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Composition of Dissolved Organic Matter in Pore Waters of Anoxic Marine Sediments Analyzed by $^1$H Nuclear Magnetic Resonance Spectroscopy

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Marine sediments are globally significant sources of dissolved organic matter (DOM) to the oceans, but the biogeochemical role of pore-water DOM in the benthic and marine carbon cycles remains unclear due to a lack of understanding about the molecular composition of DOM. To help fill this knowledge gap, we used $^1$H nuclear magnetic resonance (NMR) spectroscopy to examine depth variability in the composition of pore-water DOM in anoxic sediments of Santa Barbara Basin, California Borderland. Proton detected spectra were acquired on whole samples without pre-concentration to avoid preclusion of any DOM components from the analytical window. Broad unresolved resonance (operationally assigned to carboxyl-rich alicyclic molecules, or CRAM) dominated all spectra. Most of the relatively well-resolved peaks (attributed to biomolecules or their derivatives) appeared at chemical shifts similar to those previously reported for marine DOM in the literature, but at different relative intensities. DOM composition changed significantly within the top 50 cm of the sediment column, where the relative intensity of CRAM increased, and the relative intensity of resolved resonances decreased. The composition of CRAM itself also changed throughout the entire length of the 4.5-m profile, as CRAM protons became increasingly aliphatic at the expense of functionalized protons. Given that pore-water DOM is generated from sedimentary organic matter that includes pre-aged and degraded material, and that DOM is theoretically subject to microbial reworking in the pore waters for centuries to millennia, these data suggest that marine sediments may be sources of CRAM that are compositionally unique from CRAM generated in the upper ocean.

Keywords: pore water, DOM, carbon, NMR, sediment, anoxic, marine, Santa Barbara Basin
INTRODUCTION

Dissolved organic matter is produced as an intermediate in the mineralization of sedimentary organic matter (Burdige and Komada, 2015). This in turn supports a net efflux of dissolved organic carbon (DOC) out of the sediments to the oceans that rivals global riverine DOC export from land (Burdige et al., 1999). Previous studies that examined the composition and reactivity of pore-water DOM have highlighted the heterogeneity of this pool and the complexity of its dynamics within the sediment column (e.g., Henrichs and Farrington, 1979; Sansone and Martens, 1982; Alperin et al., 1994; Burdige and Zheng, 1998; Burdige, 2001; Hee et al., 2001; Komada et al., 2013; Burdige et al., 2016). While these studies have contributed significantly to our understanding of DOM cycling in sediments (see review by Burdige and Komada, 2015), the sheer molecular complexity of natural organic matter (Hedges et al., 2000; Hertkorn et al., 2007; Dittmar, 2015) poses a challenge toward addressing some basic questions, such as the role pore-water DOM plays in the degradation and preservation of sedimentary organic matter, and the biogeochemical significance of benthic DOM fluxes in the marine DOM cycle.

To help address these knowledge gaps, we used $^1$H NMR spectroscopy to examine the chemical composition of pore-water DOM in anoxic sediments of Santa Barbara Basin (SBB), California Borderland. While NMR has been widely used to investigate the composition and dynamics of seawater DOM (Minor et al., 2014; Repeta, 2015 and references within), application to marine pore-water DOM has been limited in number and scope. To the best of the authors’ knowledge, only two studies in the literature report NMR spectra of pore-water DOM from marine sediments (Orem and Hatcher, 1987; Repeta et al., 2002). Orem and Hatcher (1987) used solid-state $^1$H NMR to examine the composition of high-molecular-weight (HMW) pore-water DOM collected from two nearshore marine and estuarine locations, along with samples from a number of freshwater peats, and a coastal saline lake. Repeta et al. (2002) used solution-state $^1$H NMR to examine the chemical composition HMW pore-water DOM collected from two nearshore marine locations, and a continental shelf site. The primary focus of these pioneering studies was to compare and contrast NMR spectra across contrasting depositional environments (Orem and Hatcher, 1987), or between HMW-DOM in pore waters and the water column (Repeta et al., 2002); no effort was made to examine the cycling of pore-water DOM in the sediments. Orem et al. (1986) studied solid-state $^{13}$C NMR spectra of pore-water DOM in Mangrove Lake (a coastal saline lake) as a function of sediment depth, but the depth resolution was extremely low (three samples collected from intervals as thick as ~1.9 m over a total depth range of 5.5 m), and these authors did not detect any depth variability in their spectra.

