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Carbon isotopic evidence for microbial control of carbon supply to Orca Basin at the seawater-brine interface

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Abstract. Orca Basin, an intraslope basin on the Texas-Louisiana continental slope, hosts a hypersaline, anoxic brine in its lowermost 200 m in which limited microbial activity has been reported. This brine contains a large reservoir of reduced and aged carbon, and appears to be stable at decadal time scales: concentrations and isotopic composition of dissolved inorganic (DIC) and organic carbon (DOC) are similar to measurements made in the 1970s. Both DIC and DOC are more "aged" within the brine pool than in overlying water, and the isotopic contrast between brine carbon and seawater carbon is much greater for DIC than DOC. While the stable carbon isotopic composition of brine DIC points towards a combination of methane and organic carbon remineralization as its source, radiocarbon and box model results point to the brine interface as the major source region for DIC, allowing for only limited oxidation of methane diffusing upwards from sediments. This conclusion is consistent with previous studies that identify the seawater-brine interface as the focus of microbial activity associated with Orca Basin brine. Isotopic similarities between DIC and DOC suggest a different relationship between these two carbon reservoirs than is typically observed in deep ocean basins. Radiocarbon values implicate the seawater-brine interface region as the likely source region for DOC to the brine as well as DIC.

1 Introduction

Active and diverse microbial communities have been reported in deep-sea brines of the Gulf of Mexico, Red Sea and Mediterranean Sea (reviewed in Antunes et al., 2011; Joye et al., 2010). These anoxic brines test the limits of microbial adaptation across a range of temperatures, salinities and pH conditions. Orca Basin, a deep basin located on the Texas-Louisiana slope, is the largest brine pool yet identified in the Gulf of Mexico (Pilcher and Blumstein, 2007; Shokes et al., 1977; Trabant and Presley, 1978). Although moderate conditions are found in Orca Basin brine compared to other deepsea brines (Antunes et al., 2011; Joye et al., 2005), studies of microbial abundance and activity in Orca Basin have concluded that very little physiological activity can be measured below the seawater-brine interface (Dickins and Van Vleet, 1992; LaRock et al., 1979; Tuovila et al., 1987; Wiesenburg et al., 1985). This lack of microbial activity stands in contrast to other Gulf of Mexico brines in the Gulf of Mexico (Joye et al., 2009, 2010).

The continental slope of the northern Gulf of Mexico is characterized by numerous basins and pockmarks extending from the shelf edge to the basin floor (Bouma and Bryant, 1994, and references therein). This complex bathymetry is produced by the interaction of sediments and underlying salt, which drives salt deformation (Humphris, 1978), a process also associated with brine formation and brine seepage along seafloor fault networks (Reilly et al., 1996; Roberts and Carney, 1997). Although brine seepage is considered widespread in the northwestern Gulf of Mexico, only a few brine pools have been detected and studied (Joye et al., 2005; Macdonald et al., 1990; Shokes et al., 1977). Orca Basin, an intraslope depression at a water depth of approximately 2000 m on the northwestern continental slope of the Gulf of Mexico, was the first among these to be identified (Shokes et al., 1977; Trabant and Presley, 1978). Below 2240 m depth, the basin is occupied by an anoxic, hypersaline brine. Both the geochemistry of the brine and composition of underlying sediments indicate the ~ 200 m thick brine originates from dissolution of a nearby salt exposure and lateral advection of brine to the basin (Addy and Behrens, 1980; Pilcher and Blumstein, 2007; Sheu et al., 1988; Shokes et al., 1977; Trabant and Presley, 1978). This formation mechanism is unlike many other Gulf of Mexico brines that have in situ sources of dissolved ions and reduced gases (Joye et al., 2005, 2009, 2010).

Within the Orca Basin brine, oxygen and nitrate are not detectable while phosphate concentrations are elevated compared to overlying seawater resulting from organic-matter decomposition (Leventer et al., 1983; Shokes et al., 1977; Van Cappellen et al., 1998). Sulfate concentrations, however, are not depleted within the brine suggesting a limited role for microbial sulfate reduction in the anoxic brine (Hurtgen et al., 1999; Leventer et al., 1983; Trefry et al., 1984; Wiesenburg et al., 1985) despite high concentrations of organic carbon and methane (Sackett et al., 1979). The most intense microbial activity is focused in the seawater-brine interface region (LaRock et al., 1979; Sheu et al., 1988; Van Cappellen et al., 1998; Wiesenburg et al., 1985), a particle trap where successive depletion of oxygen, nitrate and oxidized manganese has been observed (Trefry et al., 1984; Van Cappellen et al., 1998) along with narrowly defined peaks of methane, ethane, propane and hydrogen sulfide concentrations (Wiesenburg et al., 1985). These authors propose that accumulation of reduced gases and anoxia within the brine result from the long residence time of brine in Orca Basin combined with slow in situ decomposition of organic carbon (Wiesenburg et al., 1985).

