Bacterial and eukaryotic intact polar lipids point to in situ production as a key source of labile organic matter in hadal surface sediment of the Atacama Trench

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Abstract. Elevated organic matter (OM) concentrations are found in hadal surface sediments relative to the surrounding abyssal seabed. However, the origin of this biological material remains elusive. Here, we report on the composition and distribution of cellular membrane intact polar lipids (IPLs) extracted from surface sediments around the deepest points of the Atacama Trench and adjacent bathyal margin to assess and constrain the sources of labile OM in the hadal seabed. Multiscale bootstrap resampling of IPLs’ structural diversity and abundance indicates distinct lipid signatures in the sediments of the Atacama Trench that are more closely related to those found in bathyal sediments than to those previously reported for the upper ocean water column in the region. Whereas the overall number of unique IPL structures in hadal sediments contributes a small fraction of the total IPL pool, we also report a high contribution of phospholipids with mono- and di-unsaturated fatty acids that are not associated with photoautotrophic sources and that resemble traits of physiological adaptation to high pressure and low temperature. Our results indicate that IPLs in hadal sediments of the Atacama Trench predominantly derive from in situ production of microbial biomass, whereas the export of the most labile lipid component of the OM pool from the euphotic zone and the overlying oxygen minimum zone is negligible. While other OM sources such as the downslope and/or lateral transport of labile OM cannot be ruled out and remain to be studied, they are likely less important in view of the lability of ester-bond IPLs. Our results contribute to the understanding of the mechanisms that control the delivery of labile OM to this extreme deep-sea ecosystem. Furthermore, they provide insights into some potential physiological adaptation of the in situ microbial community to high pressure and low temperature through lipid remodeling.

1 Introduction

The deep ocean has been classically considered a vast “biological desert” (Danovaro et al., 2003) due to the attenuation of organic matter (OM) fluxes with increasing depth (Wakeham et al., 1984; Martin et al., 1987; Hedges et al., 2001; Rex et al., 2006). However, hadal trenches (~6000–11 000 m below sea level) contradict this paradigm (Danovaro et al., 2003; Glud et al., 2013; Leduc et al., 2016; Wenzhöfer et al., 2017), as they act as depocenters of OM (Jahnke and Jahnke, 2000) and hotspots for microbial activity (Glud et al., 2013; Wenzhöfer et al., 2016; Liu et
An alternative approach to study microbial processes and the contribution of autochthonous OM is the use of cell membrane intact polar lipids (IPLs), which although less specific than genomic markers, allow for more quantitative estimates of microbial biomass in nature (e.g., Lipp et al., 2008; Schubotz et al., 2009; Cantarero et al., 2020). IPLs are composed of a polar head group typically attached to a glycerol backbone from which aliphatic chains are attached via ester and/or ether bonds (Sturt et al., 2004). Their structural diversity is given by the modifications found in the different components of their chemical structure (e.g., polar head groups can be comprised of phosphorous, nitrogen, sulfur, sugars, and amino acids), whereas aliphatic chains (alkyl or isoprenoidal) can vary in their length (number of carbon atoms) and their degree of unsaturation, methylation, hydroxylation, and cyclization (Van Mooy and Fredricks, 2010; Brandsma et al., 2012; Schubotz et al., 2013). In bacteria and eukaryotes, alkyl chains are most commonly linked via an ester bond to the sn-glycerol-3-phosphate backbone (Koga and Morii, 2007), although some bacteria are known to produce di- and tetraether lipids (Weijers et al., 2007). The variability of membrane chemical structures underlies the adaptability of microbial lifestyles to changing environmental conditions such as nutrients, temperature, oxygen, pH, and pressure (DeLong and Yayanos, 1985; Somero, 1992; Van Mooy and Fredricks, 2010; Brandsma et al., 2012; Schubotz et al., 2013). In bacteria and eukaryotic and bacterial ester-bond IPLs are more labile than ether-bond counterparts (Logemann et al., 2011), they are suitable biomarkers to evaluate sources of labile OM in marine environments.

IPLs have been previously used as microbial markers in diverse marine settings, such as along strong redox gradients in the Black Sea (Schubotz et al., 2009) and the oxygen minimum zones (OMZs) of the eastern tropical Pacific (Schubotz et al., 2018; Cantarero et al., 2020) and the Arabian Sea (Pitcher et al., 2011), as well as in surface openocean waters of the eastern South Pacific (Van Mooy and Fredricks, 2010), the northwestern Atlantic (Popendorf et al., 2011b), and the Mediterranean Sea (Popendorf et al., 2011a), to name a few. Their utility as markers of microbial diversity and processes has also been tested in marine sediments (Liu et al., 2011, 2012; Sturt et al., 2004), such as along the Peru margin, equatorial Pacific, Hydrate Ridge, and Juan de Fuca Ridge (Lipp and Hinrichs, 2009a), and in subsurface sediment layers from the Peru margin (Biddle et al., 2006). However, to the best of our knowledge, no IPL studies have been reported for sediments of hadal trenches.

In this study, we investigate the chemical diversity and abundance of microbial IPLs as markers of one the most labile molecular fractions of OM in sediments of the deepest points of the Atacama Trench and compare them to IPL stocks in bathyal surface sediments (∼500–1200 m) and the overlying 700 m of the water column (Cantarero et al., 2020). More specifically, we evaluate possible IPL provenance (in
situ vs. allochthonous production) and the presence of unique IPL signatures of the in situ microbial community as well as evidence for molecular adaptations to the extreme conditions of the hadal region.

2 Material and methods

2.1 Study areas and sampling

The Atacama Trench is located in the eastern tropical South Pacific (ETSP) along the Peru–Chile margin, and it underlies the eutrophic and highly productive Humboldt Current System (Angel, 1982; Ahumada, 1989), which includes the fourth largest (by volume) oxygen minimum zone (OMZ) in the world (Schneider et al., 2006). In this area, while there is minimal river runoff (Houston, 2006), winds can transfer dust from the adjacent continental desert (Angel, 1982). With an extension of \( \sim 5900 \) km, the Atacama Trench is the world’s largest trench (Sabbatini et al., 2002), whereas it is geographically isolated from other trenches in the Pacific Ocean.

In this study, we investigated the diversity and abundance of bacterial and eukaryotic IPLs in a total of nine hadal surface (0–1 cm) and subsurface (1–2 and 2–3 cm) sediments (three sites between 7734 and 8063 m water depth) collected during the HADES-SO261 cruise (March to April 2018) aboard the RV Sonne (Wenzhöfer, 2019) and seven bathyal surface sediments (seven sites; 529–1200 m water depth) collected during the ChiMeBo-SO211 cruise (2–29 November 2010) aboard the RV Sonne (Matys et al., 2017) (Table 1; Fig. 1). We compare our results against IPL results from the overlying water column (0–700 m) recently reported in Cantarero et al. (2020).

Sediment samples were collected using a multi-corer (MUC) equipped with twelve 60 cm long acrylic tubes (6–10 cm diameter for bathyal sediments and 9.5 cm diameter for hadal sediments). During the HADES expedition, an autonomous lander equipped with a Sea-Bird SBE-19 plus CTD and two Niskin bottles (30 L) was used to obtain hydrographic data down to \( \sim 7850 \) m. Hadal sediments from the HADES-SO261 cruise were stored at 4°C until they were extruded and subsampled aboard at 1 cm resolution and then kept frozen at \(-20^\circ\)C until their processing in the laboratory. Further information about sample collection of bathyal and hadal sediments during the ChiMeBo-SO211 and HADES-SO261 cruises can be found in Matys et al. (2017) and Wenzhöfer et al. (2019), respectively.