To explore the linkages between pore-water DOM transformations and sediment organic matter degradation, we acquired $^1$H NMR spectra of pore-water DOM as function of depth in the top 4.5 m of the sediment column in the center of SBB. With the exception of a handful of cases that applied NMR spectroscopy to whole marine DOM samples (Lam and Simpson, 2008; Zheng and Price, 2012), NMR has primarily been used to examine marine DOM isolated by ultrafiltration (HMW-DOM), by solid-phase extraction (SPE-DOM), or by coupled reverse osmosis/electrodialysis (RO/ED-DOM). Neither ultrafiltration nor SPE recover 100% of the DOM pool, nor are they free of selective bias (Mopper et al., 2007; Minor et al., 2014; Repeta, 2015). HMW-DOM is rich in a class of semi-labile polysaccharides [acyl heteropolysaccharides (APS) or heteropolysaccharides (HPS); (Aluwihare et al., 2005; Hertkorn et al., 2006; Repeta and Aluwihare, 2006)], and overall HMW-DOM represents the relatively reactive fraction of the whole DOM pool in seawater (Benner and Amon, 2015; Walker et al., 2016a,b). This characterization also appears to hold for the HMW-DOM extracted from sediment pore waters (Repeta et al., 2002). SPE-DOM, when collected using PPL (a styrene divinylbenzene sorbent), concentrates DOM primarily based upon polarity (Dittmar et al., 2008), and is relatively rich in low-molecular-weight compounds and in carboxyl rich alicyclic molecules (CRAM) that are thought to represent a refractory component of marine DOM (Hertkorn et al., 2006; Dittmar and Stubbs, 2014; Repeta, 2015). Differences in the molecular composition of HMW-DOM and SPE-DOM are clearly reflected in $^1$H NMR spectra of these components extracted from the surface ocean, where HMW-DOM shows clear enrichment in APS/HPS relative to SPE-DOM (Hertkorn et al., 2006, 2013). Differences between HMW- and SPE-DOM are less obvious for deep-sea DOM (Hertkorn et al., 2006, 2013), consistent with the semi-labile nature of APS/HPS. Relative to ultrafiltration and SPE, RO/ED has been reported to recover a larger and more representative fraction of total DOM (Vetter et al., 2007; Gürttler et al., 2008; Kopriwnjak et al., 2009; Green et al., 2014; Helms et al., 2015), but the large sample volume requirement makes RO/ED unsuitable for the study of DOM in pore waters. A mini-electrodialysis system designed to handle small-volume samples has been developed, but appears to cause considerable DOC contamination (Chen et al., 2011).

To better understand the chemical composition of pore-water DOM, we applied $^1$H NMR to whole pore-water DOM without any isolation or pre-concentration steps. High ion strengths reduce the signal-to-noise-ratio in NMR, but this was overcome by DOC concentrations in sediment pore waters that are typically much higher than in seawater. By circumventing the physical and chemical separation of DOM from the sample matrix, we aimed to capture the depth dependent variability of the chemical composition of pore-water DOM without precluding any of the DOM components from the analytical window. However, it should be noted that our samples contain dissolved metals (including paramagnetic species such as iron) which can affect NMR relaxation, and hence peak width (Satterlee, 1990). The concentration and composition of such metals may also have varied with sediment depth (e.g., Shaw et al., 1990; Severmann et al., 2006). Furthermore, suppression of the water signal can attenuate the resonances in the regions attributed to carbohydrates and olefins (Lam and Simpson, 2008). We therefore used the data primarily to examine relative changes in DOM composition within our dataset, and secondarily to compare to published $^1$H NMR
spectra of HMW-, SPE-, or RO/ED-DOM extracted from seawater.

The main objective of this study was to use $^1$H NMR to examine the variability in pore water DOM composition as function of depth in SBB sediments. We further used our findings to evaluate the reactivity of pore water DOM at this site.

MATERIALS AND METHODS

Sampling Site and Sample Collection

Sediment cores were collected from the center of Santa Barbara Basin (SBB) in the California Borderland (34.223°N, 119.983°W) at a water depth of 590 m in August 2012 aboard the R/V Robert Gordon S. Sproul, and August 2013 aboard the R/V New Horizon. Bottom-water dissolved oxygen concentrations at this site are low (<2 µM), and the fine-grained sediments are anoxic and free of bioturbation (Hüseleman and Emery, 1961; Sholkovitz and Gieskes, 1971; Soutar and Crill, 1977). Details about sediment coring and pore water sampling can be found in Komada et al. (2016). Briefly, cores penetrating as deep as ~450 cm below the sediment-water interface were collected by gravity- and multicoring, and sectioned within 2–9 h of recovery to collect pore water. Samples for NMR analysis were passed through disposable 0.2-µm nylon filters (Whatman 6870-2502; pre-cleaned with 100 mL of UV-irradiated deionized water), collected into pre-combusted glass ampules, headspace purged with ultra-high purity N$_2$, and then flame sealed and refrigerated until further processing. Pore-water DOC data obtained from these cores (along with other geochemical data such as porosity, sedimentary particulate organic carbon content, and inorganic pore-water solutes) have been presented and discussed in detail elsewhere (Burdige et al., 2016; Komada et al., 2016); the DOC data are reproduced here in Figure 1. A total of 37 samples that spanned the entire length of this DOC profile were analyzed in the present study by $^1$H NMR (Figure 1).

