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# Isoprenoidal GDGTs and GDDs associated with anoxic lacustrine environments

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# ABSTRACT

We examined membrane-spanning archaeal lipids using ultra high pressure liquid chromatography-high resolution mass spectrometry (UHPLC-HRMS) in a suite of sediment samples from both cored sequences (Messel oil shale and Lake Chala) and surface sediments (Azorean lakes) encompassing ancient and modern (Eocene to Present) lacustrine environments. Additionally we compared the lacustrine data to those of marine (Mediterranean cored sequences, Arabian Sea surface sediments and Monterey outcrop sediments) and hypersaline sediments (Vena del Gesso marls) as well as marine suspended particulate matter (SPM) from the Black Sea. Regular isoprenoidal glycerol dialkyl glycerol tetraethers (GDGTs) and glycerol dialkyl diethers (GDDs) were the most abundant membrane-spanning lipids in all investigated settings (>90 % and 84 % respectively). Interestingly, GDGTs with a cyclohexyl ring (S-GDGTs) were also present in almost all investigated lake sediments, in relative abundances of ca. 2-7 % and, for the first time, also their S-GDD counterparts were detected (2-10 %). The producers of S-GDGTs are still unknown, however our results show that it is likely that bottom water anoxia (both seasonally induced or permanent) is the driving factor for the production of these lipids, whereas previous studies suggested euxinia was required for production. Unsaturated GDGTs (uns-GDGTs, ca. 2 %) were only detected in Lake Chala sediments and surface sediments from Azorean lakes, but without accompanying uns-GDDs. GMGTs, glycerol monoalkyl glycerol tetraethers, were present in Messel oil shale and marine samples, while GMDs were only found in Messel oil shale.

#### 1. Introduction

Archaea are widespread in both the marine water column and in underlying sediments (Brocks and Pearson, 2005; Lipp and Hinrichs, 2009; Spang et al., 2017; Summons et al., 2021) where they play important roles in global biogeochemical cycles such as the methane and nitrogen cycles (Boetius et al., 2000; Pancost et al., 2001; Könneke et al., 2005; Francis et al., 2007; Jarrell et al., 2011). They are also widespread in lacustrine environments, where they perform a similar function (Glissman et al., 2004; Bomberg et al., 2008; Auguet et al., 2010; Vuillemin et al., 2018). Archaeal membranes are characterized by isoprenoidal glycerol ether lipids, mainly *sn*-2,3-diphytanyl glycerol diethers with two  $C_{20}$  phytanyl chains (archaeol) or *sn*-2,3-dialkyl diglycerol tetraethers with two glycerol moieties connected by two  $C_{40}$  isoprenoid chains (glycerol dialkyl glycerol tetraethers [GDGTs], which can contain 0 to 8 cyclopentane moieties [i.e., GDGT-n, where n is the number of cyclopentane moieties]; cf. Fig. 1) (e.g., De Rosa and Gambacorta, 1988; Koga et al., 1993; Gambacorta et al., 1995; Sinninghe Damsté et al., 2002). Soon after their discovery, it was suggested that isoprenoidal ether lipids could provide an advantage in extreme

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environments (high temperature, high salinity, or extreme pH), on the basis that they are more chemically stable than the ester-linked membrane lipids present in Bacteria and Eukarya (Langworthy, 1977; Thompson et al., 1992; Kates, 1993). However, the discovery of wide-spread occurrence of GDGTs in non-extreme environments (Schouten et al., 2000a), in concert with the cosmopolitan detection of archaeal DNA (e.g., Fuhrman et al., 1992; De Long 1992), threw doubt on this hypothesis. Schouten et al. (2002) first suggested that the number of cyclopentane moieties in isoprenoidal GDGTs within marine sediments was related to sea surface temperature (SST), and developed the TEX<sub>86</sub> index (TetraEther indeX of 86 carbons) as a paleothermometer applicable to marine sediment sequences. Since then the TEX<sub>86</sub> index has been used to reconstruct SST over the past 190 million years of Earth history (Inglis et al., 2015; O'Brien et al., 2017).

