	1	0.07	_	NA
Background	2884	GeoB BGC	33%	trace	-1	0.05	_	NA
$21^{\circ}54.01N$ ,		GeoB BGD		trace	0.0	$\frac{0.02}{0.04}$	_	NA
95 20.24 W		GeoB BGE		trace	015	0.10	_	NA
		GC 185	0.004	0.23		0110		
		N10A1	80%	%	20.9	5.2	-18.5	HMC
GC 185		GC 185	84%	0.6.7	D			
27°46.97N.	547	N10A2	0470	%	-21.4	4.6	-18.5	HMC
91°30.47W		GC 185	71%	0.20	16.2	4.0	24.2	
		INTUBI CC 185	51 %	/0	-10.3	4.2	-24.2	HMC
		N10B2		0.44	-20 5	38	-14 5	HMC
				0.17	20.0	210	1110	mile
		GC 415 NI-A	69%	%	-32.0	5.8	-7.4	HMC
		CC 415 N1 9	83%	0.44				
			0370	%	-27.4	5.0	-4.7	HMC
GC 415		GC 415 N2-A	74%	0.48	21.2	5.0	11.2	
27°32.61N,	1045			% 0.74	-31.3	5.0	-11.5	HMC
90°59.54 W		CC 115 N2-B	74%	%	-39.3	4.9	-10.9	HMC
			6004	0.07	57.5		10.9	Inde
		GC 415 N3-A	68%	%	-24.3	4.7	2.3	HMC
		GC 415 N3-B	62%	0.10				
		0C 415 N3-D	0270	%	-21.3	5.0	3.4	HMC
		GC 415 T-1	90%	N/A	-35.14	5.09	N/A	HMC
		GC 415 T-2	85%	N/A	-34.30	5.05	N/A	HMC
		GC 415 T-3	91%	N/A	-37.78	5.40	N/A	HMC
GC 415		GC 415 T-4	89%	N/A	-36.73	3.70	N/A	HMC
27°33.48N,	951	GC 415 T-5	92%	N/A	-32.97	4.38	N/A	HMC
90°58.86W		GC 415 T-6	96%	N/A	-34.12	4.28	N/A	HMC
		GC 415 T-7	91%	N/A	-34.29	3.90	N/A	НМС
		GC 415 T-8	88%	N/A	-37.60	3.94	N/A	HMC
		GC 415 T-9	91%	N/A	-42.3	4.9	N/A	НМС

GC 415 T-10	92%	N/A	-42.8	5.0	N/A	HMC
GC 415 T-11	86%	N/A	-40.6	4.9	N/A	HMC
GC 415 T-12	86%	N/A	-41.0	4.9	N/A	HMC
GC 415 T-13	86%	N/A	-33.1	4.9	N/A	HMC

N/A = not available

Cluster analysis using k-means method, applied on  $\delta^{13}C_{CaCO3}$  and  $\delta^{34}S_{CRS}$  data produced in this study and those from published reports, suggest an optimal k between 3 to 4, after which the cost function decreases by very small amounts with each iteration (Fig. 4). Visualization of data with the help of cluster analysis and site characteristics suggests for r sr mple groups with distinctly different DIC and sulfide sourcing inferred from the paired  $\delta^{-3}C_{C,CO3}$  and  $\delta^{34}S_{CRS}$  values:

- (i) samples from methane seepage sites showing AOM as the dominant diagenetic process
- (ii) samples from oil seepage sites that st ov A JNM as a dominant process
- (iii) samples from organoclastic sul<sup>*f*</sup> ate eduction (OSR) dominated setting
- (iv) samples showing a mixed sources for DIC and sulfide showing values intermediate between the other three enclanearbers.

The decision boundaries of end-numbers were loosely defined based on cluster density and site characteristics (seep type, Table 1). While AOM and AONM end-members were clearly distinguishable, the boundary between AONM and OSR as well as OSR and mixed pool were less distinct. Section 5.4 discusses these results in detail.

![](_page_23_Figure_1.jpeg)

Figure 4: Optimum number of clusters dete. nined by elbow method from unsupervised k-means learning algorithm on  $\delta^{I3}C_{CaCO3}$  and  $\delta^{3*}\mathcal{J}_{CRS}$  data produced in this study and that from published data.

#### 5. Discussion

Our results offer a broader evaluation of the sulfate reduction processes that contribute to the DIC pool for carbonate auth genesis on the GoM seafloor and their geochemical distinctions based on C-S isotopes. S afloor and petrographic observations suggest multiple seepage events at Chapopote Knoll and that much of this biogeochemical activity occurs not only interstitially but also endolithically, where microbes inhabit the carbonate crust interiors. We were also able to produce the first U-Th based dates for authigenic carbonates from the southern GoM.

## 5.1 Petrographic Observations – Endolithic Biosignatures

Petrographic observations suggest precipitation of authigenic carbonate and sulfide minerals in a dynamic seep setting with episodic fluid flow, microbial activity, and associated cycling of C, S, and Fe (Figs. 2 and 5). Peloidal micrite exhibiting a clotted fabric (Fig. 2E) has been reported widely from seep carbonates and is indicative of early-stage cementation resulting from

microbial interactions (Cavagna et al., 1999; Peckmann et al., 2002; Flügel, 2004). Microbially oxidized crude oil coating aragonite crystals and pore spaces suggests a close association between carbonate authigenesis and hydrocarbon seepage from the subsurface. Abundant authigenic pyrite in the carbonate matrix (Fig. 2F) is suggestive of sulfate reduction coupled to anaerobic hydrocarbon oxidation, which increases carbonate alkalinity while generating dissolved sulfide, the latter forming pyrite through reaction with reactive Fe phases (Peckmann et al., 2001; Peckmann and Thiel, 2004).

