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RECEIVED 30 March 2023

ACCEPTED 03 October 2023

PUBLISHED 20 October 2023

## CITATION

Redick MA, Cummings ME, Neuhaus GF,  
Ardor Bellucci LM, Thurber AR and  
McPhail KL (2023) Integration of  
untargeted metabolomics and microbial  
community analyses to characterize  
distinct deep-sea methane seeps.  
*Front. Mar. Sci.* 10:1197338.  
doi: 10.3389/fmars.2023.1197338

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# Integration of untargeted metabolomics and microbial community analyses to characterize distinct deep-sea methane seeps

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Deep-sea methane seeps host highly diverse microbial communities whose biological diversity is distinct from other marine habitats. Coupled with microbial community analysis, untargeted metabolomics of environmental samples using high resolution tandem mass spectrometry provides unprecedented access to the unique specialized metabolisms of these chemosynthetic microorganisms. In addition, the diverse microbial natural products are of broad interest due to their potential applications for human and environmental health and well-being. In this exploratory study, sediment cores were collected from two methane seeps (-1000 m water depth) with very different gross geomorphologies, as well as a non-seep control site. Cores were subjected to parallel metabolomic and microbial community analyses to assess the feasibility of representative metabolite detection and identify congruent patterns between metabolites and microbes. Metabolomes generated using high resolution liquid chromatography tandem mass spectrometry were annotated with predicted structure classifications of the majority of mass features using SIRIUS and CANOPUS. The microbiome was characterized by analysis of 16S rRNA genes and analyzed both at the whole community level, as well as the small subgroup of Actinobacteria, which are known to produce societally useful compounds. Overall, the younger Dagorlad seep possessed a greater abundance of metabolites while there was more variation in abundance, number, and distribution of metabolites between samples at the older Emyr Muil seep. Lipid and lipid-like molecules displayed the greatest variation between sites and accounted for a larger proportion of metabolites found at the older seep. Overall, significant differences in composition of the microbial community mirrored the patterns of metabolite diversity within the samples; both varied greatly as a function of distance from methane seep, indicating a deterministic role of seepage. Interdisciplinary research to understand microbial and metabolic

diversity is essential for understanding the processes and role of ubiquitous methane seeps in global systems and here we increase understanding of these systems by visualizing some of the chemical diversity that seeps add to marine systems.

#### KEYWORDS

methane seep, deep-sea, untargeted metabolomics, microbial community, tandem mass spectrometry, environmental analysis

## 1 Introduction

Methane seeps are areas where methane leaks from vast subseafloor reservoirs, fueling dense communities of organisms. Methane seep communities, including endemic microorganisms and animals, either harness the chemical energy being released from the seafloor directly or through symbiosis, or make use of the habitat (both physical and chemical) created by seep biological processes. Alongside endemic species, methane seeps host organisms that are found throughout the margin, including species that are harvested commercially (Grupe et al., 2015; Seabrook et al., 2019). The seeps themselves are a mosaic of different habitats with varying biogeochemistry and fauna (Boetius and Suess, 2004; Bernardino et al., 2012) in which communities change over time. Over the geologic time scales that methane seepage can occur, the metabolic activity of microbial communities results in carbonate precipitation that can pave vast areas of the seafloor (Cordes et al., 2009; Bowden et al., 2013). The resulting hard substrate, termed authigenic carbonate, provides for attachment and growth of anemones, sponges, and corals at mature methane seep sites in the otherwise expansive soft sediment of the deep ocean. Methane seeps enhance biological diversity of the ocean margin by producing substrate, hosting divergent biological processes and biogeochemistry, and contributing significantly to primary production in the deep ocean. Here we add to this foundational knowledge by revealing chemical diversity, which has implications for future societal use.

Methane seeps comprise a diversity of habitats in close proximity, creating a unique opportunity for natural products (NPs) discovery. The high biological diversity of seeps is illustrated in a recent study focused on the habitat along the Oregon and Washington margins in which more than 72,000 different amplicon sequence variants (ASVs) were documented from six seep and non-seep collection sites, plus a control site (Cummings et al., 2023). At the three methane seep sites sampled, 11,631 different amplicon sequence variants (ASVs) were present only in the on-seep core samples. While on-seep cores had relatively low Chao1 diversity values, with a mean of  $452 \pm 204$  for Emyr Muil and Dagorlad seeps, the 5m off-seep cores had local peaks in diversity with values of  $861 \pm 162$ . Even in earlier studies where ASVs with 97% similarity are combined into Operational Taxonomic Units (OTUs), more than 30,000 distinct OTUs have been measured at methane seep habitats (Seabrook et al., 2018).

Oxygen, sulfide (toxic to most animals), and redox gradients typify seep habitats, and the microbial communities reflect this diverse environment through an abundance of metabolic pathways, including aerobic methane oxidation, anaerobic oxidation of methane, sulfate reduction, sulfide oxidation, manganese and iron reduction, and nitrogen fixation (Levin, 2005; Reeburgh, 2007; Knittel and Boetius, 2009; Dekas et al., 2014). Sediment communities, including those at seeps, use or must exist in the presence of metabolites produced by co-occurring taxa as defense chemicals, transient chemical signals, and/or in a multitude of possible undefined ecological roles (Dekas et al., 2016; Patin et al., 2017).

Knowledge of the diverse metabolic strategies at methane seeps resulting in a diversity of pathways involved in cellular energy production and compound biosynthesis is far from comprehensive. Most studies investigating environmental microbiomes have focused on phylogenetic identity and abundance of microbial community members, without metabolomic analyses or testing for biological activity or potential ecological roles. Technological advances in high resolution mass spectrometry and associated data analysis tools have facilitated and been driven further by targeted and untargeted metabolomics coupled to microbial community analysis in microbiome studies (Bauermeister et al., 2022), with applications in human health (Aleti et al., 2022; Zhang et al., 2023) and agriculture (Rothman et al., 2019; Sun et al., 2022; Nie et al., 2023). With continued development of computational methods to annotate structurally complex specialized metabolites in complex mixtures (Morehouse et al., 2023), these integrated approaches are increasingly applicable to detection and structural annotation of microbial specialized metabolites in environmental microbiome samples (Shaffer et al., 2022), although structural annotations in untargeted mass spectrometry experiments are putative assignments of structural class or planar molecular structure that require follow-up for valid molecular structure characterization.

Limited material from environmental sampling poses a challenge for traditional NP isolations of new compounds for structure elucidation and biological activity testing. However, untargeted metabolomics analyses of environmental samples using high resolution liquid chromatography tandem mass spectrometry (LCMS<sup>2</sup>) may require only nanograms of chemical extract from milligrams of collected sample. This approach provides data for comparative analyses when working with limited materials. Importantly, the identities of metabolites detected via untargeted

analyses remain largely unknown, with an estimated 10% average annotation rate possible using LCMS spectral databases and computational prediction tools (de Jonge et al., 2022).