$^1$H NMR Analysis

Proton detected spectra were acquired on two Bruker Avance III spectrometers: one operating at 400 MHz at the College of Sciences Major Instrumentation Cluster, Old Dominion University, and another operating at 600 MHz at Lawrence Livermore National Laboratory, both using Bruker 5 mm BBO broadband—$^1$H/X z-gradient probes, and TopSpin 3.0 software. For analysis on the 400 MHz spectrometer, 0.4-mL pore-water aliquots were added to 0.04 mL D$_2$O and 3 µL tetramethylsilane (TMS) in 5-mm glass NMR tubes (Wilmad Glass Co., NJ), and analyzed within 30 min of preparation. For analysis on the 600 MHz spectrometer, samples were prepared similarly, but placed in high-resolution, 5.0-mm Norell tubes, back-filled with argon gas, flame sealed, and refrigerated at 4°C for up to 1 week prior to analysis. Of the 37 samples collected, 21 (from 0.5 to 432 cm) were analyzed on the 400 MHz NMR instrument, and the remaining 16 samples (from 95 to 435 cm) were analyzed on the 600 MHz NMR instrument (Figure 1). Tetramethylsilane was used as an NMR chemical shift reference. $^1$H NMR spectra were collected with solvent suppression using a modified version of the WATERGATE-W5 suppression sequence described by Lam and Simpson (2008). The total acquisition time was 15 h 2 s, with a 2 s relaxation delay at ~11,000 scans and 4 Hz exponential line broadening.

Data Processing

All $^1$H NMR spectra were first normalized by total area under the spectra, then specific area integrals were taken using two approaches. First, relative abundances of major $^1$H types were determined by integrating the areas in chemical shift (δ$_{^1}$H) ranges defined by Hertkorn et al. (2013; Table 1). Second, specific regions of the spectra whose relative intensity varied significantly with depth ($p \leq 0.05$) were objectively identified by 2D correlation using sediment depth as the perturbation variable (Abdulla et al., 2010, 2013b). In this 2D correlation approach, the area under the broad unresolved band between 0.91 and 3.15 ppm was operationally assigned to CRAM (Hertkorn et al., 2006), and areas of peaks that appear atop this broad band were determined after subtracting the CRAM signal from the total area. In all integrations, area under any given portion of the spectrum was calculated by trapezoidal approximation with chemical shift increments of 0.0055 ppm. Peaks were deemed absent when they amounted to < 0.2% of the total area.

Protons corresponding to individual peaks identified by 2D correlation were classified according to literature data, and tentatively assigned to major compound classes (Table 2).
Abundances of these compound classes were further expressed in DOC units by assigning H/C ratios to each compound class (Table 2, Supplementary Material). Exchangeable protons in imino, amino, carboxyl, and hydroxyl protons are not observed by solvent-suppression $^1$H NMR due to the fast exchange rate of these protons with deuterium (from $D_2O$).

## RESULTS

### Comparison of Spectra Obtained at Different Field Strengths

Common resonances were present in the spectra acquired using the 400- and 600-MHz spectrometers, but with two key differences. First, there was a clear resolution advantage in the higher field magnet (Figure 2, Supplementary Figure 1). Second, relative intensities of singly-oxygenated units including carbohydrate (HCO, 3.1–4.1 ppm) and olefinic proton (4.9–7.0 ppm) regions were higher in the spectra obtained on the 600 MHz spectrometer shown (Figure 2), possibly due to suppression of the water peak that was broadened from radiation damping. Radiation damping perturbs relaxation times of protons in the solvent (in the this case, water) which are present in extreme excess over protons in the analyte (in this case, DOM; Krishnan and Murali, 2013). Radiation damping results in spectral artifacts including line broadening and the amplitude of this effect is dependent on field strengths and pulse power (Krishnan and Murali, 2013). Thus, when the broader water peak from the 600 MHz instrument was suppressed, it resulted in intensification of the carbohydrate-like and olefinic regions on either side of the suppressed water peak relative to the signal acquired on the 400 MHz instrument, precluding direct comparisons of data obtained on the two instruments (Figure 2). However, when the spectra are normalized only between $\delta_{^1H}$ 0.2–3.1 ppm (aliphatic and functionalized protons only), there is good agreement between the data sets (Supplementary Figure 1). Quantitative analyses were done using only spectra obtained on the 400 MHz instrument, because they covered a greater depth range than those from the 600 MHz instrument (Figure 1). Spectra from the 600 MHz instrument are presented for qualitative purposes only, although the two data sets did display similar depth trends.