The onset of brine accumulation in Orca Basin is thought to be signaled by a transition from shallow, black anoxic sediments to more deeply buried gray sediments. This transition, reported in both slope sediments and in deeper basin sediments (Addy and Behrens, 1980; Leventer et al., 1983; Meckler et al., 2008; Northam et al., 1981) has been dated by three methods: radiocarbon analysis of total carbonate (Addy and Behrens, 1980); radiocarbon analysis of foraminifera (Leventer et al., 1983; Meckler et al., 2008); and biostratigraphy (Leventer et al., 1983), yielding similar ages of approximately 7900 to 8500 calendar years for Orca Basin brine.

Here we use radiocarbon measurements of dissolved inorganic (DIC) and organic carbon (DOC) and box modeling to document slow, cumulative microbial processes driving organic carbon cycling within the basin.

2 Material and methods

2.1 Sampling

Brine from the southern lobe of Orca Basin and overlying seawater was sampled by CTD rosette at $26^{\circ}54.5'$ N, $91^{\circ}20.1'$ W from the R/V *Atlantis* during cruise AT18-2. Whole seawater was collected for radiocarbon analysis of DIC following the NOSAMS DIC sampling protocol (http://www.whoi.edu/NOSAMS_DIC_collection, accessed in 2010). Briefly, 500 mL of seawater was collected in pre-combusted glass bottles after thorough rinsing with sample seawater, poisoned with mercuric chloride and sealed with greased, ground-glass stoppers. Filtered DOC samples were collected directly from Niskin bottles into 1 L amber bottles through an in-line, pre-combusted 47 mm Whatman GF/F filter. Multiple filters were necessary to collect 200 to 300 mL brine samples. DOC bottles were capped with Teflon-lined lids and stored at -20° C until analysis.

2.2 DIC and DOC concentrations

Samples for DIC analysis were filter-sterilized (0.2 µm) and stored in gas-tight 2 mL autosampler vials at 4° C prior to analysis (Brandes, 2009) using liquid chromatographyisotope ratio-monitoring mass spectrometry (LC-IRMS) with a Delta V Plus. Samples were calibrated against sodium bicarbonate standards. Samples for DOC analysis were filtersterilized on board the ship $(0.2 \,\mu\text{m Target}^{\mathbb{R}}$ filters) and stored frozen at -20° C prior to analysis using a Shimadzu TOC-V high temperature catalytic oxidation system (Joye et al., 2010). As a precautionary measure, the Target[®] filters were rinsed by running 10-15 mL of UV-oxidized milliQ water through the filters and dried by passing 5–10 mL of dry sterile (0.2 µm filtered) air through the filter. Blanks for DOC, UV-oxidized milliQ filtered through similarly cleaned 0.2 µm Target[®] filters, were collected along side DOC samples in the field and run in the laboratory; blanks were consistently less than 0.2 µM.

2.3 δ^{13} C and Δ^{14} C analysis

Stable carbon isotopic analysis and radiocarbon analysis of DIC and DOC was performed at the National Ocean Sciences Accelerator Mass Spectrometry (NOSAMS) facility. Inorganic carbon was extracted from DIC samples with the water-stripping method (McNichol et al., 1994). δ^{13} C values were determined on a split of the sample CO₂ by dual-inlet Optima IRMS, and the remainder of the CO₂ was converted to graphite for radiocarbon analysis by AMS. This method has been successfully used to analyze DIC from brine samples previously (e.g., Ma et al., 2005; Mikucki et al., 2009). Ongoing analysis of reproducibility across all sample types at NOSAMS indicates measurement uncertainty is $\pm 0.05 \%$ for δ^{13} C values and $\pm 4.9 \%$ for Δ^{14} C of modern samples

(Elder et al., 1998; Jenkins et al., 2010). Independent δ^{13} C measurements of separate DIC samples were performed by LC-IRMS (as described in Sect. 2.2; Brandes, 2009) at Skidaway Institute of Oceanography (SkIO). Reproducibility of DIC standards by LC-IRMS method was ± 0.04 ‰.