We compare our IPL results from surface sediment in the hadal and bathyal regions against samples from the overlying water column (0–700 m) recently reported in Cantarero et al. (2020). This includes size-fractionated suspended OM (0.3–2.7 and 2.7–53 µm) at two stations and from six water depths that are representative of the dominant biogeochemical zonation associated with the OMZ of this
region: chlorophyll maximum (∼10 m), upper chemocline (∼25 m), lower chemocline (∼45 m), upper OMZ (∼60 m), core OMZ (∼250 m), and mesopelagic zone (∼750 m) (see Table 1 and Cantarero et al., 2020, for further details).

2.2 Analytical methods

2.2.1 Lipid extraction

All samples were processed, extracted, and analyzed in the Organic Geochemistry Laboratory at the University of Colorado Boulder. Sediment samples were freeze-dried before extraction. Approximately 1–2 g of dry sediment was placed in a combusted glass centrifuge tube and extracted using a modified version (Wörmer et al., 2013) of the Bligh and Dyer extraction method (Bligh and Dyer, 1959) as detailed in Cantarero et al. (2020). Briefly, before extraction, we added 1 µg of C16 PAF (C_{26}H_{54}NO_{7}P) to each sample as an internal standard. Samples were sequentially extracted using dichloromethane/MeOH/phosphate buffer (1:2:0.8 v:v:v), dichloromethane/MeOH/trichloroacetic buffer (1:2:0.8 v:v:v; 2×), and dichloromethane/MeOH (1:5 v:v; 1×). After each addition, samples were vortexed for 30 s, sonicated for 10 min, and then centrifuged for 5 min at 2000 rpm. Each extraction was then transferred to a separatory funnel where a total lipid extract (TLE) was combined and then concentrated under a gentle N₂ stream. Before analysis, the TLEs were resuspended in dichloromethane/methanol (9:1 v:v) and filtered through a 0.45 µm polytetrafluoroethylene (PTFE) syringe filter. The processing and extraction of bathyal sediments from the ChiMeBo-SO211 cruise and water-column samples from the LowpHOX-2 cruise has been reported by Matys et al. (2017) and Cantarero et al. (2020), respectively. TLEs were transferred into 2 mL vials with 200 µL inserts and dissolved in 100 µL of dichloromethane/MeOH (9 : 1, v : v).

2.2.2 IPL analysis

IPL were analyzed according to Wörmer et al. (2013) and as described in Cantarero et al. (2020) using a Thermo Scientific UltiMate 3000 high-performance liquid chromatograph (HPLC) coupled to a Q Exactive Focus Orbitrap quadrupole high-resolution mass spectrometer (HPLC-HRMS) via electrospray ionization (ESI). The HPLC program comprised a flow rate of 0.4 mL min⁻¹ using a mixture of two mobile phases: mixture A consisted of acetonitrile/dichloromethane (75 : 25, v : v) with 0.01 % formic acid and 0.01 % NH₄OH; mixture B consisted of methanol/water (50 : 50, v : v) with 0.4 % formic acid and 0.4 % NH₄OH. We used a linear gradient as follows: 1 % mixture B (0–2.5 min), 5 % (4 min), 25 % mixture B (22.5 min), 40 % mixture B (26.5–27.5 min), and the HPLC column was kept at 40 °C. Samples were injected (10 µL) and dissolved in dichloromethane/methanol (9 : 1, v : v). IPLs were separated using a Waters Acquity BEH Amide column (2.1 × 150 mm; 1.7 µm particle size) that enables class-specific separation based on their hydrophilic head group (Wörmer et al., 2013).

ESI settings comprised sheath gas (N₂) pressure 35 (arbitrary units), auxiliary gas (N₂) pressure 13 (arbitrary units), spray voltage 3.5 kV (positive ion ESI), capillary temperature 265 °C, and S-lens RF level 55 (arbitrary units). The instrument was calibrated for mass resolution and accuracy using the Thermo Scientific Pierce LTQ Velos ESI Positive Ion Calibration Solution (containing a mixture of caffeine,
MRFA, Ultramark 1621, and N-butylamine in an acetonitrile/methanol/acetic acid solution).

IPLs were identified on positive ionization mode, on both full scan and data-dependent MS², based on their molecular weights as either protonated (M + H)⁺ or ammonium (M + NH₄)⁺ adducts compounds, fragmentation patterns, and retention times, and they were compared against relevant literature (Sturt et al., 2004; Schubotz et al., 2009; Wakeham et al., 2012) and the internal database of the Organic Geochemistry Laboratory at the University of Colorado Boulder.

The peak areas of individual IPLs were integrated using the Thermo Fisher Scientific TraceFinder software using extracted ion chromatograms of their characteristic molecular ions. IPL abundances were determined with a combination of an internal standard (C₁₆:₁ PAF, Avanti Polar Lipids) and an external calibration to a linear regression between peak areas and known concentrations of an IPL cocktail comprised of 17 different IPL classes across a five-point dilution series (0.001–2.5 ng µL) (see Cantarero et al., 2020). Deuterated standards (Avanti Polar Lipids: d7-PC, d7-PG, d7-PE, and DGTS-d9) were used to correct for potential matrix effects on ionization efficiency. Despite the limited number of available deuterated standards, on average, we observed that the matrix effect accounts for a loss of ∼ 7 ± 0.6 % in ionization efficiency. Therefore, it is reasonable to assume a similar loss for other IPL classes, although this remains to be tested in future studies. We highlight the importance of using as many IPL classes as possible to account for both differences in ionization efficiency and matrix effect when performing IPL quantification in environmental samples. The relative response factors followed the order MGDG > DGTS > DGTA > PDME > PME > PG > PC > PE > SQDG > DGCC > DGDG. Lipid classes were grouped into phospholipids (PG, phosphatidylglycerol; PE, phosphatidylethanolamine; PC, phosphatidylcholine; and PME/PDME, phosphatidyl(dim)ethanoylglycerol), glycolipids (MGDG, monoglycosyldiacylglycerol; DGDG, diglycosyldiacylglycerol; and SQDG, sulfoquinovosyl-diacylglycerol), betaine lipids (DGTA, diacylglyceryl hydroxymethyl-trimethyl-β-alanine; DGTS, diacylglyceryl trimethylhomoserine; and DGCC, diacylglycerylcarnboxy-N-hydroxymethyl-choline), and other lipids (glycosidic ceramides, Gly-Cer; PI, phosphatidylinositol; and OL, ornithine lipids). In addition, we use DAG to designate a diacylglycerol and AEG to designate an acyletherglycerol, and we describe short and long chains to refer to combined alkyl chains of C₂₈–₃₆ and C₃₆–₄₄, respectively (Rézanka and Sigler, 2009; Schubotz et al., 2009; Brandsma., et al., 2012).

2.3 Statistical analyses

We used the Bray–Curtis similarity coefficient (Mirzaei et al., 2008) to produce hierarchical clustering of the abundance of classes and molecules of IPLs; two types of p values were available: approximately unbiased (AU) p value and bootstrap probability (BP) value with the number of bootstrap replications of 10 000 (Suzuki and Shimodaira, 2006). We performed non-metric multidimensional scaling (NMDS) (Warton et al., 2012) to examine the dissimilarity between the IPLs in each sample. The calculated distances to group centroids were based on the Bray–Curtis dissimilarity from the IPL abundance matrix, and the significance of the associations was determined by 999 random permutations. Significance tests of the multivariate dissimilarity between groups were made using analysis of similarity (ANOSIM), where complete separation and no separation among groups is suggested by R = 1 and R = 0, respectively (Clarke and Gorley, 2015). Statistical differences in the numbers of carbon atoms and double bonds were identified by ANOVA and Tukey’s HSD (honestly significant difference) post hoc test. We used similarity of percentage (SIMPER) analysis to identify the percentage contributions of IPLs which accounted for > 90 % of the similarity within each cluster. The multivariate statistical analyses as well as other statistical analyses were calculated using the vegan package (Oksanen et al., 2013) of open-source software R version 3.6.2 within the ggplots package (Warnes et al., 2015).