Untargeted LCMS<sup>2</sup> surveys coupled with computational tools permit generation of hypotheses from exploratory data to direct further investigation and prioritization of organisms or metabolites for targeted analyses. Open access tools such as GNPS, SIRIUS, and CANOPUS enable predictions of metabolite structure or structure class based on tandem mass spectrometry (MS<sup>2</sup>) fragmentation patterns. GNPS is a web-based mass spectrometry platform for sharing and comparative networking of MS<sup>2</sup> data (Wang et al., 2016). The downloadable software framework SIRIUS (Dührkop et al., 2021) incorporates CSI : FingerID and CANOPUS for prediction of molecular formulas and structure classifications using MS<sup>1</sup> isotope and MS<sup>2</sup> fragmentation patterns (Djombou Feunang et al., 2016; Kim et al., 2021). Here we present the results of a coupled metabolome-microbiome approach using untargeted high-resolution LCMS<sup>2</sup> analysis and bacterial 16S rRNA amplicon sequencing of environmental samples to assess metabolite detection and explore the chemical and biological relationships between two methane seep habitats that differ significantly in their gross morphology (Figure 1). The 16S rRNA sequence data used for our comparative analyses of the Emyn Muil and Dagorlad methane seeps comprise a subset of the larger ASV dataset reported by Cummings et al. (2023).

The Dagorlad seep site is an expansive microbial mat, likely indicative of a recent onset of seepage, and the Emyn Muil seep site is a large carbonate outcrop consistent with seepage that has been ongoing for hundreds to thousands of years. These two sites potentially represent the opposite ends of seep succession. By comparison with nearby off-seep habitats, as well as a deep-sea non-seep control site, we address the overarching question of whether different seep microbiomes can be distinguished by

untargeted metabolomics of environmental samples. We also present a correlation analysis of the subset of putative peptides with actinobacterial 16S rRNA sequence since Actinobacteria are well-known to produce biologically active peptides (Zhao et al., 2018). Our results thus may provide preliminary insight on the potential for discovery of new biotechnological or biopharmaceutical compounds from deep-sea methane seeps. Our long term goal is to better understand the role of seeps in marine biological diversity and future pharmaceutical and biotechnological research.

## 2 Methods

### 2.1 Sample collection

Sediment cores were collected from two methane seeps along the Cascadia Margin using the ROV Hercules on *E/V Nautilus* cruise NA121, in partnership with the Ocean Exploration Trust. Here we focused on one core collected directly at the seep and one core collected 5 m away from areas of active seepage at each of two contrasting seep sites (Figure 1). These areas of active seepage had the highest microbial diversity present in the region (Cummings et al., 2023). An additional core from the Slope Base site of the Ocean Observatories Initiative (OOI) Regional Cabled Array was collected at a water depth of 2,900 m by ROV JASON during routine maintenance of the site (44.5152°N, -125.3898°W). The OOI site located 300 km south of the other sampling sites was included to compare a deeper non-seep site in the same geographical region. Sediment push cores were extruded shipboard and sectioned into 1 cm-thick sections ranging from 0 (surface) to 10 cm deep in the core, with the exterior discarded to prevent sediment smearing from impacting the measured microbial community and metabolome. Core sections were

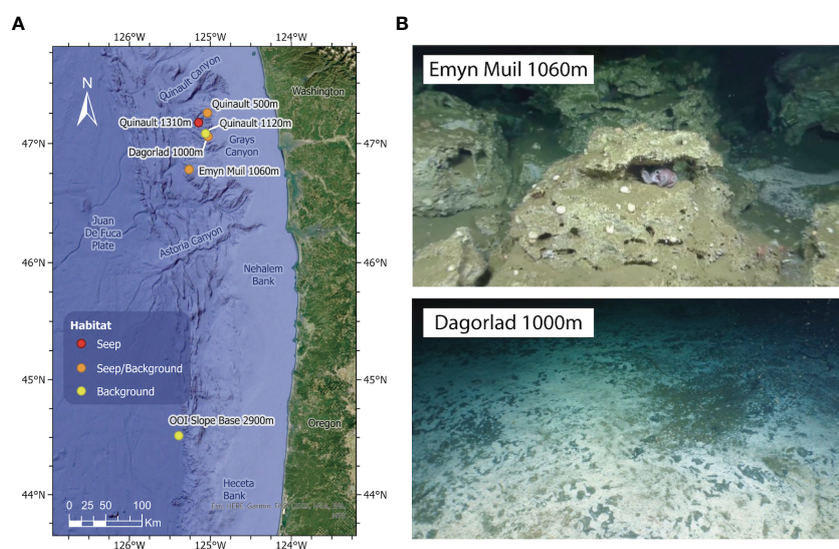


FIGURE 1

(A) Map indicating the locations of the three sampling sites (older seep Emyn Muil, younger seep Dagorlad, OOI Slope Base control). The two seeps are approximately 50 miles off the Washington coast on the Juan de Fuca Plate. (B) Images of the two methane seep sites sampled: upper is the older seep, Emyn Muil (1,060 m), characterized by carbonate deposits; lower is the younger seep, Dagorlad (1,000 m), characterized by soft sediments and extensive microbial mats. This figure is reproduced in part with permission from Cummings et al. (2023, <https://doi.org/10.7717/peerj.15119/fig-1>).

subdivided for metabolite and 16S rRNA analysis and frozen at  $-80^{\circ}\text{C}$  until analysis. Samples for metabolite profiling were only handled with metal core slicing devices and were stored frozen in glass vials.

## 2.2 Microbial community analysis

Microbial analysis of sediment samples was performed as described in [Seabrook et al. \(2018\)](#). In brief, DNA was extracted from  $\sim 0.5$  g frozen sediment using the QIAgen PowerSoil kit according to the kit protocol. The V4 region of the 16S rRNA gene was amplified and sequenced following the Earth Microbiome protocol, including the updated primers ([Parada et al., 2016](#)). Sequencing was carried out at the Oregon State University Center for Quantitative Life Sciences on the Illumina MiSeq platform, with 250-bp paired-end reads. Bioinformatics of the 16S rRNA analysis was carried out as part of a larger study using a Qiime2 pipeline as described in [Cummings et al. \(2023\)](#). The actinobacterial taxa from the resulting ASV table were selected for separate analyses.

## 2.3 Chemical extraction and LCMS<sup>2</sup>

Each sediment core subsection (48 in total) was lyophilized, resuspended in HPLC grade methanol (15 mL), sonicated (30 min) and left to extract overnight at room temperature. The solvent supernatants were then filtered using  $0.2\ \mu\text{M}$  PTFE syringe filters, dried *in vacuo*, and the resulting 48 extracts were weighed. High-resolution LCMS<sup>2</sup> profiling of all samples was performed using an Agilent 1260 Infinity II LC system with detection by an Agilent 6545 TOF mass spectrometer. For this analysis, the sample extracts were reconstituted at 10 mg/mL in LCMS grade methanol of which  $5\ \mu\text{L}$  were injected. A reversed phase C18 porous core column (Kinetex C18,  $50 \times 2.1$  mm,  $2.6\ \mu\text{m}$  particle size,  $100\ \text{\AA}$  pore size, Phenomenex, Torrance, USA) was used for chromatographic separation. A binary gradient system with mobile phase consisting of  $\text{H}_2\text{O}$  (solvent A) and acetonitrile (ACN, solvent B), and flow rate of  $0.4\ \text{mL}/\text{min}$  was used. After injection, the samples were eluted with a linear gradient from 0–5.5 min at 15% B, 5.5–15 min 15–80% B, 15–20 min 100% B. Initially the first 5.5 minutes were sent to waste but as no peaks were seen until after 8 minutes, 7.5 min of eluent was sent to waste to divert salt from the mass spectrometer for subsequent runs. Data dependent acquisition (DDA) of MS<sup>2</sup> spectra was performed in positive mode.