Developments in liquid chromatography-mass spectrometry (LC-MS) methodology (e.g., Hopmans et al., 2000; Wörmer et al., 2013, 2015) have facilitated the subsequent discovery and tentative identification of a wide range of GDGTs other than the 'regular' GDGTs (i.e., GDGT 0-8). For example, Liu et al. (2012c) described a range of novel hydroxy isoprenoidal ether lipids in marine sediments, the OH-GDGTs, which were thought to mainly originate from planktonic archaea (Huguet et al., 2013; Elling et al., 2014). OH-GDGT based indices show potential as a tool to reconstruct past surface water temperature in polar regions (e.g., Yermak Plateau close to the Arctic Ocean: Kremer et al., 2018; Prydz Bay in Antarctica: Liu et al., 2020). Glycerol monoalkyl glycerol tetraethers (GMGTs; also referred to as H-shaped GDGTs) were first reported in the hyperthermophilic methanogen Methanothermus fervidus (Morii et al., 1998; Koga and Morii, 2005) and the thermophilic euryarchaeote Aciduliprofundum boonei (Schouten et al., 2008a). Naafs et al. (2018) showed that GMGTs may be used as paleotemperature indicators in tropical peats and immature coals (lignite) from mid-latitude regions. Unsaturated GDGTs (uns-GDGTs) have been reported in a suite of marine environments by Zhu et al. (2014) and, based on their abundances and distributions, the authors suggested that they constitute an important archaeal lipid group in euxinic basins (the Black Sea Basin) and an

investigated cold seep site (site off the coast of Pakistan, in which focused upward migration of methane stimulates high rates of anaerobic oxidation of methane). Uns-GDGTs were also reported in the lacustrine surface sediment of a meromictic marl lake (Fayetteville Green Lake, near Syracuse in New York: Liu et al. 2016a). Unsaturations have been frequently reported in diphytanyl glycerol diethers (DGDs, e.g., archaeol), where the proportion of uns-DGDs in cultured halophilic archaea increases with increasing ambient NaCl concentration (Maestrojuan et al., 1992; Hafenbradl et al., 1993; Dawson et al., 2012). However, reports of uns-GDGTs in marine and lacustrine sediments are relatively sparse. Biphytanes which contain a cyclohexane ring, instead of ones with the more commonly reported cyclopentane ring, were first described by Chappe et al. (1980) and Chappe (1982) after HIdegradation (refluxing with hydroiodic acid in order to release and analyse the ether-bound hydrocarbon skeletons of GDGTs) of Messel oil shale lacustrine sediments. The GDGTs containing these biphytane moieties were first reported by Liu et al. (2016a) in surface sediment from Fayetteville Green Lake (FGL), the Messel pit (horizon between 2.5 and 3.5 m), sediments from Salt Pond, Woods Hole salt marshes at Trunk River (41°32'4.8"N, 70°38'30.4"W) and Little Sippewissett marsh (41°34'33.5"N, 70°38'14.9"W). These were named S-GDGTs (where 'S' stands for both 'sulfidic' and 'six-membered ring') as they were hypothesized to be produced by chemoautotrophic archaea that live in euxinic conditions.

In the past decade, glycerol dialkyl diether (GDD; cf. Fig. 1), analogues to several GDGT classes have also been tentatively identified. Regular GDDs have been observed in marine subsurface sediments from various geological settings (e.g., the Mediterranean Sea, margins of Peru, Namibia, Cascadia etc. by Liu et al., 2012a) and Mediterranean sediments from the Messinian period (Knappy and Keely, 2012), OH-GDDs in marine (Liu et al., 2012b) and coastal wetlands (Lü et al., 2019), and GMDs, the diether analogues of GMGTs, in sediments deposited under hydrothermal conditions (surface sediments from the Red Sea; Bauersachs and Schwark, 2016). It has been postulated that GDDs are either degradation products of GDGTs or biosynthesized by



**Fig. 1.** Structures of membrane-spanning archaeal lipids: regular isoprenoid GDGTs (-0 to Cren); S-GDGTs; H-shaped (GMGTs & GMDs) and unsaturated (uns-GDGT-1 & uns-GDD-1) GDGTs and GDDs. Note: the position of bridge in H-shaped and double bond in unsaturated GDGTs/GDDs is speculative.

archaea, either as membrane components or as GDGT intermediates (Liu et al., 2012a; Yang et al., 2014; Meador et al., 2014; Coffinet et al., 2015; de Bar et al., 2019).