Evidence for multiple seepage episodes include distinctly younger carbonate cementation around pore spaces and fractures through clasts (Fig. 2 and 5). We propose that fracture-filling carbonate cement is evidence for multiple seepage events as well as potential endolithic activity. It has been shown that seep carbonates can serve as a unique m crocial habitat for endolithic activity capable of continued anerobic hydrocarbon oxidation and a uthigenic carbonate aggregation even after their formation (Marlow et al., 2014; Marlow et al., 2015; Yanagawa et al., 2019). Marlow et al. (2015) used the term 'autoendolithic' for sthe endolithic activity because that microbial metabolism induces mineral precipitation and leads to the formation of rock that remains inhabited by the same kind of microbes. We also observed pervasive fractures, potentially caused by *in situ* brecciation of semi-indurate.' sectiments and clasts, resulting from pressure induced by rising hydrocarbons (e.g., Hovland et 1, 1987; Beauchamp and Savard, 1992; Campbell et al., 2008). These features were subsequently filled by aragonitic seams and are commonly observed together with hydrocarbon incrisions. These may indicate episodic seepage and subsequent carbonate precipitation (ven's via anaerobic hydrocarbon oxidation (Fig. 5A-E). Hence we propose that these fract re-filling signatures could be indicative of autoendolithic structures reflecting self-entombment of hydrocarbon-oxidizing microbes (Marlow et al., 2015, Fig. 5F). In addition, Campeche seep carbonates were shown to entrap methane and other hydrocarbons  $(C_2 \text{ up to } C_6)$  in the crystal spaces of carbonate minerals (Blumenberg et al., 2018). The presence of trapped gases may trigger anaerobic hydrocarbon oxidation coupled with sulfate reduction (considering the aragonite crusts are formed near the sediment-water interface with high porosity, allowing sulfate diffusion), resulting in carbonate authigenesis and potentially endolithic activity. Future work using fluorescence microscopy along with high-resolution atomic force microscopy is expected to shed more light on these unique biosignatures.

![](_page_25_Figure_1.jpeg)

Figure 5: Evidence for common endolithic activity in fractures of micritic cements at Chapopote Seeps (A-E) and interpretative drawing of autoendolithic microbe-rock relationship (F). Hydrocarbon oxidation by the microbial communities inhabiting the fractures of the seep carbonates induce additional carbonate precipitation, leading to self-entombment of

hydrocarbon-oxidizing microbes (autoendolithic activity). White arrows mark sites of potential mineralization induced by autoendolithic activity and the yellow arrow in Figure E marks a fracture Figure F is a schematic representation for relative locations and microbe-rock interactions of autoendolithic organisms adapted from Marlow et al., 2015. Circles represents endoliths and gray shading represents successive zones of autoendolithic carbonate precipitation. Refer to section 5.1 for discussion.

#### 5.2 Temporal Variability of GoM Seeps

Since authigenic carbonates may serve as a reliable geological (rchive for seepage histories including C and S cycling, constraining their precipitation ages should provide us a glimpse of the temporal patterns, causes, controls, and consequences of seep systems. U-Th dating of seep carbonates is a proven tool for constraining past seep activity at diverse seep settings globally (Aharon et al., 1997; Teichert et al., 2003; Watanabe et al., 2008; Bayon et al., 2009; Mazumdar et al., 2009; Liebetrau et al., 2010; Wirsig et al., 2017; Berndt et al., 2014; Crémière et al., 2016; Prouty et al., 2016; Mazzini et al., 2017; Saue, et al., 2017; Chen et al., 2019; Himmler et al., 2019; Judd et al., 2019; Wei et al., 2020).

To the best of our knowledge, our U-Th dates are the first for authigenic carbonates from the southern GoM. Only a few studies (A' a on et al., 1997; Feng et al., 2010) have reported U-Th dates for seep carbonates in the CoM. Also, Bian et al. (2013) constrained long-term seepage variability using <sup>14</sup>C dates on bivalve shell materials cemented in the seep carbonates. These studies were limited to the northern GoM, and so the timing and duration of seepage in the southern GoM is unknown. Sediment supply, sea-level changes, and salt deformation are suggested as the major factors that controlled recent hydrocarbon seeps in the GoM (Roberts and Carney, 1997).

Previous U-Th age-dates from northern GoM seeps along the lower continental margin and from upper bathyal depths to the abyssal plains showed strong evidence of discontinuous fluid flux in these basins during the late Quaternary (Aharon et al., 1997; Feng et al., 2010). Interestingly, the ages of many authigenic carbonate samples there clustered around 12 ka, suggestive of a role played by sediment loading and salt tectonics on glacial-interglacial time scales. Lower sea level would induce higher sediment loading on continental slopes. Rising sea level associated with deglaciation, in contrast, should reduce sediment loading in deeper waters by shifting deposition

landward to cause salt tectonic adjustment and the development of fault conduits for hydrocarbon seepage (Aharon et al., 1997; Feng et al., 2010; Roberts and Feng, 2013). Our age dates fall within that last deglaciation time frame, with the oldest being 13.5 ka and the youngest being 4.6 ka (BP).