DOC was converted to CO₂ by the UV oxidation method (Beaupré et al., 2007). Brine samples were diluted with preirradiated MilliQ water (dilution factors: 2.9, 3.0) and a split of sample CO₂ was reserved for δ^{13} C analysis by dual-inlet Prism IRMS. The remainder was graphitized for radiocarbon analysis by AMS. Sample results are all corrected for process blanks determined using the method of Currie et al. (2000) and Santos et al. (2007). Measurement of modern and dead standards indicates processing introduces the equivalent of $0.21 \pm 0.2 \,\mu$ mol modern carbon and $0.33 \pm 0.28 \,\mu$ mol dead carbon, and we observe an overall reproducibility of $\pm 0.3 \,\%$ for DO¹³C and ± 7 to 10 % for modern DO¹⁴C.

3 Results

3.1 Salinity and dissolved oxygen

Salinity and dissolved oxygen profiles are illustrated in Fig. 1. Oxygen depletion begins above the sharp salinity increase at approximately 2100 m, defining the upper limit of the seawater-brine interface region. Salinity increases sharply to the brine value at approximately 2240 m resulting in an estimated thickness of 140 m for the seawater-brine transition region. This is the same thickness reported by Sackett et al. (1979), suggesting the interface is very stable with little change observed since the late 1970s.

3.2 Dissolved Inorganic Carbon

DIC concentrations also show a similar pattern to that observed in 1976-1977 with a doubling of concentrations between overlying seawater (2.0-2.1 mM) and brine (3.9-4.1 mM; Sackett et al., 1979). This transition occurs over approximately 50 m, paralleling salinity rather than oxygen concentration (Fig. 1a). For both seawater and brine, DIC concentrations reported by Sackett et al. (1979) are slightly higher and more variable than our values (Fig. 1a). Smallscale gradients in DIC concentration are not expected because Orca Basin brine is convectively mixed (Shokes et al., 1977; Wiesenburg et al., 1985). Our low-precision estimates of DIC concentration obtained from two brine depths during CO₂ extraction for radiocarbon analysis also agree within analytical uncertainty arguing against DIC gradients within the brine. Combined, these observations give confidence to our DIC profile with its absence of small-scale variability throughout the depth profile.

By all three measures, brine DIC is significantly depleted in ¹³C relative to overlying seawater (Table 1, Fig. 1b), indicating substantial contributions from remineralized organic carbon or methane. Sackett et al. (1979) present a detailed profile documenting the transition between overlying seawater and brine δ^{13} C values, and here we describe the present isotopic state of the DIC reservoir using two independent methods: CO₂ extraction followed by offline δ^{13} C analysis by dual-inlet IRMS (NOSAMS) and LC-IRMS (SkIO). Both methods indicate a DIC reservoir that is more depleted in ¹³C than any value reported by Sackett et al. (1979). Similar to DIC concentrations, brine δ^{13} C_{DIC} values measured at NOSAMS agree within analytical uncertainty. δ^{13} C_{DIC} values determined at SkIO also agree closely within the brine, which contrasts with the ~4 ‰ range observed previously (Fig. 1). As described above, the small-scale variability recorded by Sackett et al. (1979) is not expected within a well-mixed brine lending credibility to our more recent analyses.

Despite internal consistency, our $\delta^{13}C_{DIC}$ values are significantly different than those reported previously and also significantly different from each other (Fig. 1b). We believe the 2 or 4 ‰ difference is more likely to be a result of a methodological bias rather than reflect a decadally dynamic DIC reservoir. Our parallel $\delta^{13}C$ measurements may differ due to the difference in chemical composition between brine samples and seawater calibration standards. Since we cannot reconcile our NOSAMS $\delta^{13}C_{DIC}$ values (-20.0±0.05‰) with values measured at SkIO (-18.1±0.1‰), or the average $\delta^{13}C_{DIC}$ value (-16.4±1‰) reported by Sackett et al. (1979), we include the average $\delta^{13}C_{DIC}$ value from both of our measurement methods as well as that reported by Sackett et al. (1979) in calculations and the Discussion below.