## $^1$H NMR Spectra of Whole Pore-Water DOM

$^1$H NMR spectra of whole pore-water DOM featured general characteristics that are common to spectra reported for SPE-, HMW-, and RO/ED-DOM extracted from seawater (e.g., Hertkorn et al., 2006, 2013; Dittmar et al., 2008; Koprivnjak et al., 2009; Abdulla et al., 2013a; Repeta, 2015). All spectra obtained here showed broad, unresolved resonances attributed to aliphatics ($\sim$0–1.9 ppm), functionalized protons including CRAM ($\sim$2–3 ppm), singly-oxygenated units including carbohydrate ($\sim$3–4 ppm), and to a lesser extent olefinic and aromatic protons ($\sim$5–10 ppm; Figure 2). Broadness of these peaks reflects significant signal overlap of many protons of similar resonance frequencies, consistent with a complex, heterogeneous mixture of molecules. Atop the broad envelope were relatively well-resolved peaks (or clusters of peaks) that have been observed in SPE-DOM (e.g., Dittmar et al., 2008; Hertkorn et al., 2013), HMW-DOM (e.g., Hertkorn et al., 2006; Abdulla et al., 2013a; Repeta, 2015), and RO/ED-DOM (Koprivnjak et al., 2009) albeit at different relative intensities: (1) at $\sim$1.2 ppm (attributed to methyl proton in 6-deoxysugar; Quan and Repeta, 2007); and (2) at $\sim$2.1 ppm (attributed to acetate derivatives). The latter peak is particularly prominent in HMW-DOM in surface seawater and in semi-labile polysaccharides (e.g., Hertkorn et al., 2006; Abdulla et al., 2013a; Repeta, 2015).

Examined more closely, $^1$H NMR spectra obtained here exhibited several peaks within the aliphatic range (0–1.9 ppm) that were previously observed in RO/ED-DOM spectra, but not in SPE- and HMW-DOM spectra (this is particularly evident in the spectra acquired on the 600 MHz spectrometer shown in Figure 2B). First, in addition to the $\sim$1.2 ppm peak that typically dominates this range of the SPE- and HMW-DOM $^1$H
of SBB sediment pore waters, but overall showed little depth variability.

### Synchronous 2D Correlation Analysis

To further investigate the depth variability in the $^1$H NMR spectra of pore-water DOM, we used 2D correlation to objectively identify chemical shift ranges whose relative intensities changed significantly with depth ($p \leq 0.05$). All but one of the distinct features in the 0.2–3.1 ppm range of these spectra appeared as an auto-correlation peak (autopeak) along the diagonal of the 2D correlation map, indicating that their intensities changed significantly with depth (Figure 4). The exception was the shoulder centered around 0.7 ppm, whose absence along the diagonal indicated that its depth variability was not significant. Autopeaks along the diagonal were arbitrarily labeled C–M (Table 2). Two of these autpeaks (broad bands G and K centered around 1.6 and 2.5 ppm, respectively) fell in regions of the spectra that featured only unresolved resonance without a clear peak (Figure 2). These were therefore considered subsections of the broad CRAM envelope, and from hereon, are referred to as “bands” instead of “peaks.” Accordingly, areas of bands G and K were determined by calculating the total area under the spectrum over the corresponding $\delta$H ranges.

Cross peaks that appear in the off-diagonal in the 2D correlation map allowed gross separation of peaks/bands into two groups: Group-1, which consisted of peak J, band G, and band K; and Group-2, which consisted of peaks C, E, H, L, and M, and band N (Figure 4). Major positive cross peaks (red)
were observed among Group-1 resonances, indicating that these peaks/bands co-varied with depth. Cross peaks within Group-2 were also positive, but the correlation was not as strong as compared to Group-1 resonances. Cross peaks between Groups 1 and 2 were negative (green).

**Peak Integration and Depth Profiles**

To better visualize variability in DOM composition, areas under peaks C–M were integrated and plotted against sediment depth. The area corresponding to CRAM protons was also calculated by summing the areas under the unresolved envelope (Figure 5, Supplementary Figure 2). The majority of compositional variation occurred within the top 50 cm, where relative intensities of Group-1 resonances (J, G, K) and CRAM increased, and relative intensities of Group-2 resonances (C, E, H, L, M, N) decreased (Figures 3, 5). Band O declined with depth, similar to Group-2 resonances (Figure 3). Peak L was absent below 150 cm as was peak M in two deep samples. Other than variability near the surface, band P showed no clear depth trend (Figure 3).

Protons assigned to CRAM dominated throughout the profile, accounting for 50% of the total integral in the uppermost sample to 70% at depth (Table 3; Figure 5). Bands G and K amounted to 5–7% and 15–25% of CRAM, respectively. While the relative abundance of CRAM protons increased with depth by ~30%, areas of bands G and K doubled, suggesting that these are more reactive regions of the CRAM envelope. The next most prominent resonances after CRAM were bands N and P, which accounted for 9–15% and 4–9% of the total integral, respectively (Table 3).