While the northern GoM is dominated by sediment loading from the Mississippi river, southern GoM receives strong discharge from the Grijalva-Usumacinta River, the second largest river draining into the GoM following the Mississippi (Salas-de-León et al., 2008). The salt activity of the southern GoM is likely analogous to that of the Texas-Louis iana slope owing to its shared geological history (Garrison and Martin, 1973). Shallow sedime and salt structures associated with regional salt tectonics are shown to have dominant contained and salt structures associated with regional salt tectonics are shown to have dominant contained and seep distribution along the Campeche Knolls (Ding et al., 2008; Ding et al., 2010) These factors suggest possible fault activation of hydrocarbon seepage linked to changes in sectiment loading in the Campeche Bay during deglaciation, similar to the controls in the northern GoM (Feng et al., 2010; Roberts and Feng, 2013). A case for sediment-loading-driven and the cause for seep initiation in southern GoM, however, demands additional study. Nevertheless, our U-Th age data from the southern GoM combined with existing northern GoM data strongly suggest that seeps along GoM slopes were particularly active during the last deglaciation.

#### 5.3 Carbon sources for DIC and CaCO<sub>3</sub>

Based on the average  $\delta^{13}C_{CaCO_3}$  value of -25%, a predominantly crude oil source for carbon was suggested for Chapopote  $\infty$  carbonates (up to 90%) by Naehr et al. (2009). This inference was derived using carbon is tope mass balance based on the isotopic composition of end members including methane ( $\delta^{13}C_{CH4} = -55\%$  [MacDonald et al., 2004]), crude oil (average  $\delta^{13}C$  value for GoM basin = -27% [Anderson et al., (1983]), sedimentary organic matter ( $\delta^{13}C = -20\%$  [Goñi et al., 1998]), and DIC from seawater ( $\delta^{13}C = +1\%$  [Aharon et al., 1991]). The headspace methane isotope measurement ( $\delta^{13}C_{CH4} = -55\%$ ) from Chapopote Knoll is indicative of a thermogenic ( $\delta^{13}C_{CH4}$  range -30% to -50%) and biogenic [ $\delta^{13}C_{CH4}$  range -60% to -90%] admixture (MacDonald et al., 2004; Naehr et al., 2009). Measured  $\delta^{18}O_{CaCO3}$  values ranged from +3.3 to +5.9 ‰, averaging +4.5‰. This was higher than the calculated  $\delta^{18}O_{CaCO3}$  value (+3.02‰) based on the measured bottom water temperature (4.02°C) and pore fluid  $\delta^{18}O$  values (+0.09‰) of background samples, according to Kim et al. (2007) (Supp. Table 2). Porewater <sup>18</sup>O-

enrichment sourced from shallow gas hydrate dissociation could explain this difference (Davidson et al., 1983; Naehr et al., 2009). These observations agree with Formolo et al. (2004), who suggested that  $\delta^{13}$ C of DIC (and CaCO<sub>3</sub>) at oil seep settings such as our study site may be swamped by AONM signals despite the presence of shallow pockets of gas hydrates, while the  $\delta^{18}$ O signals of adjacent porewater (CaCO<sub>3</sub>) are isotopically sensitive to <sup>18</sup>O-enrichment from shallow hydrate dissociation.

#### 5.3.1 Crude oil oxidation (AONM)

Multiple lines of evidence point to crude oil oxidation coupled with sulfate reduction as the dominant DIC source for carbonate precipitation of Chapopote Kinol.

- (1) the  $\delta^{13}C_{CaCO3}$  values averaging -25‰ are closer to a rule of signature;
- (2) the presence of abundant biodegraded crude oil w thin the carbonate samples (Fig. 2);
- (3) widespread crude oil degradation signals from organic geochemistry studies on asphalts, oily sediments, and seeping crude oil (Schrberz et al., 2011b);
- (4) an abundant unresolved complex or gar ic mixture (UCM) in the carbonate pore-filling crude oil (Naehr et al., 2009), with an ex vated baseline for the C15 to C30 range indicative of degradation involving loss of lobile n-alkanes and isoprenoids from crude oil (Mansour and Sassen, 2011; Schubotz et al., 2011b);
- (5) recently reported evidence from Chapopote seep sediments for a symbiotic microbial consortium composed of thort-chain alkanes degraders with contributions from sulfate-reducing bacteria (Last Pérez et al., 2019);
- (6) distinctive phase-specific enrichment of rare earth elements and U in Chapopote carbonates compared to methane-derived carbonates (Smrzka et al., 2016); and
- geochemical batch modeling suggestive of sulfate-driven crude oil oxidation contributing to carbonate precipitation at Campeche Knolls (Smrzka et al., 2019).

These multiple lines of evidence developed over the past two decades at Campeche Knolls, along with our petrographic observations and  $\delta^{13}C_{CaCO3}$  data, allow us to use Chapopote seep carbonates as a template for recognizing authigenic carbonates sourced via AONM in other regions.