## 2.4 Chemoinformatic analysis of metabolomic data

For LCMS<sup>2</sup> data analysis, raw data (Agilent .d format) were converted to .mzML files using MSconvert (ProteoWizard); data pre-processing to extract MS<sup>1</sup> and MS<sup>2</sup> features was performed using MZmine3 ([Pluskal et al., 2010](#)). The parameters used in MZmine3 are listed in [Supplementary Table S1](#). Mass spectrometric data are available via MassIVE (MassIVE ID MSV000090543) ([MassIVE, n.d.](#)). The MZmine3 peak list was exported as an .mgf file and uploaded to SIRIUS 5 GUI ([Dührkop et al., 2019](#)). SIRIUS

([Dührkop et al., 2019](#)), CSI : FingerID ([Dührkop et al., 2015](#)), and CANOPUS ([Djoumbou Feunang et al., 2016](#); [Dührkop et al., 2021](#); [Kim et al., 2021](#)) analyses were performed using the parameters in [Supplementary Table S2](#). The MZmine3 generated feature list and quantification table were also analyzed using the GNPS platform ([Wang et al., 2016](#)). Peak areas of LCMS (MS<sup>1</sup>) features were summed by CANOPUS-assigned NPClassifier pathways to generate a pie chart for each core, representing the distribution of features across major structure classes ([Figure 2](#)). The fatty acid and terpenoid pathways were combined to form “Lipids and Lipid-like molecules”. Peak areas of LCMS (MS<sup>1</sup>) features were summed by CANOPUS-assigned NPClassifier pathways and stacked bar plots ([Figure 3](#)) showing the proportions of each pathway by core/depth were generated using the R package ggplot2 ([Torondel et al., 2016](#); [Wickham et al., 2019](#); [R Core Team, 2022](#)).

## 2.5 Statistical analyses of microbial and metabolomic data

Comparative analyses were performed on the microbial ASV, actinobacterial ASV, and LCMS ion feature peak area datasets using multivariate analysis in PRIMER v7 ([Clarke and Gorley, 2015](#)). Data were  $\text{Log}(X+1)$  transformed and Bray–Curtis similarity was used to construct a resemblance matrix. Significant patterns in the resulting data were identified via similarity profile (SIMPROF) analysis, which is also available in R package. SIMPROF identifies significant differences within datasets without *a priori* groupings and without an arbitrary cutoff of overall similarity. We used a significance level for SIMPROF groupings of 5%. We generated SIMPROF groupings for each MS<sup>2</sup>, total bacterial community, and actinobacteria only. We also visualized these data using non-metric multidimensional scaling (NMDS). Data were processed to visualize significant Spearman’s correlations between LCMS features and microbial ASVs in heatmaps using the R ([R Core Team, 2022](#)) script provided in the supplemental data repository and performed previously ([Neuhaus et al., 2022](#)). In brief, the MS ion feature list was filtered to generate a list containing only the subset of features annotated as potentially peptide by CANOPUS in either ontology. Correlation analysis with the family level microbial data was then performed separately for this peptidic metabolite subset in which features (microbial family or MS feature) observed in less than 10% of samples were removed prior to correlation. Microbial amplicon sequencing data were transformed to proportions by dividing the counts in each sample by the sample’s total sequencing depth. The dataset was mean centered and scaled before Spearman’s correlations were calculated (R package “Hmisc”) between each pair (microbe~metabolite) of features. P values were adjusted according to the Benjamini-Hochberg procedure to control the false discovery rate. Adjusted p values  $< 0.05$  were considered significant.

## 3 Results

High resolution LCMS<sup>2</sup> and 16S rRNA amplicon sequence data were curated for each of ten core sections from four sediment cores (40 samples in total). One core was collected on-seep and one core