Upon conversion of relative proton abundance to DOC concentration (Supplementary Material), all peaks/bands showed a general increasing trend with depth (Figure 6; Supplementary Figure 3), reflecting the fact that DOC increased steadily with depth (Figure 1). Group-1 resonances (J, G, K; Table 2), CRAM, and band P increased smoothly with depth, suggesting steady net production of these substructures in the sediment column. In contrast, Group-2 resonances (Table 2) showed greater depth variability. Peaks E, H, L, and M showed clear enrichment in the uppermost sample, possibly due to high rates of production near the sediment-water interface. Peak H showed an acute drop immediately below the core top, and then a step-like increase at ~150 cm near the base of the sulfate-methane transition zone, a major redox-cline.

Relative abundances of these peaks/bands expressed in DOC units were similar to those calculated using proton resonance intensities, except for a small boost in olefins and aromatics due to their low assigned H/C ratios (Table 3). When binned into major compound classes, CRAM accounted for 53–75% of total DOC, followed by carbohydrate (11–21%; C, J, L, N), aromatics (6–14%; P), olefins (4–9%; O), and protein/peptide (1–5%; E, M). This dominance of CRAM (~50–70% of protons and DOC) is comparable to, or higher than, what has been reported for HMW-DOM in surface and deep seawater (23% and 50% of 13C NMR signal, respectively; Hertkorn et al., 2006), SPE-DOM in freshwater (~62% of 1H NMR signal in Lake Ontario surface water; Lam et al., 2007), and SPE-DOM in surface and deep seawater (50 and 56% of 13C NMR signal, respectively; Hertkorn et al., 2013).

**DISCUSSION**

Composition and Inferred Reactivity of Pore-Water DOM in SBB

The chemical composition of pore-water DOM clearly varied with depth (Figure 4), with the greatest change occurring in the top ~50 cm (Figures 3, 5). This variability can be broadly characterized as a relative increase in the unresolved CRAM envelope, that was counterbalanced by relative decreases in
Group-2 resonances (peaks C, E, H, L, M, band N) and band O. Band P (aromatic protons) showed little depth variability in relative intensity. Regardless of the sources and identities of the compounds that give rise to Group-2 resonances and band O, their declining relative intensities and the increasing dominance of the unresolved CRAM envelope is consistent with an overall increase in molecular diversity with depth in the sediment column, and therefore, with increasing microbial organic matter degradation during early diagenesis.

**Group-2 Resonances**

Group-2 resonances identified by 2D correlation analysis (Figure 4) included a number of peaks (C, E, H, L, M; Table 2). Peaks are expected to arise from specific substructures, making it possible to link them to particular biomolecules or their derivatives. Bands N and O have also been characterized to arise from biomolecules (carbohydrate-like structures, and linear terpenoids and unsaturated alicyclics, respectively; e.g., Hertkorn et al., 2013). Relative intensities of all Group-2 resonances and band O declined sharply in the top ~50 cm of the sediment column (Figures 3, 5), suggesting that these substructures are associated with labile compounds that undergo rapid turnover near the sediment surface. However, with the exception of peak L which was not readily discerned below 135 cm, all of these resonances persisted throughout the profile. This then means that Group-2 resonances and band O are not only associated with labile compounds that occur only near the sediment-water interface, but also components of more stable molecules that accumulate with depth in the sediment column. This overall pattern is also evident upon conversion of relative intensities to absolute DOC concentrations (Figure 6). All Group-2 units and band O increased with depth in the sediment column, consistent with net accumulation of semi-refractory to refractory DOM constituents (Hansell, 2013; Burdige et al., 2016). In addition to net accumulation with depth, DOC concentrations corresponding to peaks E, H, L, and M were highest at the sediment surface (0.5 cm), and reached low (and in some cases near-constant) values within the top 50 cm (Figure 6). This type of enrichment at the sediment-water interface has been observed for highly labile DOM such as dissolved free amino acids (Henrichs and Farrington, 1987; Burdige and Martens, 1990) and is predicted by diagenetic model calculations (Burdige, 2002; Burdige et al., 2016). The depth distribution of Group-2 resonances described here also appears consistent with net accumulation of carbohydrates directly determined in sediment pore waters (Burdige and Komada, 2015).