#### 5.4 C-S Isotope Systematics of the AOM and AONM settings

To test the veracity of AONM signals at Campeche Knolls, we also analyzed sediments from GC 415 and GC 185 and gathered published data from multiple seep settings in the GoM (Formolo and Lyons, 2013; Feng et al., 2016; Sun et al., 2020). This extensive database allowed us to compare and contrast the C-S isotope systematics of sulfide and DIC produced via sulfate driven AOM and AONM as recorded in GoM seep CaCO<sub>3</sub> (Table 2 and Supp. Table 1). Along with the help of cluster analysis and site characteristics, we were able to define two dominant end members based on  $\delta^{13}C_{CaCO3}$  versus  $\delta^{34}S_{CRS}$  plots (Fig. 6):

- (1) AOM-dominant seepage sites with strong <sup>13</sup>C-depletion in  $CaCO_3$  ( $\delta^{13}C_{CaCO_3} < -40\%$ ) indicating a methane source for carbon; and <sup>34</sup>S<sub>CRS</sub> values with <sup>34</sup>S-enrichment ( $\delta^{34}S_{CRS}$ >0%) suggestive of dissolved sulfide production and sulfide mineral precipitation associated with AOM (e.g., Jørgensen et al., 2004, Bolowski et al., 2013);
- (2) Crude oil oxidation sites dominated by A Nivi as expressed in moderate <sup>13</sup>C-depletion in  $\delta^{13}C_{CaCO3}$  ( $\delta^{13}C_{CaCO3} = -25.8\% \pm 1.5$ ) and  $\delta^{34}S_{CRS}$  values with relatively strong <sup>34</sup>S-depletion ( $\delta^{34}S_{CRS} < -15\%$ ), suggraving DIC sourced from AONM. These samples include virtually all (98%) of the Chap( p(4) seep carbonates in this study and those from northern GoM sites GC 185 and GC 2.32, crude oil seep sites (Feng et al., 2009; Sun et al., 2020).

The data points lying between the two end members indicate mixing from multiple carbon sources and biogeochemical processes (DIC and sulfide pools from AOM, AONM, sedimentary organic matter degradation, DIC sourced from methanogenesis and the water column, as well as likely  $\delta^{34}$ S signals from secondary sulfur cycling (e.g., Raiswell, 1987; Naehr et al., 2000; Formolo and Lyons, 2013; Crémière et al., 2020). Overall, as explained below, we seem able to identify sources of carbon for the DIC pool and suggest the particular sulfate reduction pathway that generates DIC and dissolved sulfide, namely AOM, AONM, or OSR. This observation also agrees with previously observed disparities between measured rates of sulfate reduction and methane oxidation (Joye et al., 2004; Bowles et al., 2011). Those data suggest that sulfate reduction is not driven primarily by AOM—thus highlighting the complex C-S relationship, with sulfate reduction coupled to both AOM and AONM at GoM seeps.

![](_page_30_Figure_1.jpeg)

Figure. 6: Cross plot of  $\delta'^3 C_{CaCO3}$  vs.  $\delta^{34}S_{CRS}$  values from authigenic calcium carbonate (CaCO3) and selected shell samples found within surface sediments at various seep sites in the Gulf of Mexico. We sampled sediment at Chapopote Knoll (blue triangles) and measured isotopic values of sediments from GC 185 and GC 415. Other data are taken from the literature. Symbols refer to different types of seepage: sites with dominant oil seepage are represented by crosses (Chapopote Knoll, GC 232, and GC 185), sites with dominant methane seepage are shown with triangles (AT 340, GB 260, GC 238, and GC 852), and remaining sites with a mixture of oil and methane seepage are shown with circles. Vertical dashed line shows the  $\delta^{13}C_{CH4}$  value for Chapopote headspace methane samples. The decision boundaries (shaded areas) of end-members were loosely defined based on the cluster density and site characteristics. Refer to Table 1 for site descriptions, Supp. Table 1 for data, and section 5.4 for discussion.

The  $\delta^{13}C_{DIC}$  signatures for AOM and AONM varies primarily because the  $\delta^{13}C$  of the substrates – CH<sub>4</sub> vs oil – are very different (section 5.3) and this affect the carbon isotopic composition of the DIC pool markedly (Formolo et al., 2004; Naehr et al., 2009; Mansour and Sassen, 2011; Roberts and Feng, 2013). However, the possibility of multiple carbon sources (section 5.3; e.g., methane, crude oil, seawater DIC, sedimentary organic matter) and associated processes (AOM, AONM, OSR, methanogenesis) leading to varying degrees of carbon-isotope fractionations of DIC at seeps (Sassen et al., 2004; Meister et al., 2019) can limit the application of  $\delta^{13}C_{CaCO3}$  as a single proxy to evaluate the DIC sourcing at seeps. Hence, we highlight that the combination of paired  $\delta^{13}C_{CaCO3}$  and  $\delta^{34}S_{CRS}$  offers additional constraints to disurguish the DIC and sulfide sourcing through sulfate reduction coupled to the oxidation of different carbon substrates (mainly methane vs crude oil).