3 Results

3.1 Hydrographic conditions

A physical–chemical characterization of the water column during the ChiMeBo-SO211, LowpHOX-2, and HADES-SO261 cruises has been reported in Matys et al. (2017), Cantarero et al. (2020) and Vargas et al. (2021), respectively. Briefly, the potential-temperature–salinity–dissolved oxygen (θ-s-Ο₂) diagrams revealed an oxygenated and well-mixed water mass occupying the deeper parts of the Atacama Trench (Fig. S1). However, the upper 1000 m shows variability in temperature (12–23°C), salinity (34.4–34.8 psu), and oxygen (0.5–267 µM). More stable physical–chemical conditions are apparent in the mesopelagic and bathypelagic zone of the Atacama Trench between 1000 and 4000 m (temperature ~ 2.3°C, salinity ~ 34.6 psu, oxygen ~ 120.6 µM). Below 4000 m, average conditions were characterized by a potential temperature ~ 1.8°C, salinity ~ 34.7 psu, and oxygen ~ 143 µM (Fig. S1).

3.2 IPLs in surface sediments of the Atacama trench

3.2.1 Distribution of IPL classes by polar head groups

The 16 sediment samples from bathyal and hadal regions statistically grouped into four clusters based on their dominant polar head group classes (Fig. 2, chemical structures in Fig. S2). Clusters 1 and 2 had approximately unbiased (AU) p values of 91 % and 88 %, respectively. Cluster 3 had the
highest AU $p$ value of $\geq 97\%$, whereas cluster 4 had the lowest AU $p$ value of $61\%$. The cluster analysis revealed a degree of spatial heterogeneity between bathyal and hadal depths and between the top three centimeters of hadal sediments, which results in the lack of a clear separation between hadal and bathyal environments. In addition, the 0–1 cm hadal sediments at the A4 station were un-clustered, consistent with a distinct distribution pattern of IPL classes. Cluster 1, composed of only hadal samples from three different stations and depths, included phospholipids as the most abundant IPL class (Fig. 2). Clusters 2, 3, and 4, composed of mixed bathyal and hadal samples, were mostly differentiated by changes in the relative abundances of non-phosphorous IPLs including betaine classes. The un-clustered sample was characterized by the lowest relative abundance of phospholipids and the highest relative abundance of betaine lipids (especially DGCC).

### 3.2.2 Distribution of individual IPLs

An overview of the most important IPLs contributing to dissimilarity between samples was obtained through a SIMPER analysis based on the Bray–Curtis coefficient within each cluster (Fig. 3). Samples in cluster 1 were on average $59.5\%$ similar, with 14 individual IPLs contributing $50.6\%$ of the total similarity. This cluster exhibited a high contribution of PE-DAG (32 : 1, 33 : 1, and 34 : 2), PG-DAG (36 : 2), and DGCC (26 : 0, 27 : 0, and 28 : 0) molecules (Table 2). Additionally, this cluster exhibited a large diversity of PC molecules, although with a low relative abundance (Fig. 3). Samples in cluster 2, on the other hand, which includes mainly bathyal stations, were on average $58.8\%$ similar and exhibited a high contribution of PC-DAG (35 : 0, 32 : 1, 36 : 2, 33 : 1, and 35 : 1) (Table 2). While this cluster shows a wide range of molecules, including PG, PE, and MGDG, their relative contributions are low (Fig. 3). Samples in cluster 3 were on average $57.3\%$ similar and included three bathyal and one hadal station. This cluster exhibited a high contribution of DGCC (42 : 6) and PC-DAG (35 : 0, 33 : 2, 30 : 1, and 29 : 2) molecules (Table 2). Samples in cluster 4 were on average $63.6\%$ similar and exhibited a high contribution of PC-DAG (30 : 2, 33 : 2), DGCC (42 : 6), MGDG (28 : 0), and PE-DAG (33 : 2 and 31 : 2) molecules (Table 2). The un-cluster sample (hadal sediment of 0–1 cm at A4 station) is mainly composed by the DGCC 42 : 6 (Fig. 3). In general, phospholipids showed a wide distribution and were found across all sediment samples. The total dissimilarity between cluster 1 and 2 was $59.17\%$, with PC-DAG-35 : 0, PE-DAG-32 : 1, PI-AR, PG-DAG-36 : 2, DGCC 27 : 0, PC-DAG-36 : 2, PC-DAG-34 : 1, PC-DAG-32 : 1, DGCC 26 : 0, and PC-DAG-35 : 1 contributing $32.4\%$ of it (Table 2). The total dissimilarity between cluster 1 and 4 was $62.5\%$, with DGCC 42 : 6, PC-DAG-30 : 2, PE 32 : 1, PC-DAG-35 : 0, PG-DAG-36 : 2, PC-DAG-33 : 2, and DGCC 27 : 0 contributing $37.62\%$ of it (Table 2).

### 3.3 Distribution of alkyl chains based on length and degree of unsaturation

The difference in the total number of acyl carbon atoms in both alkyl chains, rather than in individual fatty acids, and in the number of acyl double bonds within each cluster is shown in Fig. 4. Statistical differences of IPLs classes within each cluster were obtained through a Tukey HSD post hoc test at a significant level of $p<0.05$ (Fig. 4a, b). The average number of carbon atoms in the diglyceride moieties of IPLs in the cluster 1 presented that DGCC, MGDG, others, PC, and PG were all distinct from one another ($n = 283$; $P < 0.05$; Fig. 4a). PG and others were characterized by relatively long alkyl chains (35–36 C atoms, respectively) and DGCC for shorter alkyl chains (32 C atoms). In general, cluster 1 exhibited a wide range of chain lengths among DAGs (28–36 C atoms). Cluster 2 showed a narrower range than cluster 1 (30–36 C atoms). This cluster also displayed no statistical difference ($p > 0.05$) among IPL classes (Fig. 4a), following pairwise comparisons with Tukey’s HSD post hoc test, despite the wide range of DGCC structures. For cluster 3, while it exhibited low variability in betaine lipids, it also revealed the highest number of carbon atoms in DGCCs (42). On the contrary, cluster 4 presented high viability in DGCCs, which did not exceed 42 carbon atoms. Within the phospholipid class, PG showed the highest number of carbon atoms in all clusters; the mean we observed was 34 carbon atoms and a range of 32–37 (Fig. 4a). The un-cluster sample (hadal sediment of 0–1 cm at A4 station) was characterized by relatively longer alkyl chains (up to 42 C atoms) than cluster 1 (Fig. 4a).