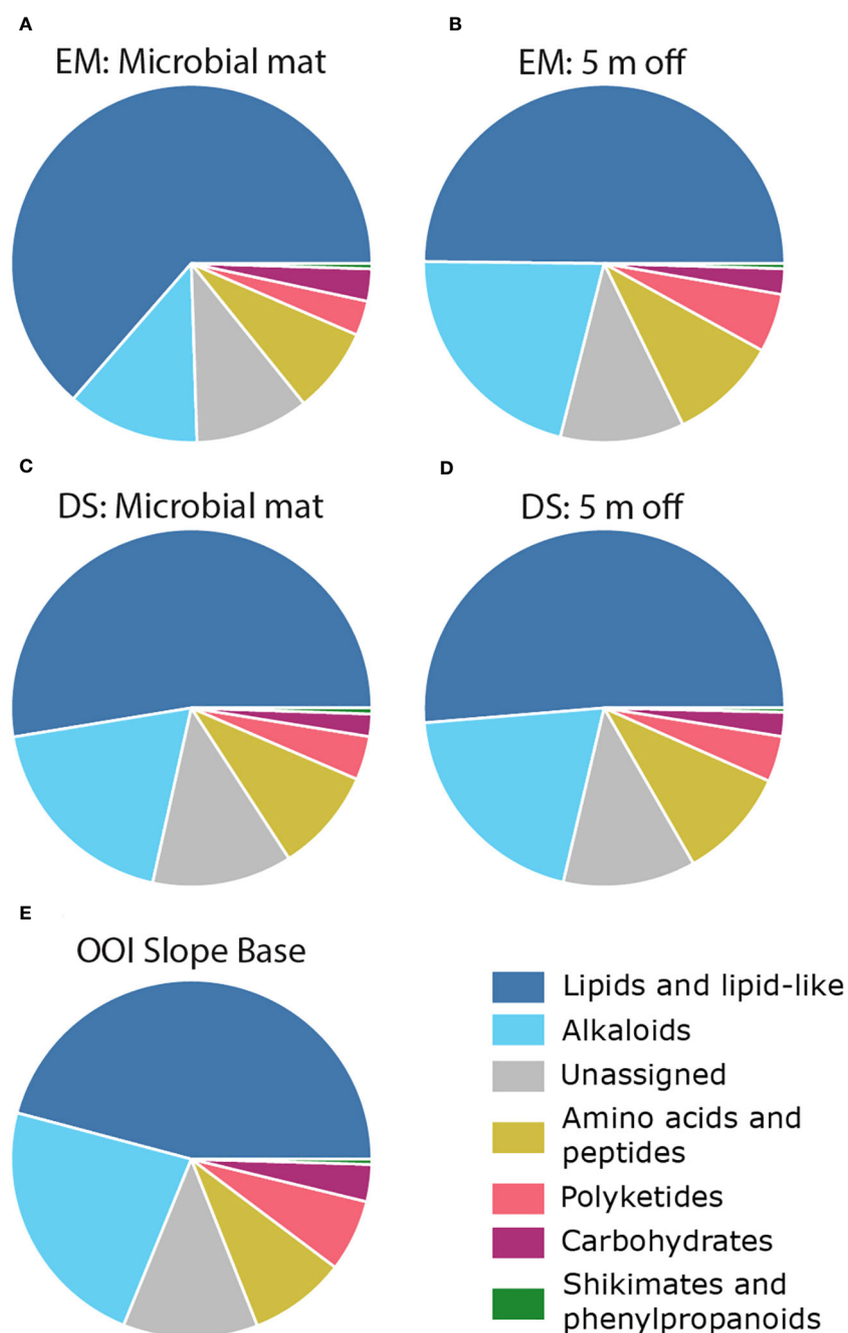


FIGURE 2

Pie charts of all MS features assigned to structural classes based on NPClassifier Pathways designated by CANOPUS. Terpenoids and fatty acids have been combined to form the class of “lipids and lipid-like” compounds. The plots display proportions based on MS<sup>1</sup> peak areas summed by sample collection site. (A) Emyn Muil (EM) on-seep microbial mat. (B) Emyn Muil (EM) 5 m off-seep. (C) Dagorlad Seep (DS) microbial mat. (D) Dagorlad Seep (DS) 5 m off-seep (E) Non-seep control site, OOI slope base 2900 m.

was collected 5 m off-seep from each of two seep sites with distinct geomorphologies (Figure 1) and variable geochemistry (Cummings et al., 2023). Paired data were similarly obtained for depth sections of a non-seep OOI core as a background control (8 samples in total). Throughout this manuscript we use the term “older” for the carbonate-dominated Emyn Muil seep and “younger” for the soft-sediment Dagorlad seep as it provides context for the results, however, we do not know the time of onset of seepage at the sites sampled. At both sites, soft sediment habitats were sampled,

although the sediment cores collected from the Emyn Muil seep were next to an expansive carbonate outcrop indicative of persistent seepage in the region for hundreds to thousands of years.

### 3.1 Microbial community

A total of 1,423,130 sequences belonging to 17,217 microbial ASVs were detected from the four methane seep sediment cores

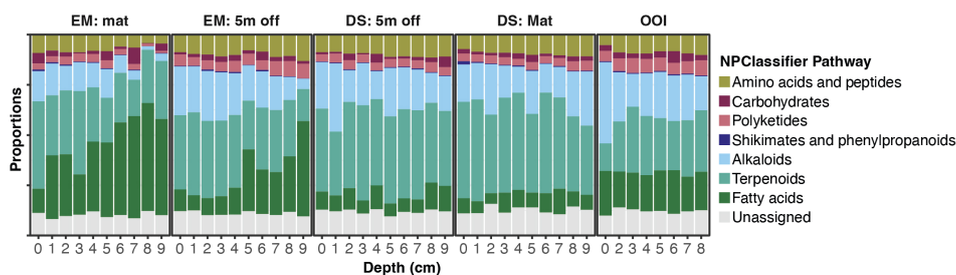


FIGURE 3

Stacked bar graphs displaying proportions of MS features assigned to structural classes based on NPClassifier Pathways designated by CANOPUS for core sections from each of the 5 different cores. EM: Mat is Emyn Muil on-seep mat core; EM: 5 m off is Emyn Muil 5 m off-seep core; DS: Mat is Dagorlad Seep on-seep mat core; DS: 5 m off is Dagorlad Seep 5 m off-seep core; OOI: non-seep is the OOI slope base core.

selected. The ASVs comprised 686 families in total. Actinobacterial sequences were a minor component of the total ASVs present. Overall, 2,046 sequences were identified within the Actinobacteria, representing 142 distinct ASVs. The Actinobacteria ASVs represented 27 families in total, although two families accounted for 85% of all ASVs, with half (54%) of those belonging to WCHB1-81 and the others falling outside of currently identifiable families.

### 3.2 Untargeted metabolomics

Processing of LCMS<sup>2</sup> raw data with MZMine3 resulted in a total of 1,304 mass features, which were assigned to 119 out of the 672 NPClassifier (Kim et al., 2021) structural classes across all seven NP biosynthetic pathways using CANOPUS (Dührkop et al., 2021). The metabolite profiles for cores from both methane seep sites (Figures 2A–D) exhibited a higher proportion of lipids and lipid-like molecules than the non-methane seep OOI site (Figure 2E). The chemistry of the microbial mat at the older seep with hard carbonate substrate clearly comprised the highest proportion of lipids and lipid-like molecules of all the sites (Figure 2A). The distribution of metabolites among NPClassifier pathways was most similar between the 5 m off-seep cores from each site (Figures 2B, D). While the relative proportion of features assigned to each structure class did not change markedly for non-lipid metabolites, the specific metabolites within each class varied significantly between sites.

Stacked bar plots of MS<sup>1</sup> feature peak areas summed by their NPClassifier assignments display the proportions of each biosynthetic class by depth in each core (Figure 3). The plots for both cores from Emyn Muil displayed the highest overall proportion and greatest variation in fatty acids and terpenoids, with the proportion of fatty acids generally increasing with depth in core. The on-seep microbial mat had a remarkably high proportion of fatty acids, and a concomitantly lower proportion of alkaloids and amino acids/peptides, especially with increasing depth in the core. In contrast, the Dagorlad cores displayed the lowest abundance of fatty acids with proportionally more terpenoids, and a relatively uniform distribution of metabolite biosynthetic classes between the on-seep microbial mat and 5 m off-seep cores. These data are summarized in Figure 4, in which

MS feature counts, summed peak areas and NPClassifier classes are summarized in parallel for each core by depth. The older Emyn Muil seep displayed greater variability in the abundance (peak areas, Figure 4A) and number (counts, Figure 4B) of LCMS features, with deeper core sections containing fewer and less abundance of features than shallower sections. In contrast, the number and abundance of LCMS features in both the control OOI slope base and the younger Dagorlad seep samples were less variable across depths. The number of distinct metabolite structural classes annotated for each sample using the NPClassifier ontology in CANOPUS was also determined and revealed consistent trends with depth (Figure 4C). Comparison of the shallower samples at the two seep sites revealed similar numbers and abundance of LCMS features (Figures 4A, B), with fewer structural classes assigned within the deeper samples from the older Emyn Muil seep (Figure 4C).

### 3.3 Integrated analysis of microbial and metabolite data

Separate NMDS plots of total microbial ASV and MS<sup>1</sup> data for all core samples exhibited corresponding patterns of variation. The subset of actinobacterial ASV abundances was also analyzed separately from the rest of the microbial ASV data because of the known capacity of Actinobacteria for specialized metabolism, resulting in structurally diverse compounds with antibiotic and other biological activities (Zhao et al., 2018).