The  $\Delta^{34}S_{sulfate-sulfide}$  ( $\delta^{34}S_{seawater sulfate} - \delta^{34}S_{porewater sulfid}$ ) depends on the rate at which seawater sulfate exchanges with subsurface diagenetic horizon: and the isotopic fractionation by sulfate-reducing microbes (Kaplan and Rittenberg, 1964; Yeahicht and Canfield, 1997; Wehrmann et al., 2011; Kump, 2012; Fike et al., 2015). In a comparative study of sulfate reduction rates from a gas seep, oil seep, and non-seep background sites in GoM, Aharon and Fu (2000) reported highest sulfate reduction rates at methode scep (0.27 to 2.51 µm SO<sub>4</sub><sup>2</sup> cm<sup>-3</sup> day<sup>-1</sup>) followed by oil seep (0.01 to 0.22 µm SO<sub>4</sub><sup>2-</sup> cm<sup>-3</sup> day<sup>-1</sup>) and non-seep background sediments (0.0043 µm SO<sub>4</sub><sup>2-</sup> cm<sup>-3</sup> day<sup>-1</sup>). These results suggests that the carbon substrate undergoing oxidation coupled to sulfate reduction has an important control on the sulfate reduction rates and associated isotope fractionations that can impact the  $\delta^{34}S_{sulfide}$  value (Aharon and Fu, 2000; Sun et al., 2020).

In a typical marine setting lacking significant methane transport, OSR would predominate. In such settings, slow sulfate reduction rates lead to high fractionations, which, under non-limiting sulfate conditions (open-system), would result in low  $\delta^{34}S_{sulfide}$  values (Jorgensen, 1979; Canfield, 2001; Lin et al., 2017b). In contrast, sites with higher subsurface methane fluxes often show higher rates of sulfate reduction due to AOM and exhaustion of sulfate in the local sulfate pool (closed-system) within the sediment column, resulting in high rates of sulfide mineral precipitation around the SMTZ with enriched <sup>34</sup>S signals (Peckmann et al., 2001; Jørgensen et al., 2004; Wang et al., 2008; Lim et al., 2011; Peketi et al., 2012; Borowski et al., 2013; Deusner et al., 2014; Peketi et al., 2015; Li et al., 2016; Lin et al., 2016a; Lin et al., 2016b; Li et al., 2017;

Lin et al., 2017a; Fan et al., 2018; Wu et al., 2019; Argentino et al., 2020). Provided sufficient Fe availability to form iron sulfide minerals,  $\delta^{34}S_{CRS}$  would be a good recorder for the porewater sulfide since isotopic fractionation associated with the sulfide mineral formation from dissolved sulfide is negligible (Price and Shieh, 1979; Butler et al., 2004). In addition to the closed-system behavior, the enriched  $\delta^{34}S_{CRS}$  signals we observe at methane seep sites are also suggestive of small instantaneous fractionations associated with rapid rates of sulfate reduction resulting in isotopically heavy sulfide that gets captured by available Fe early in the process close to the sediment-water interface (E.g., Kaplan and Rittenberg, 1964; Chanton, 1985; Lyons, 1992; Lyons, 1997; Gilhooly et al., 2016).

Oil seepage settings, where AONM occurs, could result in achaic reduction rates higher than those associated with OSR but lower than AOM (Aharon and Fu, 2000; Aharon and Fu, 2003). Thus, AONM yields fractionations larger than those associated with AOM as well as less efficient removal of sulfate. The net result would be relatively depleted <sup>34</sup>S signals in sulfide minerals. Some of the variability within the obset ed low  $\delta^{34}S_{CRS}$  values (-10 to -39‰) can be attributed to the diversity and concentration of hydrocarbon compounds involved and related differences in sulfate reduction rates microbial species and metabolic diversity, as well as varying impacts of secondary sulfide vicing such as sulfide oxidation and disproportionation (Kemp and Thode, 1968; Canfield and Thamdrup, 1994; Detmers et al., 2001; Habicht and Canfield, 2001; Lyons and Gill 2010; Sim et al., 2011; Gallagher et al., 2012; Leavitt et al., 2013; Deusner et al., 2014; Con ngelo-Lillis et al., 2019; Jørgensen et al., 2019; Pellerin et al., 2019; Bazzaro et al., 2020; Crémière et al., 2020). Overall, however, highly negative <sup>34</sup>S<sub>CRS</sub> cluster ( $\delta^{34}S_{CRS} < -15\%$ ), linked to AONM seem distinct from the dominantly heavier isotopic cluster associate with AOM. It is also noteworthy that  $\delta^{13}C_{CaCO3}$  and  $\delta^{34}S_{CRS}$  values for Chapopote Knoll, GC 232, and GC 185-the three sites with oil seepage and AONM-show similar  $\delta^{13}C$  and  $\delta^{34}S$  signals (Fig. 6). Hence, we interpret the high  $\delta^{34}S_{sulfide}$  and very low  $\delta^{13}C_{DIC}$  values in our compilation to be sourced predominantly by AOM and the relatively low  $\delta^{34}S_{sulfide}$  and moderately low  $\delta^{13}C_{DIC}$  values to be sourced primarily from crude oil oxidation and OSR (Fig. 6).