In this study we examined GDGT and GDD diversity in a range of recent and ancient lacustrine settings that are either seasonally or permanently anoxic as well as in various anoxic marine settings using ultra high pressure liquid chromatography-high resolution mass spectrometry (UHPLC-HRMS). We specifically focused our study on those GDGTs and GDDs whose occurrence has been linked to anoxia, to further constrain their use as potential proxies for anoxic environmental conditions.

# 2. Materials and methods

#### 2.1. Samples and study sites

We analysed sediment cores from two geographically distinct crater lakes characterized by permanent anoxia (Messel paleolake in Germany and Lake Chala in Kenya) as well as modern surface sediments from a variety of lakes (of volcanic origin and other natural depressions) from the Azores Archipelago. For comparison, sediments (both core and surface) and suspended particulate matter (SPM) from several marine environments characterized by anoxia or anoxic events were also studied (cf. Table S1 for the full list of samples).

#### 2.1.1. Lakes

**Messel oil shale.** Messel paleolake is an ancient maar crater lake that was formed by a phreatomagmatic eruption (Felder and Harms, 2004). The samples for this study (n = 18) were obtained from a core drilled in the Messel oil shale sequence (borehole number 5 – Darmstadt, Germany; Matthess, 1966; Rullkötter et al., 1988; Franzen and Schaal, 2000) that is Middle Eocene in age (47.8 Ma; Mertz and Renne, 2005; Lenz et al., 2011). It is noteworthy that Messel paleolake in the Middle Eocene was ~  $46^{\circ} - 47^{\circ}$ N, therefore experiencing a tropical climate (Grein et al., 2011). The total organic carbon content (TOC) of Messel oil shale is 27 % on average (Bauersachs et al., 2014).

Azorean lakes. Surface sediment samples (n = 18) from seven Azorean lakes and one coastal lagoon in the mid-Atlantic Azores Archipelago (Portugal) were analyzed. The Azores are particularly rich in lentic habitats, with 88 lakes (Porteiro, 2000), ranging between 0.4 m (Lagoa do Caldeirão Norte on São Miguel Island) and 115 m depth (Lagoa Negra on Flores Island), respectively. The lower water column of the deeper lakes is known to experience seasonal anoxia (i.e., depletion of oxygen in the bottom waters due to temperature gradient that stratifies the water column during summer months) (Gonçalves, 2008; Ribeiro et al., 2014; Raposeiro et al., 2018; Ritter et al., 2022). Surface sediments were collected during a sampling campaign in 2018 and stored at -40 °C at the NIOZ. TOC, as determined in this study, varied between 4 % (Lake Azul on São Miguel Island) and 21 % (Lake Lomba on Flores Island).

Lake Chala. Lake Chala (Kenya/Tanzania) is a permanentlystratified (meromictic) tropical crater lake (Verschuren et al., 2009; Buckles et al., 2014, 2016). In September 2016 the International Continental Scientific Drilling Program (ICDP) project DeepCHALLA (Verschuren et al., 2013) recovered a continuous sediment sequence spanning the last ca. 250 kyr (Martin-Jones et al., 2020; Maitituerdi et al., 2022) from below its profundal zone. For this study, 168 samples from the core were examined ( $\sim$ 20 – 63 m and 119 – 190 m) of which 14 % contain turbidites (sediments originating from a more peripheral location that end up at the drill site after bottom-slope failure; cf. Table S1 for full list of samples).

2.1.2. Marine and hyper saline sediment and suspended particular matter Monterey Formation. Marine sediments from the Miocene Monterey Formation (n = 10; 5.9 – 18.4 Ma, Schouten et al., 2000b, c; Isaacs et al., 2001; Isaacs and Rullkötter, 2001) were investigated. These sediments

are outcrop samples from the Naples Beach section, California (USA) obtained in 1989 and 1990 in the framework of the Cooperative Monterey Organic Geochemistry Study (CMOGS) (Isaacs, 1992). Based on the integrative CMOGS geological and palaeontological analysis, sediments from Naples Beach represent a unique occurrence (Isaacs and Rullkötter, 2001 and references therein). TOC varied between 1 and 17 % (Isaacs and Rullkötter, 2001).