The radiocarbon contents of brine DIC at 2300 and 2473 m depth agree within analytical uncertainty, showing no variation with depth (Table 1, Fig. 1c). Our values are in general



Fig. 1. Depth profiles of salinity (gray solid line) and dissolved oxy-

gen concentrations (gray dashed line) are plotted for the south basin

of Orca Basin brine along with (A) concentrations of DIC (filled

circles, this study; open circles from Sackett et al., 1979) and DOC

(filled diamonds, this study; open diamonds from Sackett et al., 1979); (**B**) δ^{13} C values of DIC and DOC; and (**C**) Δ^{14} C values for DIC and DOC. Error bars are smaller than symbols for this study

 $(\pm 0.04-0.5 \text{ }\% \text{ for } \delta^{13}\text{C} \text{ values and } \pm 1-3 \text{ }\% \text{ for } \Delta^{14}\text{C} \text{ values}).$

Depth (m)	$\delta^{13} \mathrm{C}_{\mathrm{DIC}}(\%)$	±	$\delta^{13} \mathrm{C}_{\mathrm{DOC}} \ (\%)$	±	$\Delta^{14} \mathrm{C}_{\mathrm{DIC}} \ (\%)$	±	$\Delta^{14}\mathrm{C}_{\mathrm{DOC}}(\%)$	±
2050 ^b	0.93	0.04						
2050 ^a	0.81	0.05	-23.3	0.3	-93.1	3.6	-481.9	2.4
2130 ^b	0.72	0.04						
2150 ^b	0.41	0.04						
2160 ^b	-0.13	0.04						
2180 ^b	-1.17	0.04						
2190 ^b	-2.40	0.04						
2200 ^b	-3.27	0.04						
2210 ^b	-3.97	0.04						
2215 ^b	-4.49	0.04						
2220 ^b	-5.35	0.04						
2230 ^b	-6.25	0.04						
2240 ^b	-17.75	0.04						
2250 ^b	-18.20	0.04						
2300 ^b	-17.99	0.04						
2300 ^a	-20.07	0.05	-24.0	0.3	-515.5	2.5	-586.3	1.8
2473 ^b	-18.05	0.04						
2473 ^a	-19.97	0.05	-24.0	0.3	-520.0	2.1	-587.1	1.7

Table 1. Isotopic composition of Orca Basin brine and overlying water.

^a Analyzed at NOSAMS. ^b Analyzed at SkIO.

agreement with $\Delta^{14}C_{\text{DIC}}$ values reported previously (decay corrected to $-509 \,\%$, $-496 \,\%$ and $-492 \,\%$ from Sackett et al., 1979; Stuiver and Polach, 1977), although these values were not reported with sample depths. Both the $\Delta^{14}C$ and $\delta^{13}C$ values of DIC from overlying seawater are similar to World Ocean Circulation Experiment (WOCE) values from 2000–3000 m depth in the North Atlantic (WOCE line A22, 1997).

3.3 Dissolved organic carbon

DOC concentrations within the brine $(320-349 \,\mu\text{M})$ are greater than those in the overlying seawater $(50-59 \,\mu\text{M})$ by a factor of seven and are in good agreement with DOC concentrations reported by Sackett et al. (1979; Fig. 1a). The δ^{13} C values of DOC are also very similar to those reported by Sackett et al. (1979) and do not show any variation with depth. The carbon isotopic contrast between DOC in the brine and overlying seawater is considerably smaller than for DIC (Table 1, Fig. 1b and c). The DOC from overlying seawater is also isotopically similar to DOC measured in the North Atlantic at similar depths (Druffel et al., 1992).

4 Discussion

Comparison between our isotopic results and similar values reported three decades ago (Sackett et al., 1979) suggests that the Orca Basin lacks a dynamic carbon cycle. Multiple studies of the sediments underlying Orca Basin have converged on a 7900 to 8500 yr timescale for the establishment of anoxic, hypersaline conditions in the basin (Addy and Behrens, 1980; Leventer et al., 1983; Meckler et al., 2008). For this study, we compared our radiocarbon values with the most recent foram record reported by Meckler et al. (2008) because it includes a foraminiferal radiocarbon date in close proximity to the black-gray sediment transition depth and because radiocarbon analysis of individual foraminiferal species has confirmed stratigraphic continuity in this location (Flower et al., 2011). Adjusting these reported radiocarbon ages by the more recent, smaller reservoir age from the nearby Flower Garden Banks (Wagner et al., 2009) results in an 8700 ± 100 yr calendar age for Orca Basin brine. Establishment of the brine, however, is not necessarily coupled to the establishment of its dissolved carbon reservoir. This brine age requires a greater depletion of ¹⁴C than was actually measured in brine DIC, pointing towards the input of more modern carbon to the DIC reservoir in the Orca Basin.