Peak L (tentatively assigned to amino sugar; Table 2) was prominent in samples at depths <135 cm, but was largely non-detectable in deeper horizons (c.f., Figure 2; Supplementary Figure 2). The depth at which peak L was no longer observed roughly coincides with the base of the sulfate-methane transition zone (SMTZ) in SBB sediments (125 ± 10 cm; Komada et al., 2016) that demarcates the sulfate-reducing zone above and methanogenic sediments below. SMTZs in diffusion-dominated sediments are characterized as regions of high rates of microbial activity including anaerobic methane oxidation, and with microbial assemblages that are unique compared to adjacent sulfate-reducing and methanogenic sediments (Parkes et al., 2005; Biddle et al., 2006; Harrison et al., 2009). The near-absence of peak L below the SMTZ suggests that this peak is associated with molecules that are either produced only within the sulfate-reducing zone, or that its rate of loss exceeds the rate of production within the methanogenic zone.

Relative intensity of peak H (tentatively assigned to methanethiol functional group) also showed a break in distribution across the SMTZ (Figure 5); it was higher by a factor of two in the methanogenic zone (~145 cm) than in the SMTZ and the sulfate-reducing zone (p < 0.01; the top sample at 0.5 cm was excluded from this comparison). This subsequently resulted in a step-like increase in the absolute concentration of peak H in DOC units across the SMTZ (Figure 6). Peak H was the best resolved of all resonances detected in this study and was observed in all samples (c.f., Figure 2). This peak is also clearly visible in the 1H NMR spectra of RO/ED-DOM extracted from the surface waters of the western Atlantic ocean (Koprivnjak et al., 2009). However, to the best of the authors’ knowledge, this peak has
not been identified in the literature, nor has it been reported in
$^1$H NMR spectra of SPE- or HMW-DOM. More work is needed
to evaluate the identity and biogeochemical significance of the
compound(s) associated with this peak.

Resonances Attributed to Carbohydrates (C, J, N)
Polysaccharides make up a major component of marine and
freshwater HMW- and SPE-DOM (Hertkorn et al., 2006; Lam
et al., 2007; Repeta, 2015), and sediment pore-water DOM
(Burdige and Komada, 2015). A considerable fraction of this
compound class has been hypothesized to occur in the form of
N-acetyl amino polysaccharides, or acyl heteropolysaccharides
(APS; Aluwihare et al., 1997, 2005). The widespread occurrence
of APS in DOM is supported by the uniformity in the
hydrolysable neutral monosaccharide composition, and by the
recurring features in $^1$H and $^{13}$C NMR spectra obtained from
HMW-DOM collected from a wide range of environments
including freshwaters and sediment pore waters (Repeta et al.,
2002; Repeta, 2015). As expected, the spectra obtained here
consistently show resonances centered around $\delta_H$ 1.2 ppm (peak
C), 2.1 ppm (peak J), and between 3 and 4 ppm (band N),
assigned here to methyl group of deoxysugars, methyl group of
N-acetyl amino sugars, and singly oxygenated units including
carbohydrates, respectively (Table 2; Figure 2). Combined, these
peaks account for 10–20% of bulk DOC (Table 3), similar to
the relative abundances reported previously for lakes and
rivers (Repeta et al., 2002). The relative abundances of these
substructures in our whole pore-water samples in DOC units
(C: J: N = ~1:1:6; Table 3) are also similar to the ratio reported
for seawater and marine pore-water HMW-DOM (~1:1:8;
Aluwihare et al., 1997; Repeta et al., 2002).

In SBB pore-water DOM, it is however unlikely that C, J,
and N resonances belong to a common macromolecule as
hypothesized in the past (Aluwihare et al., 1997, 2005), because
their relative intensities showed distinct depth variability. 2D
correlation analysis shows that the relative intensities of C and
N declined with depth, while J increased with depth (Figure 4A;
also see Figures 3, 5). The relative increase in peak J with depth
suggests comparatively greater stability of N-acylated moieties
relative to their non-acylated counterparts. This finding supports
the work of Abdulla et al. (2010) who used 2D correlation to
analyze $^{13}$C NMR and FTIR spectra of HMW DOM collected
across an estuarine salinity gradient. They found polysaccharides
to be split into two groups: an acylated component (APS),
and a non-acylated component (heteropolysaccharides, or HPS).
Furthermore, they found relative abundance of HPS to vary with
salinity, while the relative abundance of APS remained largely
invariant, leading them to hypothesize that HPS is more dynamic
than APS.

CRAM
CRAM increased with sediment depth in terms of both relative
proton abundances (Figure 5) and in absolute amount as DOC
(Figure 6). This increase was also associated with changes in
composition, as evidenced by the fact that the relative
intensities of CRAM constituents G and K doubled, while the
relative intensity of CRAM resonance as a whole increased
by only ~30% over the length of the profile (Figure 5). To
more closely examine how CRAM composition varied with
depth, the CRAM envelope (Figure 2) was sectioned into
the $\delta_H$ intervals corresponding to peaks C, E, H, J, L, and
M in addition to bands G and K. Area integrals of these
sections were expressed as fractions of total CRAM area and
plotted as function of depth (Figure 7; the sum of these
sections accounted for ~94% of the total CRAM envelope).
The topmost 3 samples of the profile (from 0.5 to 11 cm)
were excluded from this analysis, because they exhibited large
fluctuations likely due to low signal-to-noise ratio resulting from
FIGURE 5 | Depth variability in the fraction of total area attributed to peaks (C–M) and CRAM obtained on the 400 MHz spectrometer. Bands G and K were considered to be components of CRAM. Depth profiles of bands N, O, and P are given in Figure 3. Analogous profiles obtained on the 600 MHz spectrometer are shown in Supplementary Figure 2. Dashed lines indicate the upper and lower boundaries of the sulfate-methane transition zone (SMTZ, 125 ± 10 cm; see section Group-2 Resonances).