*Sapropels.* Eastern Mediterranean sapropel sediments from the Pleistocene, ranging in age from 121.5 to 128 kyr BP were analysed (n = 12; S5; Rush et al., 2019; Bale et al., 2019a). These sediments were sampled from a piston core taken at a water depth of 1760 m in the eastern part of the basin (Levantine Basin) (core 64PE406-E1) on R/V *Pelagia* in 2016. TOC content is up to 12 %.

*Western Mediterranean.* Western Mediterranean sediment core sampled at Station 8 (core 64PE407-W8) (n = 3; Weiss et al., 2019) at 1963 m water depth. The core was retrieved during a cruise 64PE407 on R/V *Pelagia* in 2016.

**Arabian Sea.** Arabian Sea surface sediments from various water depths (900 – 3000 m) collected from the Murray Ridge in the Arabian Sea in January 2009 (n = 3; Lengger et al., 2012).

**Vena del Gesso.** Marls were analysed from the Vena del Gesso outcrop (Gessoso-solfifera formation) deposited during the Mediterranean Miocene salinity crisis (n = 12; Messinian – 7.2 Ma, Vai and Ricci Lucchi, 1977; Sinninghe Damsté et al., 1995). Vai and Ricci Lucchi (1977) identified 14 distinguishable evaporitic cycles (i.e., beds) that formed in the Vena del Gesso Basin; the samples investigated in this study belong to four different cycles, namely the most developed bed IV, beds V, VII and XI. The TOC ranges from 1.2 to 2.5 %.

**Black Sea water column.** SPM (n = 15) was analysed from the suboxic and euxinic redox zones of the Black Sea obtained during two cruises in 2013 (PHOXY cruise; 100 to 2000 m water depth) and 2017 (64PE418 cruise; 90 to 2000 m water depth) on board of R/V *Pelagia* (Bale et al., 2021; Ding et al., 2021).

#### 2.2. Sample extraction

The majority of the samples were extracted using a modified Bligh-Dyer (BD) method (Sturt et al., 2004; with additional modifications described in Bale et al., 2021). Samples extracted for this study include Messel oil shales, the Azorean lake surface sediments, Vena del Gesso marls, Monterey sediments, Western Mediterranean sediment core and Arabian Sea surface sediments. The Mediterranean sapropels and Black Sea SPM were extracted as described in previous studies (Rush et al., 2019; Bale et al., 2019a; Bale et al., 2021). A mixture of methanol: dichloromethane:phosphate buffer (MeOH:DCM:PB 2:1:0.8, v:v) was used for ultrasonic extraction (2  $\times$  10 min) and combined supernatants were phase-separated by adding DCM and PB to obtain a final solvent ratio of 1:1:0.9 (v:v). The organic phase containing the intact polar lipids (IPLs) was collected and the aqueous phase re-extracted three times with DCM. The whole extraction process was then repeated on the residue but with a mixture of MeOH:DCM:TCA (aqueous trichloroacetic acid solution; pH 3; 2:1:0.8, v:v). The organic extracts were combined and dried under a stream of N2 gas. Before analysis the extracts were redissolved in a mixture of MeOH:DCM (9:1, v:v) and filtered through 0.45 µm regenerated cellulose syringe filters (4 mm diameter; Grace Alltech).