Greatest variability in both metabolites (Figure 5A) and microbes (Figure 5B) was evident around the older carbonate Emyn Muil seep compared to the relatively clustered samples of the younger soft sediment Dagorlad seep, and the distinct microbial community and chemistry of the non-seep OOI slope base. We note that each point represents both a core location and depth within that core, making statistical analysis using *a priori* groupings (i.e. PERMANOVA) inappropriate to apply. Hence our application of exploratory significance testing using similarity profile (SIMPROF) analysis. The SIMPROF method examines whether the similarities observed in the data are smaller and/or larger than those expected by chance (null hypothesis testing). The MS<sup>1</sup> data showed more

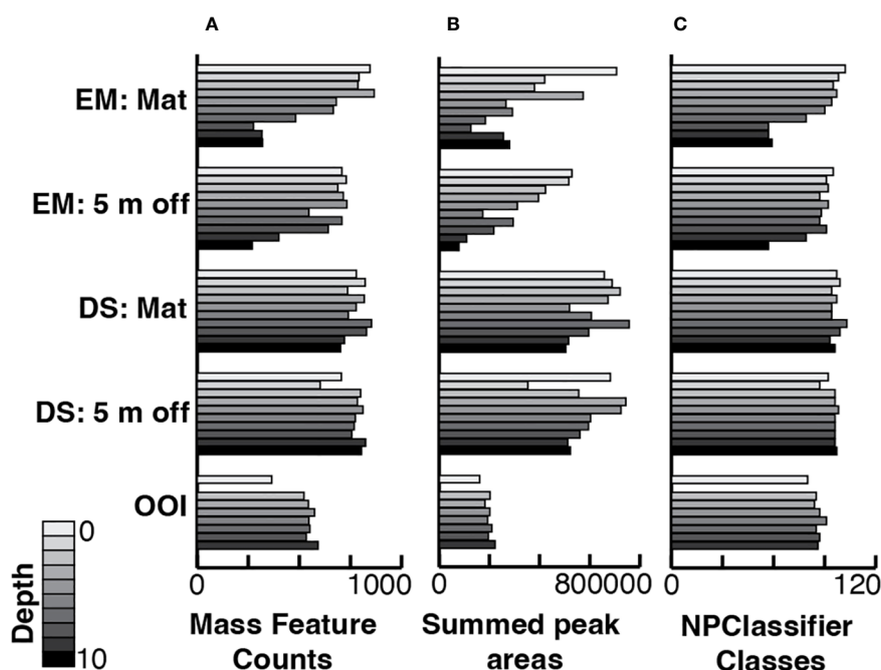


FIGURE 4

Bar graphs displaying per sample, stacked by core and sample depth in core, (A) the number of mass features, (B) the sum of all LCMS peak areas from the feature list generated in MzMine3, and (C) the number of distinct NPClassifier classes in each sample. EM: Mat is Emyu Muil on-seep mat core; EM: 5 m off is Emyu Muil 5 m off-seep core; DS: Mat is Dagorlad Seep on-seep mat core; DS: 5 m off is Dagorlad Seep 5 m off-seep core; OOI: non-seep is the OOI slope base core.

variability in both feature counts (on-seep mat:  $602.9 \pm 233.6$ ; 5 m off-seep:  $613.5 \pm 160.2$ ) and abundances (on-seep mat:  $3.48E5 \pm 1.78E5$ ; 5 m off-seep:  $3.05 \pm 1.61E5$ ) at the older Emyu Muil seep, with deeper sections yielding lower feature counts and abundances than shallower ones (Figure 4). The LCMS data for the two Dagorlad seep cores varied less, in both feature counts (on-seep mat:  $777.3 \pm 51.8$ ; 5 m off-seep:  $762.5 \pm 65.6$ ) and abundances (on-seep mat:  $6.24E5 \pm 9.02E4$ ; 5 m off-seep  $5.85E5 \pm 1.15E5$ ). This is further reflected in the greater spread in core section LCMS data for the two Emyu Muil cores compared to the Dagorlad cores (Figure 5A). Additionally, the LCMS data formed less distinct clusters (Figure 5A) compared to the corresponding microbial ASV data (Figure 5B) of the sediment cores, reflecting the shared chemistry of metabolisms between different microbial taxa. In both the ASV and LCMS data plots, the OOI slope base core samples clustered separately from the on-seep and 5 m off-seep samples (Figures 5A, B).

The 5 m off-seep cores possessed greater ASV richness than their corresponding on-seep microbial mat cores. The Emyu Muil sediment core from 5 m off-seep had greater microbial ASV richness (6,948 ASVs) than the on-seep microbial mat core from the same site (4,427 ASVs). The Dagorlad seep cores had less ASV richness than the corresponding cores at the Emyu Muil seep but again, greater richness was seen in the 5 m off-seep core (5,802 ASVs) than the on-seep mat (1,338 ASVs). Sediment samples were generally clustered by core using microbial ASV data (Figure 5B). However, the samples from the younger Dagorlad seep cores clustered more tightly than those from the older Emyu Muil seep, where deeper depths were

less clustered than shallower depths (Figure 5B). The NMDS plot of core samples based on the actinobacterial ASVs (Figure 5C) displayed the least separation between the cores, likely due to the low overall counts of Actinobacteria. Nevertheless, it is noteworthy that actinobacterial ASV richness appeared to be less diverse for the on-seep microbial mat cores (EM:28, DS:12) than for the 5 m off-seep cores (EM:78, DS:35).

Clustering by similarity using the SIMPROF routine is a method of exploratory data analysis that detects patterns using null hypothesis testing. In summary, a Bray-Curtis similarity matrix for the data is compared, using the pi statistic, to a mean permuted similarity profile generated from iterative random shuffling of the raw data. This SIMPROF method was performed on microbial ASV abundance data, the actinobacterial ASV subset, and LCMS peak area (abundance) data for the sediment samples. Sediment core schematics, representing the spatial distribution of SIMPROF groupings for the three datasets by depth of core section sample, are presented in Figure 6, together with a template core on which section depths are labeled (Figure 6A). In general, for Emyu Muil and Dagorlad seep sites, changes in metabolites with depth, indicated by bold line boundaries between different colored sections, are consistent with changes in microorganisms with depth (Figures 6C–F). In contrast, the metabolite and microbial datasets for the non-seep OOI control site do not correlate well in terms of variation with depth, and there are overall fewer changes/boundaries (Figure 6B). In numerous instances for the two seep sites, as indicated by circled letters, SIMPROF groupings match between the LCMS and total microbial ASV datasets. These