TABLE 3 | Relative peak intensities observed in spectra acquired on the 400 MHz instrument.

| Peak/Band | C | E | H | J | L | M | N | O | P | CRAM |
|-----------|---|---|---|---|---|---|---|---|---|------|
| % areaa   | 2–6 | ≤3 | 1–2 | ≤1 | ≤2 | 9–15 | 3–6 | 4–9 | 2–5 | 8–17 | 50–70 |
| % DOCb    | 1–5 | ≤3 | 1–2 | ≤1 | ≤2 | 8–13 | 4–9 | 6–14 | 2–5 | 9–18 | 53–75 |

a Fraction of the total area under the normalized spectrum. Ranges reflect minimum and maximum values observed in the profile.

b Fraction of total DOC. Ranges reflect minimum and maximum values observed in the profile.

the low DOC concentration near the sediment surface (e.g., Figures 1, 2).

The fraction of CRAM protons resonating in the aliphatic δH ranges (corresponding to peaks C and E, and band G) increased significantly with depth (p < 0.05), while protons resonating in functionalized δH ranges (corresponding to peaks K and L, and band M) decreased significantly with depth (p < 0.05; Figure 7). Overall, slightly over half of the CRAM protons fell in the functionalized category, but this dominance declined as the relative abundance of aliphatic protons increased with depth (Figure 7). CRAM protons that resonated near the boundary between aliphatic and functionalized δH ranges (corresponding to peaks H and J) did not change significantly with depth (p < 0.05), and accounted for a ∼constant fraction of the total CRAM envelope.

These results clearly indicate that CRAM (operationally defined here as the broad envelope in the chemical shift region 0.91–3.15 ppm) is dynamic not only in terms of its total concentration, but also in its molecular composition. While functionalized protons dominated the total CRAM envelope throughout the profile, their decline relative to aliphatic protons suggest that CRAM that are rich in aliphatic protons may contain more stable molecules than CRAM rich in functionalized protons.

Bulk Chemical Composition of SBB Pore-Water DOM Relative to Seawater DOM

NMR studies of seawater DOM have shown widespread occurrences of two major DOM components: APS/HPS, the semi-labile component; and CRAM (Repeta, 2015). The relative contributions of these two components vary with water depth, and by the method used to extract DOM from seawater (Benner et al., 1992; Aluwihare et al., 2002; Hertkorn et al., 2006; Koprivnjak et al., 2009; Broek et al., 2017). However, both the molecular composition of APS/HPS (Aluwihare et al., 1997), and the shape of the broad CRAM envelope (e.g., Hertkorn et al., 2006; Lam et al., 2007; Lam and Simpson, 2008) show close agreement across marine and some freshwater environments.
Therefore, $^1$H NMR spectra of HMW-DOM and SPE-DOM commonly show strong similarity within each group (e.g., Hertkorn et al., 2013; Repeta, 2015).

$^1$H NMR spectra of pore-water DOM from SBB sediments reported here showed key differences compared to spectra of HMW-, SPE-, and RO/ED-DOM in the literature listed above. Possible explanations for this observation are: (1) SBB pore-water samples were analyzed as whole, whereas HMW-, SPE-, and RO/ED-DOM were subject to fractionation during isolation and concentration prior to NMR analysis; (2) SPE- and RO/ED-DOM were subject to chemical alteration during sampling, but SBB pore-water DOM was not; (3) SBB pore-water DOM is compositionally different from seawater DOM; or (4) SBB pore-water DOM spectra are influenced by the occurrence of dissolved metals (Satterlee, 1990) and by suppression of the water signal (Lam and Simpson, 2008), while other are not. These four factors are not mutually exclusive.

Despite these differences, SBB pore-water DOM and seawater DOM clearly have common constituents. First, SBB pore-water DOM displayed resonances that are also ubiquitous in marine DOM (peaks C, J, and band N). Second, the relative abundances of major $^1$H types observed in SBB pore-water DOM are similar to those reported for seawater SPE-DOM in the eastern Atlantic Ocean (Hertkorn et al., 2013; Table 1). Third, the overall decline in relative abundance of functionalized protons, and relative increases in HCO and olefinic protons with increasing depth in the sediment column (Figure 3) are similar to depth trends reported for the eastern Atlantic Ocean SPE-DOM (Hertkorn et al., 2013), implying overall similarity in relative reactivity across different $^1$H types in the water column and sediments.