The Lake Chala sediments have been extracted previously in the same manner as described for the Lake Chala sediments (core depths between 60 and 65 m) in Baxter et al. (2019). Briefly, extraction was done using ASE Dionex<sup>TM</sup> apparatus with a DCM:MeOH mixture 9:1 (v: v) at high temperature (100 °C) and pressure (7.6 × 10<sup>6</sup> Pa). The total lipid extracts (TLEs) were first dissolved in a small amount of DCM/ methanol (v:v 1:1) and run on a Na<sub>2</sub>SO<sub>4</sub> column to remove the excess water and dried under N<sub>2</sub> gas. TLEs were than separated into apolar, ketone and polar fractions using Al<sub>2</sub>O<sub>3</sub> column chromatography with eluents *n*-hexane:DCM (9:1, v:v), *n*-hexane:DCM (1:1, v:v) and DCM: MeOH (1:1, v:v), respectively. All Chala polar fractions were dried under

 $N_2$  gas and filtered using a PTFE 0.45  $\mu m$  filter. The polar fractions were analysed for the purpose of this study.

# 2.3. Hydrogenation

Selected acid-hydrolysed Bligh Dyer (BD) extracts of Messel oil shale sediments (from 72 and 76 m core depth; Table S1), as well as a polar fraction of one surface sediment from Lake Azul (SS4; Table S1) were hydrogenated to aid in the identification of unsaturated GDGTs. Hydrogenation was done in ethyl acetate (EtOAc) with 1 drop of concentrated acetic acid by bubbling  $H_2$  over PtO<sub>2</sub> as catalyst (Aldrich) for 1 h at room temperature, followed by overnight stirring (Schouten et al., 2008a). GDGTs and GDDs. Stepped normalized collision energy (NCE) of 15, 22.5 and 30 was used for fragmentation. Distributions of GDGTs and GDDs were determined by integration of peak area of the summed mass chromatograms of the protonated, ammoniated and sodiated molecules  $([M + H]^+, [M + NH_4]^+, [M + Na]^+)$  and their first isotopologue within 3 ppm relative mass tolerance.

#### 2.6. Indices and Statistics

The ring indices of GDGTs, GDDs, S-GDGTs and S-GDDs were calculated using a modified equation of He et al. (2012). The ratios and calculations that were carried out on the core lipid data are as follows: Ring index GDGTs =

$$(\% GDGT-1) + (\% \ S-GDGT-1) + 2*(\% GDGT-2) + 2*(\% S-GDGT-2) + 3*(\% GDGT-3) + 4*(\% GDGT-4) + 5*(\% Crenarchaeol) + 2*(\% S-GDGT-2) + 2*(\% S-GDGT-2) + 3*(\% GDGT-3) + 4*(\% GDGT-4) + 5*(\% Crenarchaeol) + 2*(\% S-GDGT-2) + 3*(\% GDGT-3) + 4*(\% GDGT-4) + 5*(\% Crenarchaeol) + 2*(\% S-GDGT-2) + 3*(\% GDGT-3) + 4*(\% GDGT-4) + 5*(\% Crenarchaeol) + 2*(\% S-GDGT-2) + 3*(\% GDGT-3) + 4*(\% GDGT-4) + 5*(\% Crenarchaeol) + 2*(\% S-GDGT-2) + 3*(\% S-GDGT-3) + 4*(\% GDGT-4) + 5*(\% Crenarchaeol) + 3*(\% S-GDGT-3) + 4*(\% S-GDGT-4) + 5*(\% Crenarchaeol) + 3*(\% S-GDGT-3) + 3*(\%$$

100

and Ring index GDDs =

 $\frac{(\% GDD - 1) + (\% S - GDD - 1) + 2*(\% GDD - 2) + 2*(\% S - GDD - 2) + 3*(\% GDD - 3) + 4*(\% GDD - 4) + 5*(\% Cren - GDD)}{100}$ 

# 2.4. Elemental analyses

TOC, total nitrogen (N<sub>tot</sub>) and total sulfur (S<sub>tot</sub>) contents were measured for the Azorean lake surface sediments (n = 18). The analysis was done on decalcified, powdered and freeze-dried samples (ca. 30 mg for lakes Azul and Verde, ~10 mg all others) using an Elementar Vario Isotope Cube directly linked to an Isotope Ratio Mass Spectrometer (IRMS) Elementar Isoprime vision. Samples for TOC analysis were pretreated with excess 2 M hydrochloric acid (HCl) to remove carbonates and were put on a shaker overnight. They were then neutralized with bidistilled water and freeze-dried. TOC was measured in duplicate with a sample reproducibility of < 0.5 %.