Overall, the degree of unsaturation (i.e., number of double bonds) within clusters was variable (Fig. 4b). Cluster 1 predominantly consisted of fully saturated and mono-unsaturated IPLs, except for PG that showed 2 double bonds on average. In cluster 2, the fatty acids of DGCCs were distinctly variable, although they exhibited 2 unsaturations on average. A similar pattern was observed in DGDGs with an average of 2.5 unsaturations (Fig. 4b). DGTs, MGDG, PC, and SQDG showed zero to 1 unsaturation, whereas DGTA, PE, and PG exhibited between 1 and 2.5 unsaturations. Cluster 3 showed more than 5 unsaturations on average for DGCC, unlike other IPL classes that did not exceed 2 unsaturations. In cluster 4, PG and DGCC presented $\sim 3$ and $\sim 5$ unsaturations on average. Also, on average, DGDG and SQDG exhibited 2 unsaturations, MGDG and others were mono-unsaturated, and DGTS was saturated (Fig. 4b). Additionally, the ratio of total unsaturated fatty acids to total saturated fatty acids in IPLs increased from (on average) $\sim 0.9$ in
Table 2. Similarity percentage (SIMPER) analysis. The average abundance and contribution of IPLs that explain the main differences among the sediment samples is based on the hierarchical clusters shown in Fig. 2.

<table>
<thead>
<tr>
<th>Group cluster 1</th>
<th>Cluster 1: average similarity = 59.53</th>
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<td>IPLs</td>
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<td>DGCC-26 : 0</td>
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<td>DGCC-27 : 0</td>
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<td>PE-DAG-32 : 2</td>
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<td>DGCC-28 : 0</td>
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all water-column samples (2–76 bar) to \(\sim 2.7\) in the bathyal (54–113 bar) and hadal sediments (777–810 bar) (Fig. 5).

### 3.4 Unique IPLs in hadal sediments of the Atacama Trench

Water-column particles and bathyal–hadal sediments shared 242 (96.1 \%) IPL structures (Fig. 6a), while hadal sediments and water-column particles shared 14 (0.02 \%), and hadal and bathyal sediments shared 55 (3.6 \%). Of all the analyzed IPLs reported in this study, eight of them were unique to the Atacama Trench sediments and are not present in shallower sediments or the overlying water column. They include five glycolipids (SQDG-42: 11, SQDG-23: 0, DGDG-35: 1, DGDG-35: 2 and DGDG-37: 1), two phosphatidylinositolts (PI-diOH-Ext-AR and PI-OH-AR), and one ornithine lipid (OL-37: 6). While unique to hadal sediments, their total concentration was low (\(\sim 53.32\) ng per gram of sediment), and they contributed \(\sim 0.00012\) \% of the total IPL pool (Fig. 6a). We then performed a cluster analysis to compare IPLs in deep-sea surface sediments against IPLs reported in the overlying water column (Cantarero et al., 2020; Fig. 6b). Cluster 1 comprised samples from the core OMZ in the free-living fraction (AU \(p\) value of 100 \%). Cluster 2 comprised samples from both the upper and lower oxyclines (\(\sim 14–60\) m) as well as from the chlorophyll maximum (AU

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**Table 2. Continued.**

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Figure 2. Cumulative bar charts of the fractional abundance of IPL classes in each surface sediment sample from the bathyal and hadal regions (left panel). Samples were grouped according to arithmetic mean (UPGMA) hierarchical clustering based on Euclidean distances. The \( p \) values are shown at branches, with AU in red and BP in green (right panel). Clusters 3 with an AU \( \geq 95\% \) confidence are indicated by the red rectangles (left) and are considered to be strongly supported by the data.

\( p \) value of 99\%). Cluster 3 comprised bathyal and hadal samples (AU \( p \) value of 99\%). Cluster 4 mostly comprised the deepest water-column sample (mesopelagic region at 750 m) and hadal samples (AU \( p \) value of 98\%; Fig. 6b). We also compared IPLs in hadal and bathyal sediments against the pool of IPLs reported as diagnostic of the planktonic community inhabiting the chlorophyll maximum in the upper water column (Cantarero et al., 2020) and thus assess their export and stability through their transit to the deep sea. Notably, these IPLs from this region of the water column only represent \( \sim \) 0.001\%–0.005\% and 0.002\%–0.03\% of the total IPL pool in hadal and bathyal sediments, respectively (Fig. S3).

We found a high degree of heterogeneity in total IPL concentrations among sites and different sediment levels (0–1, 1–2, 2–3 cm) in the Atacama Trench, which were an order of magnitude higher than bathyal sediments (see Fig. S4a, b). Hadal sediments at station A10 (7734 m) showed a large range of phospholipid concentrations (\( \sim 47–2698 \) ng per gram of sediment) (Fig. S4b). Although the highest total IPL abundances were observed at hadal station A10 (Fig. S4b), the greatest diversity in IPL composition was observed in the 0–1 cm of the hadal station A4, previously referred to as unclustered (see Fig. 2). The most abundant IPL class in hadal sediments was phospholipids, PCs (\( \sim 41–2698 \) ng per gram of sediment), PEs (\( \sim 26–1813 \) ng per gram of sediment), and PGs (5–937 ng per gram of sediment). The concentration of IPLs normalized by total organic carbon (TOC) (ng IPL per gram of TOC) showed maximum values in the hadal station A10 (\( \sim 497 \) µg IPL per gram of TOC), followed by lower values in the hadal stations A5 and A4 of \( \sim 291 \) and \( \sim 75 \) µg IPL per gram of TOC, respectively (Fig. S5).

4 Discussion

4.1 Potential sources of phospholipids

PG (phosphatidylglycerol)

Phospholipids are common constituents of cellular membranes in most microorganisms (Ratledge and Wilkinson, 1988). Since PGs play an essential role in photosynthesis (Wada and Murata, 2007), they have therefore been mainly

identified in algal and bacterial photoautotrophs (Dowhan, 1997; Sato et al., 2000; Gombos et al., 2002). However, their biological origin is highly diverse and also includes heterotrophic bacteria (Oliver and Colwell, 1973; Van Mooy et al., 2009; Popendorf et al., 2011b; Carini et al., 2015; Sebastián et al., 2016), methylo trophs (Batrakov and Nikitin, 1996), methanotrophic bacteria (Makula, 1978), Pelagibacter ubique (Van Mooy et al., 2009), and barophilic bacteria (e.g., DB21MT-2 and DB21MT-5) isolated from sediments from the Mariana Trench (Fang et al., 2000).

The hierarchical cluster analysis on variations in the relative abundance of PGs suggests that several compounds maintained a similar proportion in bathyal and hadal sediments, which differs from the water column (Fig. S6). Most PGs in the bathyal and hadal sediments have long acyl carbon chains (C_{34}–C_{41}), and they show odd- and even-numbered polyunsaturated fatty acids (Figs. 2, S6). The average chain lengths of even-numbered n-C_{18}, n-C_{20}, and n-C_{22} fatty acids, mostly in PCs and PGs, are indicative of algal inputs (Kaneda, 1991; Thompson, 1996; Bergé and Barnathan, 2005; Brandsma et al., 2012). However, since these PGs were not dominant in the water column, a source from deeper environments is likely. Specifically, PG-DAG-36 : 2, PG-DAG-35 : 2, PG-DAG-36 : 5, PG-DAG-37 : 2, and PG-DAG-41 : 4 are the dominant constituents of this IPL class in hadal–bathyal sediments (Figs. 7, S6). PG-DAG-36 : 2 has been described in surface waters of the North Sea and also detected in picoeukaryotes (Brandsma et al., 2012) and in heterotrophic bacteria in surface waters of the open South Pacific Ocean (Van Mooy and Fredricks, 2010). However, these PGs are not dominant in the water column near the Atacama Trench (Cantarero et al., 2020). On the other hand, PG-DAG-35 : 2, PG-DAG-36 : 5, PG-DAG-37 : 2, and PG-DAG-41 : 4 are not commonly reported in water-column studies. Thus, it is possible that PGs present in the Atacama Trench sediments derive from in situ microbial production, although downslope and lateral transport of labile OM cannot be ruled out. PG-DAG-36 : 2 (Fig. 3) is the PG contributing most to the dissimilarity within the cluster containing only hadal sediments (cluster 1 in Fig. 2). Thus, this lipid appears to be more representative of in situ microbial production in this environment.