The aragonitic composition of Chapopote carbonates is strongly supportive of a shallow diagenetic origin in the presence of sulfate, low phosphate concentration, and high Mg/Ca ratios

(Burton and Walter, 1990; Burton, 1993). Aloisi et al. (2002) suggested that porewater sulfate concentration can play an important role in determining seep carbonate mineralogy since sulfate inhibits Mg-calcite precipitation and favors aragonite formation. Alternatively, Mg-calcite precipitation maybe catalyzed by the presence of sulfide ions, which accelerate Mg dehydration and incorporation into the mineral lattice (Zhang et al., 2012; Zhang et al., 2013; Lu et al., 2018; Smrzka et al., 2021). Mansour (2014), in a compilation based on seep carbonates from multiple GoM sites, suggested that aragonite is the dominant mineralogy at sites of crude oil oxidation in comparison to methane seep settings due to relatively lower sulfate reduction rates during crude oil oxidation. This relationship was also reported for crude oil oxid, tion at site GC 185 by Feng et al., (2009) and GC 232 by Sun et al. (2020). Aragonite min. ralc gy of authigenic carbonates reported from Chapopote corroborate to these findings (Fig. 7. AOM-dominated samples, in contrast, showed a diverse mix of carbonate composition spanning aragonite, high-Mg calcite, and low-Mg calcite (Fig. 7). Sufficient Fe availability is iron sulfide mineral precipitation is necessary when considering  $\delta^{34}S_{CRS}$  as a faith full recorder of porewater sulfate reduction processes, which is not always the case in mony GoM seep settings, especially below the surface-most sediment layers (Formolo and Lyons, 2013). Hence aragonite samples, preferably formed closer to the sediment-water in rface (e.g., Feng et al., 2016), where high interstitial sulfide concentrations and sufficien. He availability are more likely, serves ideal targets to evaluate the AOM versus AONM listinction based on  $\delta^{13}C_{CaCO3}$  and  $\delta^{34}S_{CRS}$ .

While the AOM and AONM  $s_{12}$  hals were clearly distinguishable, the boundary between crude oil oxidation and OSR is not clearly defined based on existing data. The  $\delta^{34}S_{CRS}$  data fail to show a clear distinction between OSR and AONM (Fig. 6). The  $\delta^{13}C_{CaCO3}$  results also do not clearly demarcate the two processes likely due to combination of similar values between  $\delta^{13}C$  values of organic matter and oil, slight fractionation between organic matter and DIC, and mixing of DIC sources. This challenge is exacerbated by the fact that our samples and most of the literature data compiled considered here are from seep sites. For example, data from site GC 234 taken from Formolo and Lyons (2013), despite belonging to a seep site, plot with the OSR cluster, with only a hint of seep-induced C-S signals (average  $\delta^{13}C_{CaCO3} = -11.9\%$  and  $\delta^{34}S_{CRS} = -15.3\%$ ). This contradiction comes as no surprise because the data likely reflect a combination of OSR, AONM, and Fe-limitation (Formolo and Lyons, 2013)—pointing to the need for additional paired

 $\delta^{13}C_{DIC}$ - $\delta^{34}S_{sulfide}$  data from both OSR and seep-dominant sites from the GoM to better constrain the end members as well as the frequent cases of mixed signals.

There are a few additional arguments that lend support to the possibility of distinguishing between OSR and AONM in our cross plot. The carbonate content for the samples that plot in OSR cluster is very low (e.g., average carbonate content ~24% Supp Table. 1) in comparison to the crude oil seep samples (average carbonate content = ~80%). This observation is in agreement with reports suggesting that authigenic carbonate precipitation decreases at low sulfate reduction rates, due to the dominant effect of H<sup>+</sup> over bicarbonate (Meister 2013; Zhang, 2020). Further, samples in the OSR cluster show minimal <sup>18</sup>O-enrichment (Fi<sub>5</sub>. 3C, Supp. Table 1), unlike a seep-dominant AONM setting where the  $\delta^{18}O_{CaCO3}$  sign 2. ... e often influenced by <sup>18</sup>O enrichment sourcing from localized shallow gas hydrate dissociation (Formolo et al., 2004), deep hydrocarbon rich brines (Mansour and Sassen, 2011), and less likely) clay mineral dehydration (Roberts and Feng, 2013). These observations, along v the cariched  $\delta^{13}C_{CaCO3}$  values than AONM samples, suggest that the data from GC 234 p'o. or in the OSR cluster are likely influenced more by OSR than AONM.

Recent studies have shown that multip.'e sulfur isotope signatures of sulfide minerals ( $\Delta^{33}$ S and  $\delta^{34}$ S) could be an effective tool to digiting 'ish between OSR and AOM (Lin et al., 2017b; Gong et al., 2018; Lin et al., 2018; Crémie.'e et al., 2020; Liu et al., 2020). Such efforts, along with incorporation of new data from additional seep and OSR dominant sites, will improve the definition of end-member being aries. The relative roles of aerobic and anaerobic pathways of hydrocarbon oxidatio. in sur ace/near-surface sediments also remain fodder for future research as related to carbonate precipitation and dissolution (Aller, 2014). Redox-sensitive proxies such rare earth element patterns and biomarker records of aerobic hydrocarbon oxidizers preserved in the carbonate matrix will aid in these distinctions (Feng et al., 2009; Birgel et al., 2011; Smrzka et al., 2020)

![](_page_35_Figure_1.jpeg)

Figure 7: Cross plot of  $\delta^{13}C_{CaCO}$  vs.  $\delta^{14}S_{CRS}$  values of various phases of authigenic calcium carbonate (CaCO<sub>3</sub>) and selected shell samples as aragonite (triangles), and low-Mg and high-Mg calcite (squares and crosser, respectively), and of authigenic pyrite found within surface sediments of oil seep sites, Chapopote Knoll, Gulf of Mexico; and Northern GoM seep sites plotted in Fig. 6. GC 25.<sup>4</sup> and not have the mineralogy information hence is not included in the plot. Vertical dashed line shows the  $\delta^{13}C_{CH4}$  value for Chapopote headspace methane samples. The shaded areas on the top left and bottom right represent the same AOM and AONM+OSR endmember pools in Fig. 6.