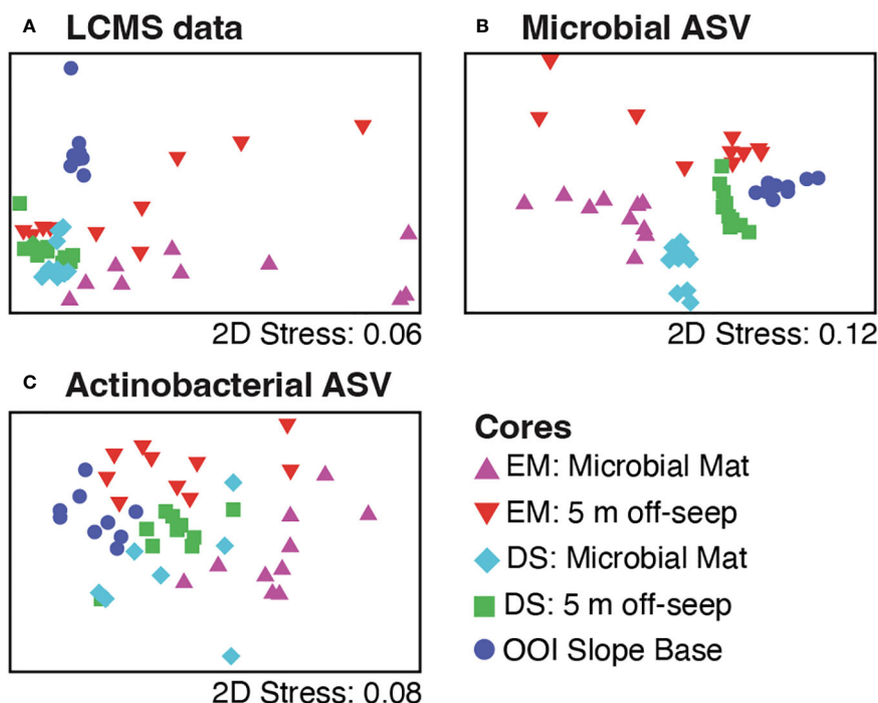


FIGURE 5

Non-metric multidimensional scaling (nMDS) plots of the data for 10 core sections from five different cores, including on-seep mat (Eymn Muil, EM, purple; Dagorlad Seep, DS, cyan), 5 m off-seep (EM, red; DS, green), and non-seep OOI control (dark blue) using (A) LCMS data, (B) all microbial (16S rRNA) ASV data, and (C) actinobacterial ASV data only.

correlations imply the presence of distinct microbial populations with correspondingly distinct chemistry in these core sections. There is less correlation in the change with depth of LCMS versus actinobacterial ASV data alone, which may again reflect the relatively low overall counts of actinobacterial ASVs. Somewhat correlated changes in depth between actinobacteria and LCMS datasets are apparent for the 5 m off-seep sites, for example, between 4–7 cm core section depth in the Eymn Muil samples and variously throughout the Dagorlad samples.

Significant positive and negative microbe–metabolite Spearman’s correlations occurred between microbial ASVs assigned at the family level and LCMS features annotated as putative peptides in either ontology (Supplementary Figure S1). The Log(X+1) normalized abundances of each of these features, separated by sample core depth for each site, is presented in Figure 7. Many individual LCMS features in this peptide dataset were simultaneously categorized in other NPClassifier pathways by CANOPUS, as anticipated for metabolites substituted with various functional groups, or with low probability peptide annotations. The 407 mass ion features annotated as “peptide” ranged from  $m/z$  361.2448 to 1239.9522. While the overall proportion of peptides did not change much between cores (Figure 4), at the feature level there are differences in the pattern of these metabolites present in the different cores. Peptidic metabolites significantly correlated to microbes were predominant in the upper two-thirds of the older Eymn Muil on-seep microbial mat core with relatively few correlated metabolites distributed throughout all core sections

(Figure 7). By comparison, the Eymn Muil 5 m off-seep core possessed a greater proportion of significantly correlated peptidic metabolites distributed from the surface to deeper in the core. The Dagorlad on-seep microbial mat and 5 m off-seep core profiles were comparable to each other and contained a majority of significantly correlated peptidic metabolites throughout the core, as was also found in the non-seep OOI core (Figure 7). The corresponding abundance of significantly correlated peptide metabolites plotted on the right-hand y axis indicates a decrease in peptide abundance with core section depth for the two Eymn Muil carbonate seep cores, but not for the soft sediment Dagorlad seep cores. This is in agreement with the trend across all metabolites (Figure 2). Notably, positive correlations were observed between actinobacterial families and putative cyclic peptides (Table 1). The complete dataset of correlations between features annotated as peptidic and microbial families can be found in the Supplementary Material (Supplementary Table S3, Supplementary Figure S1).

## 4 Discussion

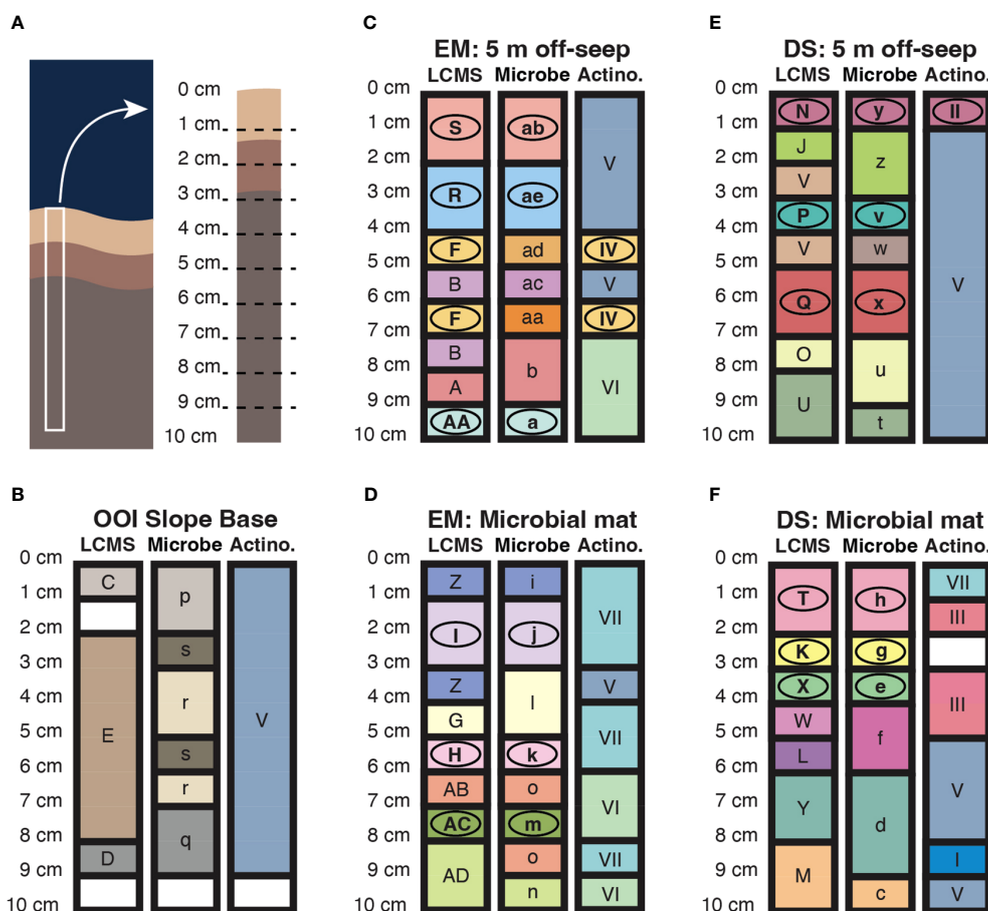
Our coupled metabolomics and bacterial community survey revealed distinct chemistry at methane seep sites compared to non-seep sites at the level of microbial family and metabolite structure class. The microbial family level was chosen for our analyses to facilitate data handling and visualization, and for consistency with our previous report (Cummings et al., 2023). Furthermore,

variations in specific molecular analogues within a series of biosynthetically related molecules (molecular family) may occur at the microbial genus, species or strain level, however, presence or absence of biosynthetic genes encoding a biologically important molecular family is most often noted at the microbial genus or higher taxonomic level (Donia et al., 2014). Chemical diversity changed across sites and core depth in relation to the changes in microbial diversity (Figures 5, 6). While the proportions of structure superclasses did not vary markedly across sites (Figures 2, 3), individual features and subclasses of compounds varied considerably between samples (e.g., for peptides, Figure 7).