**PE (phosphatidylethanolamine)**

PE and its methylated derivatives (PME, PDME) have been predominantly reported in membranes of diverse bacterial sources, including heterotrophic bacteria (Van Mooy and Fredricks, 2010; Schubotz et al., 2018), nitrifying/denitrifying bacteria (Goldfïne and Hagen, 1968), sulfate-reducing bacteria (Rütters et al., 2001; Sturt et al., 2004), sulfur-oxidizing bacteria (Barridge and Shively, 1968; Imhoff, 1995; Wakeham et al., 2012), methanotrophic bacteria (Makula, 1978; Sturt et al., 2004), and barophilic bacteria (Fang et al., 2000).

PEs showed a similar distribution in bathyal and hadal sediments (Fig. S7), where they are dominated by long-chain (C_{36}–_{44}) polyunsaturated fatty acids, contrary to the shorter chains (C_{28}–_{36}) of saturated and monounsaturated fatty acids present in the water column. PE-DAG-32 : 1, PE-DAG-32 : 2, and PE-DAG-33 : 1 are the dominant PE compounds of bathyal and hadal sediments (Fig. 7). These IPLs have been previously reported in heterotrophic bacteria (Van Mooy and Fredricks, 2010; Brandsma et al., 2012). On the
Figure 4. Total number of acyl carbon atoms (a) and acyl double bonds (b) in IPL classes across the distinct clusters shown in Fig. 2. The letters “a” and “b” indicate the presence of statistically distinct groups ($p<0.05$) from both ANOVA and post hoc Tukey HSD tests, respectively.

other hand, fatty acids in PEs including monounsaturated and polyunsaturated (e.g., C$_{20.5}$ and C$_{22.6}$) have been reported in barophilic bacteria isolated from sediments from the Mariana Trench (Fang et al., 2000). Thus, although we cannot confidently rule out other sources, it is possible that PEs present in the AT sediments predominantly derive from in situ production by barophilic heterotrophic bacteria. PE-DAG-32 : 1, PE-DAG-32 : 2, and PE-DAG-33 : 1 (Fig. 3) are the PEs that contributed most to the dissimilarity within the cluster containing only hadal sediment samples (cluster 1 in Fig. 2). Thus, this cluster appears to be representative of in situ microbial production in this environment.

PC (phosphatidylcholine)

PCs were amongst the most diverse (43 structures: Fig. S8) and abundant phospholipid class in hadal sediments (Fig. S4). PC is the major membrane-forming phospholipid in eukaryotes (Lechevalier, 1988; Sohlenkamp et al., 2003; Van Mooy et al., 2006; Van Mooy and Fredricks, 2010). Additionally, PC has been reported to be a major DAG in zooplankton, from protozoa to copepods and krill (Patton et al., 1972; Mayzaud et al., 1999; Lund and Chu, 2002). However, genomic data indicate that more than 10% of all bacteria possess the genetic machinery for PC biosynthesis (Sohlenkamp et al., 2003). PC has also been reported in nitrifying bacteria (Lam et al., 2007), phototrophic bacteria (Koblížek et al., 2006; Van Mooy et al., 2006), and barophilic bacteria (Fang et al., 2000). In surface sediments of the Black Sea (2000 m), PCs were related to algal material rapidly exported from surface waters (Schubotz et al., 2009).

Hadal and bathyal sediments, in addition to two OMZ core stations, were clustered in the PC class (AU $p$ value of 97%; Fig. S8). This cluster showed PCs with long (C$_{33}$–38) and polyunsaturated fatty acids (up to 10 unsaturations). The dominant constituents were PC-DAG-35 : 0, PC-DAG-30 : 2, PC-DAG-30 : 1, PC-DAG-33 : 2, PC-DAG-35 : 1, PC-DAG-29 : 2, PC-DAG-32 : 1, and PC-DAG-36 : 2 (Figs. 7, S8). PC-
DAG-36 : 2 and PC-DAG-30 : 1 have been associated with phytoplankton detritus (Schubotz et al., 2009) and bacteria (Brandsma et al., 2012), whereas PC-DAG-32 : 1 has been associated with picoeukaryotes (Brandsma et al., 2012).

Since the most abundant PCs in cluster 1 have not been reported as dominant structures in any specific environment before, they are possibly produced in situ, although downslope and/or lateral transport cannot be ruled out. Among bacteria, those membranes reported to contain PC belong to the alpha and gamma subgroups of the Proteobacteria (Sohlenkamp et al., 2003). Given that these bacterial groups are abundant in trench samples from Puerto Rico (Eloe et al., 2011), the Mariana Trench (Nunoura et al., 2015), and recently in the Atacama Trench (Schauberger et al., 2021), it is possible that PCs present in high abundance in the Atacama Trench are consistent with high abundance of Proteobacteria in these regions. Given their general known association and abundance in Atacama Trench sediments (Fig. S4), they likely derive primarily from bacterial but also possibly from fungal or metazoan sources that have not yet been studied and to a lesser extent from phytoplankton. Indeed, fungal strains isolated from the water column and sediment in the ESTP off Chile reported high levels of polyunsaturated fatty acids and PCs (Gutiérrez et al., 2020), whereas a high fungal diversity associated with denitrification potential was reported in the Yap Trench (Gao et al., 2020). The latter suggests that eukaryotic PCs in hadal sediments could be much more diverse in origin than previously thought.

**PME/PDME (phosphatidyl(di)methylethanolamine)**

PME/PDMEs have been observed in association with methanotrophic bacteria (Makula, 1978; Goldfine, 1984; Fang et al., 2000); sulfide-oxidizing bacteria (Barridge and Shively, 1968); sulfate-reducing bacteria, mainly Desulfobulbus spp. (Rossel et al., 2011), Proteobacteria (Oliver and Colwell, 1973; Goldfine, 1984); and barophilic bacteria from the Mariana Trench (Fang et al., 2000). Additionally, the occurrence of PME-DEG at some hadal stations suggests the presence of sulfate-reducing bacteria (Rütters et al., 2001; Sturt et al., 2004).

PME/PDMEs exhibited their lowest abundance (~10 ng per gram of sediment) in sediment samples compared to other phospholipids (Fig. S4b). In the bathyal and hadal sediments they were clustered (AU p value of 97 %) and dominated by PDME-DAG-33 : 1, PME-DAG-37 : 2, PME-DAG-34 : 2, PME-DAG-31 : 1, and PME-DEG-33 : 0 (Fig. S9a). PME-DEG-33 : 0 has been shown to correlate with high NO3− in the overlying water column of this area (Cantarero et al., 2020), which could suggest a potential association with denitrification processes. These structures have also been reported in the deep chemocline of the Cariaco Basin (Wakeham et al., 2012), suggesting a potential chemoautotrophic and/or heterotrophic source. The distribution of these com-

Figure 6. Comparison of IPLs in bathyal and hadal sediments (this study) and the overlying water column (Cantarero et al., 2020). (a) Venn diagram showing the number and percentage of unique and shared IPL molecules between these three environments. (b) Cumulative bar charts of IPL fractional abundances in each sample. Samples were grouped according to arithmetic mean (UPGMA) hierarchical clustering based on Euclidean distances. The cluster analysis on the right-hand side shows approximately unbiased (AU) and bootstrap probability (BP) in red and green numbers, respectively, whereas \( p \) values are shown at branching points. Clusters with AU \( \geq 95 \% \) confidence are highlighted in red on the left-hand side.