4.1 Conservative mixing of brine and seawater end-members

The depth profiles of DIC and DOC concentrations closely parallel that of salinity through the seawater–brine interface of Orca Basin (Fig. 1a), and Fig. 2 illustrates the relationship between DIC, DOC and salinity across this interface. Conservative mixing between seawater and a brine end-member source is implied by the strong linearity observed between salinity and DIC as well as salinity and DOC concentrations. Small deviations from the conservative mixing line (Fig. 2) do allow for a limited DIC source in the low-salinity region of the seawater–brine interface where extensive organic



Fig. 2. Mixing diagrams for (A) DIC and (B) DOC concentrations across the seawater–brine interface plotted along with solid lines which represent expected concentrations from conservative mixing of brine and seawater end-members.

matter degradation has been reported (Tribovillard et al., 2008; Wong et al., 1985) as well as nonconservative behavior of DOC within the brine.

The isotopic composition of this brine end-member can be calculated by constructing an isotopic mixing model. Solving the following equations:

$$m_{\text{measured}} = m_{\text{seawater}} + m_{\text{brine}},\tag{1}$$

 $m_{\text{measured}} \cdot \delta_{\text{measured}} = m_{\text{seawater}} \cdot \delta_{\text{seawater}} + m_{\text{brine}} \cdot \delta_{\text{brine}}, \quad (2)$

$m_{\text{measured}} \cdot \Delta_{\text{measured}} = m_{\text{seawater}} \cdot \Delta_{\text{seawater}} + m_{\text{brine}} \cdot \Delta_{\text{brine}}, \quad (3)$

where the concentration (m_{brine}), δ^{13} C value (δ_{brine}) and Δ^{14} C value (Δ_{brine}) of the brine end-member calculated using the average measured brine values for m_{measured} , δ_{measured} and Δ_{measured} and values from overlying seawater at a depth of 2050 m for m_{seawater} , δ_{seawater} and Δ_{seawater} , yields an δ^{13} C estimate of -40 ± 3 ‰ (NOSAMS) or -36 ± 2 ‰ (SkIO) and a Δ^{14} C estimate of -930 ± 64 %. Using the δ^{13} C_{DIC} values reported by Sackett et al. (1979) indicate a brine endmember that is a bit less ${}^{13}C$ depleted (-33 ± 2 ‰). This approach, similar to that utilized by Sackett et al. (1979), suggests that brine DIC is supplied by a source depleted in both 13 C and 14 C. The δ^{13} C value of the brine end-member likely reflects a combination of POC remineralization with typical marine values of -21 to -22 % (Northam et al., 1981; Sackett et al., 1979) and oxidation of methane having a δ^{13} C value of -72 to -75 ‰ (Sackett et al., 1979; Wiesenburg et al., 1985). This conservative mixing model, however, is unable to account for the time-evolving nature of ¹⁴C concentration and is most representative of a single modern mixing event. The three similar estimates of brine age from slope (Leventer et al., 1983; Meckler et al., 2008) and deep basin sediments (Addy and Behrens, 1980) suggest that the current brine thickness accumulated quickly, making a single modern mixing event unlikely. If any single episode (i.e., of short enough duration that DIC does not "age" significantly) mixed seawater and end-member brine to produce the current brine composition, it would have occurred at the onset of brine accumulation, 8700 calendar years (or 8078 radiocarbon years) ago. However, a long-ago mixing event is incompatible with the radiocarbon age of brine DIC (\sim 5800 radiocarbon years), which is younger than the age of the brine. Thus we conclude that an isotopic mixing model cannot accurately represent the radiocarbon content of a brine end-member and that the true brine DIC source is likely to be significantly more modern than the value calculated above.

Although it does not represent the ¹⁴C contents of reduced carbon sources which ultimately contribute DIC to Orca Basin brine, δ^{13} C results from the isotopic mixing model highlight the significant contribution that microbial oxidation of reduced carbon compounds make to the inorganic carbon pool in Orca Basin. They also do not reveal any one dominant source of reduced carbon, rather suggesting a combination of methane and sinking organic carbon. A mass balance calculation according to the following equation:

$$\delta^{13}C_{\text{brine}} = \frac{[\text{DIC}]_{\text{seawater}}}{[\text{DIC}]_{\text{brine}}} \cdot \delta^{13}C_{\text{seawater}} + f_{\text{POC}} \frac{([\text{DIC}]_{\text{brine}} - [\text{DIC}]_{\text{seawater}})}{[\text{DIC}]_{\text{brine}}} \cdot \delta^{13}C_{\text{POC}} + (1 - f_{\text{POC}}) \cdot \frac{([\text{DIC}]_{\text{brine}} - [\text{DIC}]_{\text{seawater}})}{[\text{DIC}]_{\text{brine}}} \cdot \delta^{13}C_{\text{methane.}} (4)$$