# 2.5. Ether lipid analysis

Analysis of both the BD extracts and polar fractions was carried out using UHPLC-HRMS according to Wörmer et al. (2013), with modifications described in Bale et al. (2021). Briefly, analysis was carried out on an Agilent 1290 Infinity I UHPLC, equipped with thermostatted autoinjector and column oven, coupled to a Q Exactive Orbitrap MS with a Ion Max source and heated electrospray ionization (HESI) probe (Thermo Fisher Scientific). Separation was achieved on an Acquity BEH C18 column (Waters, 2.1  $\times$  150 mm, 1.7 mm) maintained at 30  $^\circ\text{C}.$  The eluent composition was (A) methanol/water/formic acid/14.8 M NH3aq [85:15:0.12:0.04 (v:v)] and (B) isopropyl alcohol/methanol/formic acid/14.8 M NH<sub>3</sub>aq [50:50:0.12:0.04 (v:v)]. The elution program was: 95 % A for 3 min, followed by a linear gradient to 40 % A at 12 min and then to 0 % A at 50 min, this was maintained until 80 min. The flow rate was 0.2 mL min<sup>-1</sup>. Positive ion HESI settings were: capillary temperature, 300 °C; sheath gas (N<sub>2</sub>) pressure, 40 arbitrary units (AU); auxiliary gas (N<sub>2</sub>) pressure, 10 AU; spray voltage, 4.5 kV; probe heater temperature, 50 °C; S-lens 70 V. Lipids were detected using a mass range of m/z350 – 2000 and MS<sup>2</sup> spectra were obtained via data-dependent acquisition, where the top ten abundant ions per MS<sup>1</sup> scan were selected for fragmentation. Additionally, dynamic exclusion (6.0 s) was used as well as an inclusion list containing calculated m/z values for all known

Principal component analysis was performed using R (version R-4.0.4; R Core Team, 2021) using factoextra (version 1.0.7; Kassambara and Mundt, 2020) on calculated relative abundances of regular, H-shaped, *uns-* and S-GDGTs and GDDs.

#### 3. Results & discussion

# 3.1. Identification of GDGTs and GDDs

A broad range of regular isoprenoidal GDGTs and GDDs were detected in all settings (cf. Fig. 1 for structures and nomenclature). GDGTs and GDDs were tentatively identified based on their elution patterns under reversed phase chromatography (Liu et al., 2016a) and by comparison with published fragmentation spectra (Knappy et al. 2009, 2011; Knappy and Keely 2012; Liu et al., 2012b, c; Zhu et al., 2014; Bauersachs and Schwark, 2016). Fig. 2 shows the distribution of GDGTs and GDDs in a Lake Messel oil shale (79 m; Fig. 2a) and in Lake Azul surface sediment (SS4; Fig. 2b). In addition to the regular GDGTs 1-4 and crenarchaeol, several early and late eluting isomers of various GDGTs were observed (Fig. 2a, b). S-GDGT-1 (with one cyclohexyl moiety; MS<sup>2</sup> in Fig. S1b) and S-GDGT-2 (with one cyclopentyl and one cyclohexyl moiety) elute after GDGT-1 and -2 respectively (Fig. 2), as described by Liu et al. (2016a). The extracted ion chromatograms of m/z1300.307 and 1298.291 also contained a series of peaks eluting before GDGT-1 and GDGT-2. GMGT-0, -1 and -2 (Fig. 2a) were identified based on comparison of their fragmentation spectrum to published spectra (Schouten et al., 2008a; Knappy et al., 2009; Bauersachs and Schwark, 2016). Spectra of GMGTs are characterized by a series of losses of H<sub>2</sub>O (-18) from their protonated molecule and a lack of fragments relating to the loss of one of the isoprenoid chains. Liu et al. (2016a) identified early eluting peaks as unsaturated GDGTs (uns-GDGTs), with a reversed elution order compared to the regular GDGTs. As uns-GDGTs cannot be distinguished from GDGTs with ring structures in the isoprenoid chains based on mass spectrometry, selected samples were hydrogenated to confirm their identification (Fig. S2). Peaks that were no