We previously reported relative abundance of the 13 most abundant microbial families in the on-seep microbial mat and 5 m off-seep cores from Dagorlad and Emyu Muil seeps (Cummings et al., 2023). Notably, the microbial mat core from older Emyu Muil comprises a relatively large portion of anaerobic methanotrophic archaea (ANME), specifically ANME-1, not represented in the younger Dagorlad microbial mat. The Dagorlad microbial mat

instead comprises a relatively high abundance of sulfur-reducing bacteria (SRB) in the families Desulfobulbaceae and Desulfobacteraceae, with ANME-2 present. The partitioning of ANME-1 and ANME-2 in this way has been reported to be driven by methane supply (Niu et al., 2023) and suggests that the older Emyu Muil seep may experience higher methane seepage than Dagorlad seep. However, in the absence of flux data, there are a variety of possible interpretations, such as is proposed in Cummings et al., 2023. Further, nitrogen-fixing ANME-2 and SRB have been deemed responsible for the majority of new nitrogen production previously (Dekas et al., 2014). Thus other limiting nutrients or processes, besides methane seepage, may impact the community compositions observed here.

The relatively high abundance of lipids and lipid-like molecules at both methane seep sites versus the non-seep OOI control site (Figures 2, 3) may be attributed to methane seep-specialized bacteria, which are known to produce large quantities of methanotroph-specific monounsaturated fatty acids (Guan et al.,



**FIGURE 6** Core schematics representing the spatial distribution of similarity groupings according to SIMPROF analysis of LCMS features (LCMS), microbial 16S ASVs (Microbe), and actinobacterial 16S ASVs (Actino). Conserved color and label of core section within LCMS, Microbe or Actino datasets indicates that those sections belong to the same SIMPROF grouping. Circled letters indicate groupings that match across at least two datasets. (A) Diagrammatic representation of a sediment core *in situ*, in benthic substrate, alongside a representative core with depth graduations corresponding to sectioning of the core during sample processing. (B–F) Data for the non-seep OOI Slope Base, and off-seep and on-seep microbial mat for the younger Dagorlad (DS) and older Emyu Muil (EM) seeps, as labeled.

2022; Cordova-Gonzalez et al., 2020) and terpenoid compounds including carotenoids, steroids, polyprenols, and hopanoids (Salvador-Castell et al., 2019; Grivard et al., 2022). The unknown reason for the highest proportion of lipids and lipid-like molecules at the older carbonate substrate seep (Figure 2A) is presumably associated with the specific microbial community (and geochemistry) at the older carbonate substrate seep. These observations present opportunities for future hypothesis-driven research, potentially using targeted lipid analysis. These chemosynthetic habitats add to the chemical diversity of the ocean, and our exploratory work here demonstrates that even comparable habitats (soft sediment microbial mats, fueled by methane) have divergent metabolites (Figure 4), demonstrating that different chemosynthetic habitats add to the overall chemical diversity present in marine systems.

As very few MS ion features had spectral library or *in silico* predicted annotations in GNPS, we relied on CANOPUS to provide structural information for LCMS features that could be used to compare sediment cores from different sites. CANOPUS annotations are predictions that require further analysis and additional data for confirmation and comprehensive structure elucidation. CANOPUS predictions are also less accurate for molecules with masses greater than 850 Da, and SIRIUS does not currently support formula prediction or structural class prediction from doubly charged ion features. The semantic ontologies (NPClassifier and ClassyFire) used for computational assignment of structural class or 'pathway' may offer several alternatives since many chemical compounds comprise multiple substructures which fall into separate structural categories. Hence, assignment of a reasonably complex compound to only one structural class or another is subjective and requires manual assignment. Thus, metabolites are considered only at the level of biosynthetic class in this study.

Marine sediment samples posed challenges for LCMS analysis that may have resulted in an underestimation of the NPs present. Salt comprised a large portion of the mass-limited extracts obtained and potentially contributed to ion suppression, reducing the sensitivity of the mass detector (Metwally et al., 2015). To prevent large quantities of salt from entering the mass spectrometer, the LC column eluent was directed to waste for the first several minutes, meaning that more polar metabolites were likely lost to waste. The proportions of metabolites belonging to the various pathways are also likely skewed by electrospray ionization (ESI) technique and (positive or negative) mode used, as well as parameters set in the mass detection method, which affect metabolite ionizability. For example, carbohydrates may be underrepresented because many glycans do not give strong signals in positive mode due to lack of basic sites for protonation (Grabarics et al., 2022). Additionally, there is a large amount of elemental sulfur in deep-sea sediments around methane seeps, and organic polysulfide compounds may require chemical ionization (CI) or Fast Atom Bombardment (FAB) for mass detection, as is the case for lissoclinotoxins (Davis et al., 2003) and lissoclibadins (Nakazawa et al., 2007). As a result, while we have used a powerful technique coupled with multiple computational pathways, we are likely underestimating the chemical diversity present in methane seep habitats.

The correlations between LCMS features and particular bacterial families determined here will direct future investigation. The heatmap of peptidic feature correlations to microbial families (Supplementary Figure S1) emphasizes the chemical and biological complexity of methane seep sediment communities, illustrating that any one LCMS feature may have multiple positive and negative correlations with diverse microbial families and *vice versa*. The heatmap is organized by similarity of patterns in correlations, so that MS ion features with

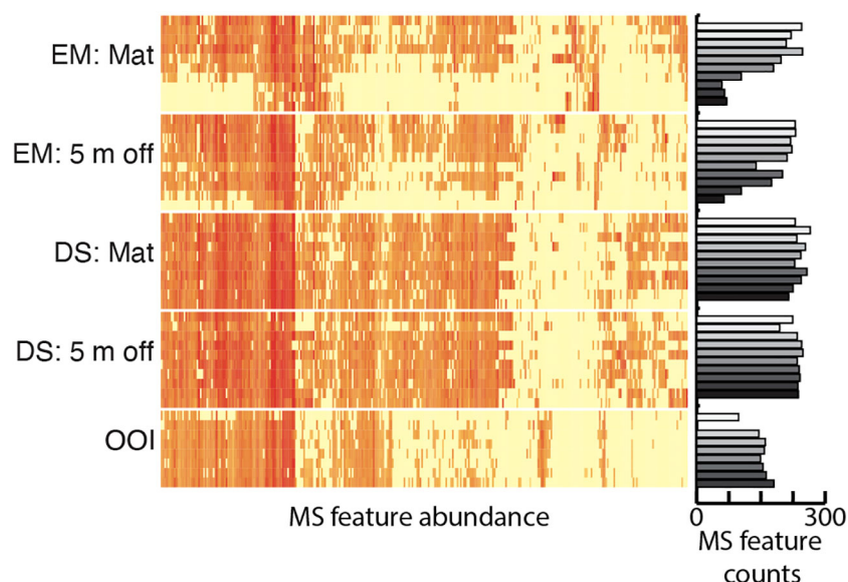


FIGURE 7

Heatmap of MS ion feature abundance representing distribution by core of putative peptidic metabolites with significant positive or negative correlations to families of microbial ASVs. Bar graph representing total numbers of features annotated as putative peptides by depth and core. For both heatmap and bar graphs, rows are stacked by depth within cores.

similar patterns of correlation across microbial families are clustered together. Thus, in the upper portion of the heatmap, there are obvious groupings of microbial families (in rows) that display consistent positive and negative correlations with groupings of metabolites, which may form the basis of hypotheses for future investigation.