#### 5.5 Implications for the sedimentary record

Carbonate systems are Earth's largest carbon reservoir, accounting for >60 million Gt C (Falkowski et al., 2000). Carbonate burial in marine sediments is an important part of carbon cycling through most of the geological history, accounting for ~80% of the total carbon removal

from Earth's surface today (Derry, 2014; Sun and Turchyn, 2014; Berg, 2018).  $\delta^{13}C_{CaCO3}$  signals from the geologic record are an important sedimentary proxy in efforts to understand the evolution of the carbon cycle and the chemical composition of the ocean-atmosphere system over geological history (Hayes et al., 1999; Berner, 2003; Katz et al., 2005). For example, a period of higher organic carbon burial would remove more <sup>12</sup>C carbon, resulting in <sup>13</sup>Cenrichment in DIC (and CaCO<sub>3</sub>) (Berner, 2003; Canfield and Kump, 2013; Lyons et al., 2014). This approach has been used to reconstruct organic carbon burial and the oxygen content of the atmosphere (Shackleton, 1985; Kump and Arthur, 1999; Berner, 2004). Authigenic carbonate precipitation was invoked as a third major carbon sink, with major impact on  $\delta^{13}C_{CaCO3}$ fluctuations in the geological record (Schrag et al., 2013). Fu ther, authigenic carbonate precipitation may result in significant carbon sequestration up esent, comparable to ~15% of carbonate accumulation on continental shelves and in the authysal ocean, respectively (Akam et al., 2020). Hence, better characterization of the sedime, ary proxies for authigenic carbonate formation is an important goal (e.g., Bjerrum and Can'ield, 2011; Bristow and Grotzinger, 2013; Zhao et al., 2016; Cui et al., 2017; Davis Ba' are tal., 2019; Jiang et al., 2019).

Our results emphasize an important additional process for C-S coupling through organic carbon decomposition and resulting authiger carbonate formation—crude oil oxidation (Formolo et al., 2004; Peckmann et al., 2007; Neer et al., 2009; Mansour and Sassen, 2011; Formolo and Lyons, 2013; Kiel and Peckmann, 2019; Smrzka et al., 2019)-to add to the well-known processes of AOM and OSR (Pradbury and Turchyn, 2019; Akam et al., 2020). Further, we explore the use of cross ploting  $\delta^{13}C_{CaCO3}$  versus  $\delta^{34}S_{CRS}$  to distinguish crude oil seepage and methane seepage proces, as in present sediments and in the geologic record, which can be confirmed using additional evidence, such as petrography and mineralogy (Mansour, 2014), biomarker analysis (particularly unresolved complex mixture suggestive of petroleum degradation; Sassen et al., 2001; Naehr et al., 2009; Mansour and Sassen, 2011), trace metal and rare earth element concentration (Smrzka et al., 2016; Smrzka et al., 2019), and total organic carbon and total organic sulfur contents (Sun et al., 2020). Our  $\delta^{13}C_{CaCO3}$  versus  $\delta^{34}S_{CRS}$  from GoM could serve as a template for assessment of carbon sources and sulfate-reduction and other early diagenetic processes from additional sites and regions. Further, authigenic carbonate records have been used to estimate past methane fluxes (e.g., the Neoproterozoic caprocks of Snowball Earth; Kennedy et al., 2001; Jiang et al., 2003). An ability to better distinguish

methane-driven carbonate authigenesis from that derived from oxidation of other hydrocarbons could allow us to better constrain the temporal variations in methane fluxes over geologic history.

#### 6. Conclusion

We examined the C-S coupling and temporal patterns for Chapopote asphalt seeps in the southern Gulf of Mexico based on CaCO<sub>3</sub> geochemistry and compared our result with data available from multiple northern GoM seep sites. Carbonate petrography of Chapopote samples showed peloidal textures and autoendolithic features, indicative of dynamic, microbially-driven biogeochemical process resulting in carbonate authigenesis. V/e present the first U-Th-based ages for seep carbonates from the southern GoM. U-Th ages for Chapopote seep carbonates ranged from 13.5 ka to 4.6 ka (BP), suggesting that Chapopote asphalt seepage has been ongoing for thousands of years. These results encourage further have stigation that GoM slopes might have experienced seep activation during the last deglacious on owing to changes in sedimentary loading and associated salt-tectonic adjustment, which activates fault conduits for deep-seated hydrocarbon seepage.