Fig. 2. Partial summed Extracted Ion Chromatograms (EIC), showing elution patterns of GDDs and GDGTs under reverse phase (RP) conditions with accurate masses of protonated, ammoniated and sodiated adducts (and their first isotopologue) indicated, of Messel oil shale sediment (79 m) (A) and Lake Azul surface sediment (SS4) (B). Structures of detected ether lipids are shown in Fig. 1. All chromatograms were smoothed with a factor Gaussian 7. NL: corresponds to Normalization Level of peak intensity.

longer detected after hydrogenation were thus identified as *uns*-GDGTs (Fig. S2b). The early eluting isomers in Messel were identified as GMGTs, while in Azul they were identified as *uns*-GDGTs. The *uns*-GDGTs observed in Azul conform to the elution pattern designated as 0:1:0 and 0:2:0 by Liu et al. (2016a) (*uns*-GDGT-1 and *uns*-GDGT-2, respectively; Fig. 1 for structures and nomenclature). This assignment complies with x:y:z naming system where x represents the number of cyclopentyl rings, y the number of double bonds and z the number of cyclohexyl rings in GDGT structure, as described in Liu et al. (2016a).

In addition to the GDGTs described above, various GDDs were detected in all investigated sediments. The distribution of GDDs in a Messel oil shale and Azul sediments largely followed the distribution of GDGTs (cf. example in Fig. 2). In addition to the regular GDDs (labelled GDD-0 to GDD-Cren; Fig. 2a, b), both a series of late and early eluting peaks were observed. The distribution of the late eluting peaks mirrored that of the S-GDGTs and they were identified as S-GDDs based on their later retention time compared to the regular GDDs, as well as their MS<sup>2</sup> spectra. The MS<sup>2</sup> spectrum of S-GDD-1 (0:0:1; Fig. S1a) exhibited an [M

#### Table 1

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Relative abundances (%) and standard deviations of GDGTs (GDDs), GMGTs (GMDs), S-GDGTs (S-GDDs) and uns-GDGTs in examined samples.
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Samples		Туре	GDGTs	GMGTs	S-GDGTs	Uns- GDGTs	GDDs	GMDs	S-GDDs
Lakes									
Messel $(n = 18)$		Lacustrine oil shale	$\textbf{90.2} \pm \textbf{5.9}$	$\textbf{2.7} \pm \textbf{1.8}$	$\textbf{7.2} \pm \textbf{4.3}$	-	$\textbf{84.1} \pm \textbf{8.9}$	$\textbf{5.5} \pm \textbf{3.6}$	$10.4 \pm 5.8$
Lake Chala ( $n = 168$ )		Sediment core	$95.0 \pm 2.5$	-	$\textbf{2.8} \pm \textbf{1.6}$	$2.1\pm1.2$	$\textbf{97.9} \pm \textbf{2.1}$	-	$\textbf{2.1} \pm \textbf{2.1}$
Azorean lakes (n = 18)		Surface sediment	$95.9 \pm 3.5$	-	$1.8 \pm 1.3$	$2.3\pm2.3$	$94.1\pm4.0$	-	$\textbf{5.9} \pm \textbf{4.0}$
Marine/hypersaline									
Monterey(Naples Beach)		Miocene marine sediments	$\textbf{99.8} \pm \textbf{0.2}$	$\textbf{0.2}\pm\textbf{0.2}$	-	-	100	-	-
(n = 10)									
E. Mediterranean ( $n = 12$ )		Pleistocene sapropel	100	-	-	-	100	-	-
W. Mediterranean $(n = 3)$		Sediment core	100	-	-	-	100	_	-
Arabian Sea (n = 3)		Surface sediment	100	-	-	-	100	_	-
Vena del Gesso (n = 12)	bed IV, V, VII & XI	Hypersaline marls	$\textbf{97.1} \pm \textbf{1.9}$	$\textbf{2.5} \pm \textbf{1.5}$	$\textbf{0.4}\pm\textbf{0.6}$	-	100	_	-
Suboxic/euxinic marine									
Black Sea (n = 15)		SPM <sup>a</sup>	100	-	-	-	100	_	_

<sup>a</sup> SPM – Suspended Particulate Matter.