The hierarchical cluster suggests that several MGDG compounds maintained a similar proportion in bathyal (AU \( p \) value of 90 \%) and hadal (AU \( p \) value of 98 \%) sediments (Fig. S10). The most abundant MGDGs in the bathyal and hadal sediments were MGDG-28 : 0, MGDG-30 : 0, and MGDG-31 : 0 (Figs. S9a and S16; Cantarero et al., 2020). Thus, and similar to other lipid classes, they most likely derive from in situ production in hadal sediments rather than from the water column, although other sources such as downslope and/or lateral transport cannot be ruled out. No particular PME/PDMEs were found to contribute to the dissimilarity between the cluster containing only hadal sediment samples (cluster 1 in Fig. 2) and other sediment samples.

4.2 Potential sources of glycolipids

MGDG (monoglycosyldiacylglycerol)

Due to their dominant occurrence in chloroplast thylakoid membranes (Murata and Siegenthaler, 1998) and particularly in cyanobacteria (Heinz, 1977; Harwood, 1998; Wada and Murata, 2007; Van Mooy and Fredricks, 2010), but also in heterotrophic bacteria (Popendorf et al., 2011b), MGDGs are probably the most abundant IPLs on earth (Gounaris and Barber, 1983).

The hierarchical cluster suggests that several MGDG compounds maintained a similar proportion in bathyal (AU \( p \) value of 90 \%) and hadal (AU \( p \) value of 98 \%) sediments (Fig. S10). The most abundant MGDGs in the bathyal and hadal sediments were MGDG-28 : 0, MGDG-30 : 1, MGDG-32 : 1, MGDG-32 : 0, and MGDG-37 : 3. MGDG-28 : 0 and MGDG-30 : 1 are ubiquitous along the oxycline of the overlying OMZ (Fig. 7; Cantarero et al., 2020). In addition, MGDG-32 : 1 has been previously reported in waters of the eastern South Pacific (Van Mooy and Fredricks, 2010). Thus, the occurrence of these MGDGs in sediment could indicate at least some export of labile OM from surface waters. On the other hand, MGDG-37 : 3 does not appear to be a dominant structure in any specific environment in the literature, which suggests a likely in situ production.
DGDG (diglycosyldiacylglycerol)

DGDGs are commonly found in membranes of eukaryotic algae and cyanobacteria (Wada and Murata, 1998; Sakurai et al., 2006; Kalisch et al., 2016). DGDGs clustered together in bathyal and hadal sediments (AU p value of 96 %), whereas their distribution differed from the water column (Fig. S11). The most abundant DGDGs in hadal and bathyal sediments of the Atacama Trench was DGDG-34 : 2 (Fig. 7), which has been previously reported in cyanobacterial strains isolated (da Costa et al., 2020) but has not been previously reported as abundant in the water column. In contrast, DGDG-30 : 0, which is widely distributed in the water column of this region (Cantarero et al., 2020), is consistently present in hadal and bathyal sediment samples although at very low abundances (Fig. 7). Thus, although DGDGs account for less than ∼ 5 % of the total IPL pool (Fig. 6b), except for station A10 (2–3 cm) where they reach ∼ 10 %, their presence in bathyal and hadal sediments is indicative of at least some export of labile OM from surface waters.

SQDG (sulfoquinovosyldiacylglycerol)

SQDGs are predominantly produced by photoautotrophs (Van Mooy et al., 2006; Popendorf et al., 2011b), including various groups of diatoms, brown and green algal chloroplast membranes (Harwood, 1998), and cyanobacteria (Siegenthaler, 1998; Wada and Murata, 1998). SQDGs have also been found in bacteria from the α- and γ-proteobacterial lineages (Benning, 1998). In the overlying water column of the Atacama Trench, Cantarero et al. (2020) suggested a higher contribution of SQDGs from cyanobacteria than algae. Also, SQDGs found in the deep Atlantic (down to ∼ 4000-5000 m) appear to indicate a source and export from surface waters (Gašparović et al., 2018).

SQDGs showed a consistent distribution in bathyal and hadal sediments, where they are dominated by long-chain (C36–44) fatty acids (Fig. S12). This is contrasting to their distribution in the overlying water column where they are dominated by shorter-chain (C28–36) saturated fatty acids (Cantarero et al., 2020). SQDG-30 : 0, SQDG-32 : 0, SQDG-30 : 2, and SQDG-38 : 4 were the dominant SQDG constituents of bathyal and hadal sediments (Fig. 7). SQDG-30 : 0 and SQDG-30 : 2 have been reported in bacteria in North Sea surface waters (Brandsma et al., 2012), in cyanobacteria of the eastern subtropical South Pacific (Van Mooy and Fredricks, 2010), and in plankton detritus from surface sediments of the Black Sea (Schubotz et al., 2009). Furthermore, SQDG-30 : 0 is abundant in surface waters of our study area, and SQDG-38:4 has been correlated with NO−3 (Cantarero et al., 2020). The observed differences in the distribution of SQDGs in deep sediments compared to the water column suggests an in situ production of previously poorly characterized compounds, in addition to at least some export from surface waters.

4.3 Potential biological sources of betaine lipids

DGTS (diacylglycerol trimethylhomoserine)

DGTSs have diverse biological origins, being found in a wide range of eukaryotes (Sato, 1992; Dembitsky, 1996; Kato et al., 1997; Van Mooy et al., 2009), photoheterotrophic bacteria (Benning et al., 1995; Geiger et al., 1999), photoautotrophic bacteria (Popendorf et al., 2011b) including cyanobacteria (Rezanka et al., 2003), and members of the α-Proteobacteria subdivision (López-Lara et al., 2003). Schubotz et al. (2018) showed DGTS with varying fatty acid compositions in the OMZ system of the eastern tropical North Pacific, especially in OMZ waters, indicating that these compounds can be biosynthesized by a wider range of source organisms than previously thought.

Consistent with other IPL classes, DGTSs of the bathyal and hadal samples were grouped in the same cluster (AU p value of 98 %) and differed from the water column (Fig. S13). However, several DGTSs are shared between surface waters (9–60 m) and deep sediments. Indeed, the most abundant DGTSs in bathyal and hadal sediments (DGTS-34 : 0, DGTS-32 : 1, DGTS-26 : 0, DGTS-34 : 1, DGTS-32 : 0, and DGTS-25 : 0; Figs. 7, S13) are also prominent in the chlorophyll maximum in the eastern subtropical South Pacific (Van Mooy and Fredricks, 2010; Cantarero et al., 2020). Therefore, their presence in hadal sediments suggest the export of some labile OM from the euphotic zone, although we cannot rule out other sources.

DGTA (diacylglycerol hydroxymethyl-trimethyl-β-alanine)

DGTA have been widely reported in eukaryotic phytoplankton (Araki et al., 1991; Dembitsky, 1996; Cañavate et al., 2017), mainly in diatoms (Volkman et al., 1989; Zhukova, 2005; Gómez-Consarnau et al., 2007), and are also especially abundant in cultures of prymnesiophytes and cryptophytes (Kato et al., 1997). DGTA have also been found in cyanobacteria (Brandsma et al., 2012) and heterotrophic bacteria (Popendorf et al., 2011a; Sebastián et al., 2016).