Using the expected δ^{13} C values for methane and sinking POC described above indicates $65 \pm 6\%$ (NOSAMS) or $72 \pm 6\%$ (SkIO) of brine end-member DIC must come from POC remineralization and the remainder from methane oxidation. Using the average δ^{13} C_{DIC} value reported by Sackett et al. (1979) results in an estimate of $80 \pm 8\%$ of brine DIC deriving from POC remineralization.

4.2 Box-model estimates of brine DIC sources

Microbially mediated DIC addition to Orca Basin argues against the model of a single mixing episode and instead implies a slow accumulation of excess DIC and therefore an uncoupling between brine formation and DIC accumulation within the brine. To better capture this model of DIC buildup and constrain the radiocarbon age of the microbially contributed DIC, we constructed a box model to represent the evolution of DIC during the 8700 yr history of Orca Basin brine. Brine DIC can be contributed by the remineralization of sinking organic carbon ("modern", with an age of contemporaneous marine organic carbon from photosynthesis in the mixed layer), methane from the seawater-brine interface ("modern") or methane diffusing upwards from sediments ("aged"). We assume a constant rate of DIC increase added to an initial DIC concentration and isotopic composition equivalent to seawater overlying the Orca Basin brine. We do not explicitly include molecular diffusion as only picomolar DIC loss can be expected over the entire lifetime of the brine with a molecular diffusivity of $0.5 \times 10^{-9} \text{ m}^2 \text{ s}^{-1}$ (Zeebe, 2011). Eddy diffusion is likely to have a somewhat greater effect than molecular diffusion, although the sharp and stable DIC gradient at the brine interface (Fig. 1a) argues against any large diffusive DIC losses from the Orca Basin brine. Because we cannot measure eddy diffusivity for the brine interface and because the large density contrast between the brine and overlying seawater will inhibit turbulent diffusion making it unlikely to have a significant influence on the evolution of the DIC reservoir, we do not include DIC loss due to eddy diffusion in the box model. ¹⁴C decay is incorporated into each timestep following Eq. (7), and the fraction of POC and methane added at each time increment is determined by the fractions calculated in Sect. 4.1. We prescribe the $\Delta^{14}C$ value of sinking POC over the lifetime of the brine as atmospheric Δ^{14} C values according to IntCal09 (Reimer et al., 2009) offset by the reservoir age from the nearby Flower Garden Banks $(-53.2 \pm 1 \%)$; Wagner et al., 2009) to account for atmospheric ¹⁴C variability through time and solve iteratively for the Δ^{14} C of methane. The following equations describe the concentration and isotopic composition of DIC at time *t* in the box model:

$$[DIC]_{t} = [DIC]_{t-1} + \frac{[DIC]_{\text{final}} - [DIC]_{\text{initial}}}{t_{\text{final}} - t_{\text{initial}}},$$
(5)

$$\delta^{13}C_{t} = \frac{[DIC]_{t-1} \cdot \delta^{13}C_{t-1}}{[DIC]_{t}} + \frac{f \cdot ([DIC]_{t} - [DIC]_{t-1}) \cdot \delta^{13}C_{POC}}{[DIC]_{t}} + \frac{(1-f) \cdot ([DIC]_{t} - [DIC]_{t-1}) \cdot \delta^{13}C_{methane}}{[DIC]_{t}}, \quad (6)$$

$$[\Delta^{14} C_t]_{AGED} = \left[\left(\frac{\Delta^{14} C_t}{1000} + 1 \right) \cdot e^{\lambda(-5)} - 1 \right] \cdot 1000, \quad (7)$$

$$\Delta^{14}C_{t} = \frac{[DIC]_{t-1} \cdot [\Delta^{14}C_{t-1}]_{AGED}}{[DIC]_{t}} + \frac{f \cdot ([DIC]_{t} - [DIC]_{t-1}) \cdot \Delta^{14}C_{POC}}{[DIC]_{t}} + \frac{(1-f) \cdot ([DIC]_{t} - [DIC]_{t-1}) \cdot \Delta^{14}C_{methane}}{[DIC]_{t}}.$$
 (8)