**Net Production of CRAM and the Reactivity of Pore-Water DOM in Santa Barbara Basin Sediments**

The majority of the compositional change we detected by $^1$H NMR occurred within the top 50 cm of the sediment column (Figure 5) with a key characteristic being an increase in relative abundance of CRAM, and decrease in relatively well-resolved peaks. The dramatic increase in CRAM resonance in the top 50 cm of the sediment column where remineralization rates are high (Burdige et al., 2016) strongly indicates net production of CRAM by the microbial community during sedimentary organic matter degradation. This corroborates the findings from laboratory experiments (Koch et al., 2005, 2014; Rossel et al., 2013; Lechtenfeld et al., 2015) and field observations (Schmidt et al., 2017) showing the importance of microbial activity in the generation of CRAM-like formulas from known biomolecules.

At depths $>\sim 50$ cm, there was comparatively less variability in DOM composition (Figure 5). A simple and plausible explanation for this muted variability in bulk composition is that the substructures detected at depth are largely associated with molecules having limited reactivity. This suggestion agrees with recent modeling work (Burdige et al., 2016) that found labile DOC and dissolved organic nitrogen (with first-order
degradation rate constant \( k = 30 \) to >200 yr\(^{-1} \) to disappear within the top \( \sim 60 \) cm, and a refractory component \( (k \sim 2 \times 10^{-4} \) yr\(^{-1} \)\) to continuously accumulate throughout the depth interval examined. At any given depth, the refractory component dominated the DOC standing stock (70–80% in the top 10 cm, and >99% below the SMTZ). Given the fact that CRAM likely consists of a highly diverse set of molecules (Hertkorn et al., 2006), limited compositional change in CRAM-dominated pore-water DOM at depth is also consistent with the molecular diversity hypothesis for DOM stability (Dittmar, 2015).

The microbial pump has been put forward as the mechanism for production of refractory DOM in the ocean (Jiao et al., 2010). While a number of studies indeed demonstrate microbial conversion of labile monomers and fresh biomass into more stable compounds that resemble CRAM (Gruber et al., 2006; Rossel et al., 2013; Lechtenfeld et al., 2015), more recent studies suggest that actual production of refractory DOM found in the deep sea may require more than a few months to years of microbial processing (Koch et al., 2014; Osterholz et al., 2015). The work presented here shows that there is net production of CRAM (Figure 6), and compositional change in CRAM within the sediment column (Figure 7). If this compositional change is a result of CRAM transformation occurring within the pore water, then the time scale for this process over the 4.5-m length profile could be on the order of 2 kyr (DOM diffusional time scale calculated from the Einstein-Smoluchowski relation assuming that the whole sediment diffusion coefficient for DOM is 50 cm\(^2\) yr\(^{-1} \); Boudreau, 1997). Furthermore, pore-water DOM is generated during degradation of sedimentary organic matter with a wide range of reactivities including microbially reworked and pre-aged components (Ohkouchi et al., 2002; Blair and Aller, 2012; Komada et al., 2013; Burdige et al., 2016). In sum, marine sediments might be sources of CRAM that are compositionally unique from CRAM generated in the upper ocean.

SBB sediments are clearly sources of DOM to the overlying water column (Figure 1), including CRAM (Figure 6). If a major fraction of DOM accumulating in these pore waters—and hence supporting the benthic DOM flux—is indeed refractory, one might then conclude that these sediments are sources of refractory DOM to the water column (also see discussions in Burdige et al., 1999, 2016). However, this would depend on the extent to which DOM that exhibits limited reactivity in anoxic sediments also shows limited reactivity in the water column. For example, the presence of molecular oxygen in the water column may speed up the degradation of some compounds (e.g., Hulthe et al., 1998; Sun et al., 2002; Reimers et al., 2013). On the contrary,
differences in microbial enzymatic capabilities (Arnosti, 2000, 2008; Teske et al., 2011; Cardman et al., 2014), or the effect of dilution (Arrieta et al., 2015) could render some compounds less susceptible to degradation upon export into the water column. Rigorous assessment of the role of sedimentary DOM in the marine DOM cycle will require structural characterization of DOM molecules allowing for cross-identification in sediments and in the water column.

AUTHOR CONTRIBUTIONS

HA, CF, and JL: generated the NMR data, and HA conducted the 2D correlation analysis; DB and TK: coordinated the project and collected the samples; CF and TK: wrote the paper with intellectual feedback from all authors.

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SUPPLEMENTARY MATERIAL

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Conflict of Interest Statement: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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