DGTA in bathyal and hadal sediments are mainly composed of longer (C28–C42) and polyunsaturated (1–12) fatty acids compared to those present in the shallowest region of the overlying water column, composed of shorter and saturated fatty acids (Fig. S14). In the overlying water column, these compounds are associated with relatively high chlorophyll and O2 concentrations (Cantarero et al., 2020), similar to North Sea surface waters (Brandsma et al., 2012). To the best of our knowledge, the dominant DGTA in hadal and bathyal sediments (Figs. 7, S14) have not been previously reported as dominant IPLs in other environments. Whereas no specific biological sources in hadal sediments are known, the structures containing between 30 and 38 carbon atoms might be characteristic of this type of environment.
DGCC (diacylglycerylcarboxy-N-hydroxymethyl-choline)

Our knowledge of DGCC sources is limited. They have been found in membranes of prymnesiophyte algae (Kato et al., 1994), mainly in *Pavlova lutheria* (Kato et al., 1994; Eichenberger and Gribi, 1997) and in *E. huxleyi* (Volkman et al., 1989; Pond and Harris, 1996; Van Mooy and Fredricks, 2010). Additionally, they have also been reported in the diatom *Thalassiosira pseudonana* (Van Mooy et al., 2009).

The most abundant IPL from the entire data set of bathyal and hadal sediments is DGCC-42 : 6 (Figs. 7, S15). This is the compound with the largest number of C atoms (42) and unsaturation (6) in all IPLs detected in this study. DGCCs with long-chain, polyunsaturated fatty acids (i.e., C_{38:6}, C_{40:10}, C_{42:11}, and C_{44:12}) have been previously reported in phytoplankton (Hunter, 2015; Van Mooy and Fredricks, 2010). However, the most abundant DGCCs in hadal sediments have, to the best of our knowledge, not been previously reported, which highlights their potential as biomarkers of deep-sea sediments. However, three hadal stations clustered in a separate group (see Fig. S15) were dominated by DGCC-27 : 0 and did not contain DGCC-42 : 6, indicating that this IPL probably derives from allochthonous sources.

**4.4 Potential biological sources of other lipids**

Glycosidic ceramides (Gly-Cer) have been reported in eukaryotic algae such as prymnesiophyte (Vardi et al., 2009) and have also been shown to be abundant in water columns of OMZ systems (Schubotz et al., 2009, 2018; Cantarero et al., 2020). In general, the overlying water column shows Gly-Cer with a ceramide chain and polyunsaturated fatty acids with C_{21–38}. However, these structures are scarce in the bathyal and hadal sediments (see Fig. S9b), which could reflect a deficient export from surface waters due to intense remineralization. On the other hand, ornithine lipids (OL), phosphatidylinositol (PI), PC-AEGs, and other unidentified phospholipids were also present in deep sediments (Fig. S9b). Some PIs and OLs have been reported in sulfate-reducing bacteria (Sturt et al., 2004; Bühring et al., 2014), whereas PC-AEGs have been reported in bacteria inhabiting water columns with reduced oxygen concentration (Schubotz et al., 2018). Thus, the high relative abundance of PC-AEG-34 : 3 in hadal and bathyal sediments (Figs. S9b and S16) could be indicative of anaerobic microbial processes. PC-AEG-34 : 3 contributed the most to the dissimilarity between the cluster containing only hadal sediment samples (cluster 1 in Figs. 2 and 3), thus suggesting an in situ microbial production, although we cannot confidentially rule out other sources.

**4.5 Allochthonous versus autochthonous IPLs in the Atacama Trench**

Given their rapid degradation after cell death (White et al., 1979; Harvey et al., 1986; Logemann et al., 2011; Schouten et al., 2010), IPLs are typically considered markers of living or recently dead cells (White et al., 1979; Harvey et al., 1986; Petersen et al., 1991; Lipp et al., 2008). The distribution of IPLs in bathyal and hadal sediments exhibits a high degree of similitude, as demonstrated by the hierarchical analysis (cluster 1 in Fig. 8a), the NMDS (Fig. 8b), and the SIMPER analysis (cluster 1 in Table S1). The deep-sea surface sediments showed weak clustering with the IPLs reported in the overlying water column by Cantarero et al. (2020) (Fig. 9a). Additionally, water-column samples exhibit a larger degree of separation than sediments (ANOSIM, *R* = 0.78; *P* < 0.01; Fig. 8b) and are broadly clustered by geochemical environments (Cantarero et al., 2020). The low abundance of IPLs characterized of organisms inhabiting the chlorophyll maximum in deep-sea sediments of the Atacama Trench (< 0.005 % of the total IPL pool; Fig. S3) suggests minimal export of labile organic compounds from the upper ocean. This result implies rapid IPL degradation during sinking in the water column, which is consistent with experimental degradation rates (Westrich and Berner, 1984; Logemann et al., 2011) and first-order POM sinking rates. Indeed, by using the experimentally calculated kinetic degradation rate constants (k') of ester-bound IPLs by Logemann et al. (2011) and the sinking rate of particles from surface waters to 4000 m (20–100 m d^{-1}; Billett et al., 1983; Danovaro et al., 2014), we calculated that ~86%–98% (k' = 0.033 and k' = 0.011) of IPLs from surface waters should degrade by the time particles reach depths of ~8000 m. These results are also in accord with studies indicating elevated benthic oxygen consumption rates resulting from intense microbial respiration of sinking OM reaching the sediment (Glud et al., 2013; Wenzhöfer et al., 2016). Thus, the pool of IPLs in hadal sediments appears to predominantly represent in situ microbial production, whereas the deep-sea microbial community in both bathyal and hadal sediments is similar despite their bathymetric zonation (~1000–8000 m).

Alternatively, we cannot rule out the possibility of new IPL production, particularly from heterotrophic and chemoheterotrophic bacteria in micro niches of sinking particles reaching the deep sea and/or downslope and lateral sediment transport.

Marine trenches receive organic carbon from a variety of sources and transport mechanisms. These include canyons and river systems that channel OM from land to coastal regions, aeolian transport, surface water productivity, and in situ production, to name a few (Wenzhöfer et al., 2016; Tarn et al., 2016; Luo et al., 2017; Xu et al., 2018; Guan et al., 2019; Xu et al., 2021). Carbon flux events can increase the delivery of particulate carbon from surface waters to the seafloor (Poff et al., 2021), whereas river discharge and ae-
oligian transport can result in enhanced terrestrial carbon (Xu et al., 2021). Mass wasting events are also known to create dynamic depositional conditions and strong spatial heterogeneity in OM distribution in marine trenches (Schauberger et al., 2021; Xu et al., 2021). While marine organic carbon appears to dominate sediments in the Japan (Schwestermann et al., 2021), Massau (Xu et al., 2020a), and New Britain (Xu et al., 2020b) trenches, the Atacama and Kermadec trenches, on the other hand, have been reported to be dominated by terrigenous OM. Since our study only focuses on the most labile component of the total lipid pool, it predominantly traces labile and indigenous OM and not recalcitrant fractions of the lipid pool. The latter warrants further investigation.

In regions like the Japan Trench, downslope sediment transport has been linked to earthquake-driven remobilization (Bao et al., 2018; Schwestermann et al., 2021). Whereas we lack sedimentological/geochemical data to discriminate whether the top 3 cm of our hadal stations represent debris flows, turbidite, or mass wasting events, ongoing work in the Atacama Trench indicates heterogenic sediment deposition along the hadal zone (Matthias Zabel, personal communication, 2021). Thus, the role of downslope transport as a mechanism to explain the high statistical similarity between bathyal and hadal sediments remains to be tested.