Using the parameters in Table 2 (with either NOSAMS or SkIO value for final δ^{13} C) and a 5 yr time resolution, this model calculates a Δ^{14} C value of $-31 \pm 102 \%$ for DIC resulting from methane oxidation (Fig. 3). Substituting in the average δ^{13} C_{DIC} value reported by Sackett et al. (1979) changes the result by less than 1 ‰ because the calculated methane source has very similar radiocarbon contents to the average prescribed POC source. It also does not increase uncertainty in the model output due to the larger uncertainties assigned to the δ^{13} C values of POC and methane (Table 2). This "modern" result points to the brine interface as the dominant source region for methane that is oxidized by the microbial assemblage in Orca Basin, resulting in DIC with a radiocarbon value similar to sinking POC (time average: -18 %,



Fig. 3. Box modeled results for the Δ^{14} C value of DIC (solid black line) over the 8078 radiocarbon-year history of Orca Basin brine plotted along with dashed lines representing uncertainty in model results. The solid gray line represents Δ^{14} C values of POC input driven by IntCal09 (Reimer et al., 2009) adjusted for a local reservoir age from the nearby Flower Garden Banks (Wagner et al., 2009).

Fig. 3, gray line). Considering the slightly ¹⁴C-depleted value compared to time-averaged POC and considering the uncertainties of this approach (Fig. 3, dashed lines), model results do not exclude a small contribution from the oxidization of aged methane diffusing into the brine from sediments (Wiesenburg et al., 1985). The importance of the brine interface as a potential DIC source to the brine below is suggested by the multiple studies that point toward high rates of microbial activity focused in this region (LaRock et al., 1979; Sheu et al., 1988; Van Cappellen et al., 1998; Wiesenburg et al., 1985) and that document extensive degradation of organic matter at the seawater-brine interface (Tribovillard et al., 2008; Wong et al., 1985). And the small deviation of DIC concentrations from a conservative mixing line (Fig. 2) compared to the deviations observed for alkalinity and Mn²⁺ (Van Cappellen et al., 1998) lend support to the low-salinity region of the interface being a slow, cumulative source of DIC.

4.3 Isotopic composition and sources of brine DOC

In the seawater overlying Orca Basin, the contrast between the radiocarbon contents of DIC and DOC is 389 ‰, not significantly different from the 300–380 ‰ contrasts observed at similar depths in the Arctic (Griffith et al., 2012), Atlantic, Pacific and Southern Oceans (Druffel and Bauer, 2000). Within the Orca Basin brine, however, the difference between the radiocarbon contents of DIC and DOC is much smaller (~70 ‰, Table 1). This smaller contrast implies sources and sinks of Orca Basin DOC that are distinct from deep-ocean refractory DOC despite δ^{13} C values that are indistinguishable (δ^{13} C_{brineDOC} – δ^{13} C_{seawaterDOC} = -0.7 ± 1 ‰).

In addition to the similarity in radiocarbon contents, the δ^{13} C values of brine DIC and brine DOC are also strikingly

Symbol	Parameter	Value	±	Source
t ₀	initial time in years	0		
$t_{\rm f}$	final time in years	8078	74	uncalibrated radiocarbon age of brine
[DIC] ₀ (µM)	initial DIC concentration	2006	20	value from overlying seawater at 2063 m
[DIC] _f (µM)	final DIC concentration	4086	41	average brine [DIC]
$\delta^{13}C_0$ (‰)	initial δ^{13} C value	0.81	0.01	value from overlying seawater at 2050 m
$\delta^{13} C_{f}$ (‰)	final δ^{13} C value	-20.02	0.01	average brine $\delta^{13}C_{\text{DIC}}$ (NOSAMS)
		-18.08	0.11	average brine $\delta^{13}C_{\text{DIC}}$ (SkIO)
$\delta^{13}C_{POC}$ (‰)	sinking POC δ^{13} C value	-22	1	Holocene organic matter ^a
$\delta^{13}C_{\text{methane}}$ (‰)	methane δ^{13} C value	-74	2	average methane values ^b
$f_{\rm POC-methane}$	fraction POC	0.65	0.06	calculated in Sect. 4.1, Eq. (4)
λ	1/true mean-life	1/8267		Stuiver and Polach (1977)
$\Delta^{14}C_0$ (‰)	initial Δ^{14} C value	-93.1	3.6	value from overlying seawater at 2050 m
$\Delta^{14}C_{POC}$ (‰)	sinking POC Δ^{14} C value	-53.2	1.0	IntCal09 values ^c minus reservoir age ^d
$\Delta^{14} C_{f}$ (‰)	final Δ^{14} C value	-517.8	2.5	average brine $\Delta^{14}C_{DIC}$

Table 2. Box model parameters, values and uncertainties.