Isotopic measurements of the authigenic n-inerals of calcium carbonate (CaCO<sub>3</sub>) and bulk sulfide minerals (native sulfur, iron monosulfices, pyrite) reveal the geochemical characteristics of interstitial DIC and dissolved sulfide. A cross plot of  $\delta^{13}C_{CaCO3}$  and  $\delta^{34}S_{CRS}$  distinguishes between carbon substrates (mechane vs. crude oil) and between the sulfate-reduction processes of anaerobic oxidation of reachane (AOM) and anaerobic oxidation of non-methane (AONM).. Authigenic carbonate sumples from asphalt seeps at Chapopote Knolls showed distinct signatures, with  $\delta^{13}C_{CaCO3}$  and  $\delta^{34}S_{CRS}$  values characterized by moderate <sup>13</sup>C-depletions ( $\delta^{13}C_{CaCO3} \sim -25\%$ ) and relatively strong <sup>34</sup>S-depletions ( $\delta^{34}S_{CRS} < -15\%$ ), indicative of DIC sourcing via AONM. AOM-dominant seepage sites, in contrast, show strong <sup>13</sup>C-depletion in CaCO<sub>3</sub> ( $\delta^{13}C_{CaCO3} < -40\%$ ) and noticeable <sup>34</sup>S.enrichment in sulfide minerals ( $\delta^{34}S_{CRS} > 0\%$ ) values, indicative of predominantly AOM sourcing for DIC. The spread of data in between the two end-members indicate mixing from multiple sources (DIC and sulfide pools from AOM, crude oil oxidation, organic matter degradation, secondary sulfide cycling, as well as DIC sourced from methanogenesis and the water column). Future work on triple S isotopes, organic S, and CAS may shed light on the full extent of sulfur cycling in these settings as well as to

improve the current field boundaries.  $\delta^{13}C_{CaCO3}$  versus  $\delta^{34}S_{CRS}$  compilation from the GoM could serve as a template for data from additional sites and regions and highlights the importance for producing more paired  $\delta^{13}C_{CaCO3}$  and  $\delta^{34}S_{CRS}$  dataset from diverse seep and OSR dominated settings to build on this template. Overall, the combination of age-dating and detailed geochemical analyses highlights the potential for better evaluation of carbonate authigenesis in seep settings, which is a significant component of marine carbon burial and an important geological archive for seepage events.

#### Acknowledgments

SA would like to acknowledge the Graduate Research Capit. from the American Association of Petroleum Geologists (AAPG), Geological Society of America (GSA), and Gulf Coast Association of Geological Societies (GCAGS). Soft arships to SA from TAMUCC Endowment Fund, Gulf Coast Chapter of Air & Waste Malagement Association (A&WMA), and Corpus Christi Geological Society (CCGS) are acknowledged for the support during this research. The Petroleum Research Fund of the American Chemical Society provided support to TWL. Christine Chen, Ben Hardt, and Gabriela perrato Marks at McGee Lab (MIT) are thanked for the assistance during the U-Th analysis. The Lyons' group at UCR is acknowledged for the support to SA during the lab visit for inotogic analysis. Pankaj Khanna, Tarini Bhatnagar, Wasif Riza, and Mustafa Kalam are thanked to the cheir assistance with Figure 1 and cluster analysis. Samples for this study were obtain d charling R/V SONNE and R/V Meteor cruises SO174/2 and M67/2. We would like to extend our marks to the entire team involved in those expeditions. We would like to thank Walter S. Borowski and an anonymous reviewer for the constructive inputs that greatly helped in improving this manuscript.

#### **Declaration of Interest Statement**

We declare that this manuscript entitled "Carbon-sulfur signals of methane versus crude oil diagenetic decomposition and U-Th age relationships for authigenic carbonates from asphalt seeps, southern Gulf of Mexico" with authors Sajjad A Akam, Timothy W. Lyons, Richard B.

Coffin, David McGee, Thomas H. Naehr, Steven M. Bates, Clay Clarkson, and Brandi Kiel Reese is original, has not been published before and is not currently being considered for publication elsewhere. We know of no conflicts of interest associated with this publication, and there has been no significant financial support for this work that could have influenced its outcome. As the corresponding author, I confirm this on behalf of all the named authors.

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## Highlights:

- We present new insights into the geological signatures of anaerobic crude oil oxidation coupled with sulfate reduction using authigenic carbonate geochemistry from southern Gulf of Mexico (Chapopote asphalt volcano, Bay of Can pec. e).
- We report the first U-Th-based ages for seep carbon and nom southern Gulf of Mexico. U-Th age-dating of Chapopote seep carbonate samples vielded ages of 13.5 ka – 4.6 ka before present (BP), suggesting that Chapopote as halt seepage has been ongoing for thousands of years.
- We report distinguishable variation in the paned  $\delta^{13}C_{DIC}$  and  $\delta^{34}S_{sulfide}$  signatures produced via sulfate-driven anaerobic ox ar ion of methane (AOM) and non-methane hydrocarbons AONM), which can be normalised through paired  $\delta^{13}C_{CaCO3}$  and  $\delta^{34}S_{sulfide-mineral}$  signals in seep carbo at s.
- We present a  $\delta^{13}C_{CaCO3}$  vs.  $\delta^{34}S_{CRS}$  ross plot to distinguish and classify the near-surface authigenic carbonate systems in the Calf of Mexico formed due to anaerobic methane oxidation and crude oil oxidation.