### 4.6 Characteristic IPLs of hadal and bathyal sediments

The IPLs that contribute most to the dissimilarity between the hierarchical cluster containing samples from the hadal and bathyal sediments (cluster 1 of Fig. 8) and the water column (cluster 2, 3, 4, and 5 of Fig. 8) are represented in Fig. 9. The most characteristic IPLs of hadal and bathyal sediments are DGCC-42:6, DGCC-27:0, DGCC-26:0, PC-DAG-35:0, PC-DAG-30:1, PC-DAG-30:2, PC-DAG-33:2, PC-DAG-32:1, PE-DAG-32:1, PE-DAG-32:2, PE-DAG-33:1, PG-DAG-36:2, and DGDG-34:2, which we propose as potential markers for these environments. Even though DGCCs have been mainly related to algae membranes (Kato et al., 1994; Van Mooy et al., 2009), they are minor components of the water column in this area, suggesting the occurrence of an alternative source. In addition to DGCCs, the two other betaine lipids, DGTA and DGTS, exhibited five IPLs that were almost exclusively present in sediment samples (DGTA-34:1, DGTA-32:1, DGTA-34:2, DGTS-34:0, and DGTS-32:1; see Fig. 11). We note that almost all the PC phospholipids in our study have not, to the best of our knowledge, been previously reported in the literature, which reinforces their use as markers of sedimentary in situ bathyal and hadal production.
The presence of a few MGDGs and SQDGs in hadal and bathyal sediments (~7 % of the total IPL pool) indicates that at least some labile OM could derive from the shallow water column (see Sect. 4.2). However, the most abundant IPLs in our sediment samples, DGCC-42 : 6, PC-DAG-35 : 0, PE-DAG-32 : 1, and PG-DAG-36 : 2 (19.8 % of the total IPL pool; Fig. S16), are almost completely absent in the overlying water column (Fig. 9). This reinforces the idea that these IPLs most likely originate from in situ microbial production in sediments. The single most abundant IPL in sediments, DGCC-42 : 6, was not present in cluster 1, which only contains hadal sediments (Figs. 2 and 3). Instead, this compound is prominent in cluster 3, 4, and 5, containing both hadal and bathyal samples. Thus, DGCC-42 : 6 and PC-DAG-35 : 0, which has the lowest relative abundance in the cluster with only hadal sediments, could be indicators of downslope transport from bathyal to hadal regions.

We acknowledge that temporal variability in IPL production in the water column and sediment and the lack of data on the largely uncharacterized hadal endemic microbial community could complicate some of the phylogenetic and source associations of IPLs and warrant further investigation. Despite this, our study represents a step forward on the characterization of labile sources of OM sustaining hadal ecosystems.

4.7 Do IPLs reveal homeoviscous adaptation to the deep-sea environment?

Environmental factors such as pH, conductivity, temperature, and pressure impact the permeability and fluidity of cell membranes (Shaw, 1974; Macdonald, 1984; DeLong and Yayanos, 1985; Somero, 1992; Komatsu and Chong, 1998; Van Mooy et al., 2009; Carini et al., 2015; Sebastián et al., 2016; Siliakus et al., 2017; Boyer et al., 2020; Allen et al., 1999). Thus, organisms adapt to changes in environmental factors to maintain physiological homeostasis by altering their fatty acid composition (DeLong and Yayanos, 1985; Fang et al., 2000; Nichols et al., 2004; Siliakus et al., 2017). For instance, the combined physiological effects of high hydrostatic pressure and low temperature on prokaryotic membranes in laboratory cultures leads to the production of unsaturated lipids (DeLong and Yayanos 1985; Fang et al., 2000; Nichols et al., 2004; Siliakus et al., 2017). However, few
Figure 9. Relative abundance of individual IPLs that contribute most to the dissimilarity between clusters of Fig. 8 derived from the SIMPER analysis (Table S1). Circle size is proportional to the relative abundance of IPL compounds per sample. Samples are organized along the vertical axis and shown in colors that match the hierarchical cluster analysis in Fig. 8. The legend shows the scale for circumference size.

studies have been conducted using culture-independent tech-
niques in search for potential adaptation mechanisms in or-
ganisms inhabiting the deep ocean (i.e., Zhong et al., 2020).
We sought to understand whether the chemical composition
of core fatty acids within different IPL classes (i.e., carbon
length and unsaturation degree) reflects the combined ef-
facts of the low temperature and high pressure typical of
hadal settings. We show that PGs are abundant in hadal sedi-
ments of the Atacama Trench (Fig. S4). Bacterial strains iso-
lated from Mariana Trench sediments contain PG as the most
abundant class of phospholipids (Fang et al., 2000), which
these authors presumed could represent a physiological re-
response to high pressure and low temperature. This has been
confirmed by subsequent studies (Winter et al., 2009; Peri-
asamy et al., 2009; Jebbar et al., 2015, Allemann et al., 2021).
Cluster 1 in the boxplot analysis (Fig. 4) likely contains the
most characteristic IPL classes of the hadal zone. In general,
the phospholipids in this cluster exhibited fatty acid chains
that are monounsaturated and saturated compared to other
environments (Fig. 4a, b). Additionally, we observed an in-
crease in the ratio of total unsaturated to saturated fatty acids
in deep sediments compared to the water column (Fig. 5),
which could reflect physiological adaptations of their bio-
llogical producers. These results are in accord with studies
indicating biosynthesis and incorporation of polysaturated
fatty acids into phospholipid membranes of piezophilic bac-
teria (DeLong and Yanoas, 1985; Baird et al., 1985; Yano
et al., 1998; Winter, 2002; Mangelsdorf et al., 2005; Winter
and Jeworrek, 2009; Allemann et al., 2021). Thus, the chem-
ical characteristics (C length and degree of unsaturation) of
the most abundant IPLs in sediments of the Atacama Trench
suggest homeoviscous adaptation to this type of environment
by their source organisms, in addition to potentially indicat-
ing the occurrence of compounds that are unique to the en-
dogenous community.

5 Conclusions

Bacterial and eukaryotic IPLs in surface hadal sediments
from the deepest points of the Atacama Trench share char-
acteristics with those in bathyal sediments and differ from
those found in suspended particles from the upper 750 m of
the water column, including the oxygen minimum zone. This
indicates that (a) most IPLs abundant in the upper water col-
umn are almost entirely degraded during their descent to the
hadal seafloor and (b) IPLs found in hadal sediments are pre-
dominantly derived from in situ microbial communities.

The most dominant ester-bound IPL structures found in
bathyal and hadal sediments show a great variety of phospho-
lipids with varying degrees of unsaturation, most of them yet
to be described, that are likely derived from as of yet poorly
characterized bacterial and/or eukaryotes sources. Hadal sed-
iments also exhibit unique glycolipid structures, such as SQDG-42 : 11, SQDG-23 : 0, DGDG-35 : 1, DGDG-35 : 2, and DGDG-37 : 1, that to the best of our knowledge have not been reported in other environments. However, these lipids are present in low abundance and represent a small fraction (∼0.00012 %) of the total IPL pool. Furthermore, elevated ratios of unsaturated/saturated fatty acids in hadal sediments are likely indicative of homeoviscous adaptation to the high pressure and low temperatures characteristic of this extreme deep-sea environment.

An improved understanding of the phylogenetic, ecological, and metabolic association of IPLs present in the Atacama Trench could be achieved in future studies by the pairing of lipidomics with genomic techniques (e.g., microbial community composition, functional groups, lipid biosynthesis), in addition to a detailed sedimentological and biogeochemical characterization of sediments.

Data availability. Biomarkers metadata that generates and supports the findings of this study and R code are available in: https://github.com/EdgartFlores/IPLs-Atacama-Trench-, last access: 3 March 2022 (https://doi.org/10.5281/zenodo.6325647, Flores, 2022).

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