^a Northam et al. (1981); ^b Wiesenburg et al. (1985); ^c Reimer et al. (2009); ^d Wagner et al. (2009).

similar ($\delta^{13}C_{brineDOC} - \delta^{13}C_{brineDIC} = 4 \pm 2\%$ (NOSAMS), $6 \pm 2 \%$ (SkIO)). This δ^{13} C similarity is not typical of open ocean basins and suggests a stronger linkage between these two reservoirs in Orca Basin than observed in other parts of the deep ocean. DOC is enriched within Orca Basin brine by approximately sevenfold compared to overlying seawater (Fig. 1a). This is a greater enrichment than is observed for DIC and points to the importance of in situ brine DOC sources. These sources could include incorporation of DOC produced from DIC by chemoautotrophy at the brine interface (Jove et al., 2009), microbial conversion of POC to DOC and disaggregation of POC to DOC. The δ^{13} C value of DOC does not allow for a significant ¹³C-depleted source from methane oxidation as it is not more ¹³C-depleted than DOC in the water column above. A fraction of brine DOC is also likely to come from deep-ocean refractory DOC incorporated with bottom water upon brine formation, but this may only represent one seventh of total brine DOC.

Constructing a conservative mixing model of brine and seawater end-members would not be appropriate because DOC is likely to be a more dynamic reservoir than DIC with potential for microbial sinks as well as sources (Fig. 2b). Still, results of the DIC box model have implications for the radiocarbon contents of the in situ DOC sources. Box model results emphasize the importance of modern DIC sources (Sect. 4.2). Modern-day brine DIC and DOC have similar radiocarbon contents (Fig. 1c) but the greater enrichment of brine DOC concentrations compared to overlying seawater and the greater ¹⁴C-depletion of overlying DOC, and presumably original end-member DOC, compared to overlying DIC implies an even more dominant "modern" source to brine DOC than DIC. Along with evidence from δ^{13} C values, this further minimizes the potential for microbial interaction with aged methane to create brine DOC.

 Table 3. Modeled apportionment of DIC sources in Orca Basin brine.

Modern brine DIC sources	Contribution	$\Delta^{14}\mathrm{C}(\%)$	$\delta^{13}\mathrm{C}(\%)$
original seawater DIC oxidized sinking POC oxidized methane from interface oxidized methane from sediments	$49 \pm 1 \% 33 \pm 4 \% 18 \pm 3 \% 0 \pm 3 \%$	-93.1 ^a IntCal09 ^b IntCal09 ^b "aged" ^e	$^{+1^{a}}_{-22^{c}}_{-74^{d}}_{-74^{d}}$

^a Overlying seawater from 2050 m, Table 1; ^b Reimer et al. (2009); ^c Northam et al. (1981); ^d Wiesenburg et al. (1985); ^e assumed.

5 Conclusions

Both the concentration and isotopic composition of dissolved carbon reservoirs in Orca Basin brine appear to be stable over the past three decades. Modeled results point toward the importance of slow, cumulative and in situ microbial contributions to the inorganic carbon pool and emphasize the importance of the brine interface as the location of this microbial activity. Although both organic carbon and methane are implicated as the ultimate sources of brine DIC, box-modeled results show limited potential for contributions from the oxidization of methane diffusing upwards from sediments. The relative importance of individual DIC sources to Orca Basin brine is summarized in Table 3.

The isotopic composition of brine DOC is similar to brine DIC suggesting both a stronger coupling between these two reservoirs than is observed in overlying seawater and distinct sources and sinks for brine DOC compared to DIC. Stable carbon isotopic values only allow for a small DOC source from microbial incorporation of methane, unlike the proportionally important source expected for DIC. Aside from anaerobic methane oxidation, carbon isotopic evidence cannot indicate specific microbial pathways important in DOC or DIC cycling. It also cannot determine the limiting factor that prevents organic carbon remineralization by microbial sulfate reduction in Orca Basin brine. Further, microbial and compound-specific isotopic investigations of the brine and seawater-brine interface are needed to form a better understanding of specific microbial processes contributing to dissolved carbon storage in the Orca Basin brine.

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