While it is not possible to confidently assign producing organisms or functional roles to (structurally unassigned) metabolites based on Spearman's correlations (Carr et al., 2019), the co-occurrence of known biologically active classes of metabolites with bacterial families known to produce those metabolites demonstrates relevance and rationale for natural products biomedical and biotechnological research. Our data reveal that several bacterial families belonging to orders that have previously yielded structurally diverse bioactive NPs are positively correlated with LCMS features of the same structural classes as the known bioactive NPs. For example, positive correlations were seen between actinobacterial ASVs and several LCMS features annotated as cyclic peptides (Table 1). Actinobacteria are well-known to produce biologically active cyclic peptides (Zhao et al., 2018). Additionally, other bacterial families within Cellvibrionales (Han et al., 2013), Campylobacterales (Rischer et al., 2018) and Rhodobacterales (Íoca et al., 2021) also have significant positive correlations with putative peptidic features (correlation data in Supplementary Table S3), and members of these orders are known producers of bioactive nonribosomal peptides. Rhodobacteraceae are reported to produce peptide-derived alkaloids (Dickschat et al., 2005a; Dickschat et al., 2005b; Nicacio et al., 2017) and several LCMS features in our methane seep cores that were annotated as both alkaloid and peptide positively correlated with Rhodobacteraceae. The exploratory analysis undertaken here leverages significant bioinformatic and analytical advancement and provides a foundation for future work to realize the

NP potential as well as the functional role of specialized metabolites in marine systems. In order to bridge the gap between specialized metabolite discovery by LCMS feature annotation and identification of biologically active metabolites relevant to human health, the adoption of computational methods is essential. Emerging tools such as NPAnalyst (Lee et al., 2022) or Pearson correlation pipelines (Nothias et al., 2018) permit mapping of biological activity observed for complex mixtures to individual components therein. The ability to discern structural and substructural motifs known to be associated with specific bioactivities further enables more targeted testing of limited sample quantities. In addition, tools such as GNPS may allow identification of other sources of the same or related metabolites of interest, facilitating access to larger quantities for more extensive biological testing and structure elucidation. The deep sea, beyond depths accessible by SCUBA, is only beginning to be included in the search for new NPs, with 75% of the new metabolites discovered found to be biologically active, and almost half having some anti-cancer potential (Skropeta and Wei, 2014). The challenges of accessing, isolating, and characterizing new organic molecules requires ongoing technological innovations and interdisciplinary collaborative research, which is well justified given the tremendous contributions to date of small organic compounds in chemical and biological research, and to global health (Cheng et al., 2014; Newman and Cragg, 2020; Carroll et al., 2022). Here we present a snapshot of the potential for NP discovery in methane fueled habitats, where we found different metabolite profiles at two sites at the same water depth in relatively close proximity. Seep biodiversity is known to vary latitudinally across the Cascadia Margin (Seabrook et al., 2018) and there is high microbial heterogeneity at <1 m spatial scale (Boetius and Suess, 2004; Cummings et al., 2023). In addition, seeps change over time, as do the biological communities that inhabit them (Cordes

TABLE 1 Correlations between families within Actinobacteriota and mass features annotated as cyclic peptides by CANOPUS.

Microbial Families within Phylum Actinobacteriota			Corr. Coeff.	Adj. p value	Feature ID	m/z	NPC Class	NPC Class Prob.
Class	Order	Family						
Unassigned	Unassigned	Unassigned	0.4781	0.018	260	420.250666	Cyclic peptides	0.888
Acidimicrobiia	Microtrichales	Ilumatobacteraceae	0.5155	0.009	259	655.395939	Cyclic peptides	0.226
Coriobacteriia	CG2-30-50-142	CG2-30-50-142	0.5934	0.001	260	420.250666	Cyclic peptides	0.888
RBG-16-55-12	RBG-16-55-12	RBG-16-55-12	0.7564	4.40E-6	260	420.250666	Cyclic peptides	0.888
Thermoleophilia	Solirubrobacterales	67-14	0.5335	0.006	260	420.250666	Cyclic peptides	0.888
Thermoleophilia	Solirubrobacterales	67-14	0.4311	0.041	1109	693.551802	Cyclic peptides	0.990
WCHB1-81	WCHB1-81	WCHB1-81	0.5703	0.003	260	420.250666	Cyclic peptides	0.888
WCHB1-81	WCHB1-81	WCHB1-81	0.4553	0.027	1112	725.577524	Cyclic peptides	0.987

et al., 2005; Bowden et al., 2013; Thurber et al., 2020). If we presume our two seep sites are at the opposite ends of seep succession, temporal patterns also contribute to the diversity of metabolites present in these habitats. Continued and increased characterization of habitats at the interface of metabolome and microbiome science has astounding potential, especially considering the more than 3,500 seeps that are present but 99% unexplored on the Cascadia Margin (Merle et al., 2021), where this research was carried out. Characterizing the role of habitat heterogeneity on deep-sea ecosystem services, such as NP discovery, will allow informed management of deep-sea habitats with an eye towards future potential balancing immediate gains.

## Data availability statement

The original contributions presented in the study are publicly available. These data can be found here: For the LCMS data, MassIVE ID MSV000090543, <https://massive.ucsd.edu/ProteoSAFe/dataset.jsp?task=0de7900cc26b433bf18049d0bdf622b>; 16S sequence data, <https://www.ncbi.nlm.nih.gov/bioproject/?term=PRJNA718843>.

## Author contributions

AT, KM contributed to conceptualization of the study and funding acquisition; all authors contributed to the investigation; MR and MC led formal analysis and data visualization with contributions by all authors; MR, KM and AT contributed to writing – original draft. All authors contributed to the article and approved the submitted version.

## Funding

This research was funded by NOAA Ocean Exploration Grant NA19OAR0110301, and by the National Institutes of Health via

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## Acknowledgments

We thank Prof. Benjamin Philmus for assistance with LCMS data acquisition.

## Conflict of interest

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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## Supplementary material

The Supplementary Material for this article can be found online at: <https://www.frontiersin.org/articles/10.3389/fmars.2023.1197338/full#supplementary-material>

### SUPPLEMENTARY FIGURE S1

Heatmap of significant positive (red) and negative (blue) correlations between putative peptidic LCMS features and families of microbial ASVs.

### SUPPLEMENTARY TABLE S3

Dataset of correlations between all features annotated as peptidic and microbial families.

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