Fig 3. Relative distributions (%) of all GDGTs (A) and GDDs (B) identified in Messel oil shale sediments, Azorean lakes surface sediments and investigated Lake Chala core sediments.

+ H]<sup>+</sup> ion at m/z 1244.281 and [M + H-H<sub>2</sub>O]<sup>+</sup> at m/z 1226.270. Additional diagnostic fragments included a fragment ion at m/z 667.660 which represents the loss of 576.620 (C<sub>40</sub>H<sub>80</sub>O; an alkyl chain with ring, presumably the cyclohexane ring). To the best of our knowledge this represents the first report of S-GDDs.

GMDs, diether analogues of GMGTs, were identified based on their

retention time and comparison with published spectra (Bauersachs and Schwark, 2016). The  $MS^2$  spectrum of GMD-0 contained a  $[M + H]^+$  ion at m/z 1244.281, with the only significant fragment ions at m/z 1226.270 and 1208.259 representing sequential losses of water. The presence of *uns*-GDDs could not be unambiguously confirmed.



Fig. 4. PCA analysis of relative abundances of all identified GDGTs and GDDs in all investigated sediments (lacustrine, marine, hypersaline) grouped as unsaturated-, H-shaped (GMGT/GMD), regular- and S-shaped. The color code for the different GDGT and GDD groups and their contribution is indicated. Component score plot for PC1 and PC2 for the total of 259 core and surface sediments of 10 lakes, marine sediments, surface sediments, sapropels, evaporites and SPM combined (204 of which are lake sediments; 55 marine). Note: Lake Chala sediments plotted here include 14% turbidites (i.e sediments originating from a more peripheral location that end up at the drill site after bottom-slope failure) that were present in the investigated section of the core.

#### 3.2. Distribution of isoprenoidal ether lipids

The relative abundance of the various classes of GDGTs and GDDs (regular, unsaturated, GMGTs and S-GDGTs) in the different sediments are summarized in Table 1. Regular GDGTs and regular GDDs were detected in all lacustrine and marine sediments with GDGTs dominating (84 to 95 % of total; Table 1; Fig. 3). uns-GDGTs (but no uns-GDDs) were detected in the Azorean lakes and Lake Chala (ca. 2 %) but not in the Messel oil shale and in the marine sediments. GMGTs were detected in the Messel oil shale sediments, the Miocene Monterey sediments, and the Vena del Gesso sediments (Table 1), while GMDs were only detected in the Messel oil shale sediments. S-GDGTs and S-GDDs were detected in all lacustrine sediments (Messel, Chala and the Azorean lakes) but not in the marine settings, with the exception of S-GDGTs that were present in two out of four Vena del Gesso beds investigated (beds VII and XI; Table 1). The detection of S-GDGTs in Messel oil shales was previously reported by Liu et al. (2016a). The average abundance of S-GDGTs (% of total GDGTs investigated in this study, Table 1) and S-GDDs (% of total GDDs investigated in this study, Table 1) were 7.2  $\pm$  1.0 and 10.4  $\pm$  1.4 respectively in Messel oil shales; 1.8  $\pm$  0.3 and 5.9  $\pm$  0.9 in the Azorean lakes surface sediments and 2.8  $\pm$  0.1 and 2.1  $\pm$  0.2 in Lake Chala sediments (Table 1; Fig. 3).

A PCA was performed on the data calculated from relative abundances of investigated GDGTs and GDDs to constrain the variability (Fig. 4). The first principal component (PCA1) accounts for 59.6 % of the total variance and the second (PCA2) for 28.7 % (Fig. 4) for 259 samples investigated (204 lacustrine and 55 marine). Regular GDGTs and GDDs

were negatively loaded on the first principal component, while GMGTs, GMDs, S-GDGTs and S-GDDs were positively loaded. *uns*-GDGTs were positively loaded on the second principal component. All the Messel oil shales clustered in the same quadrant due to the relatively high abundance of GMGTs and GMDs. The Lake Chala sediment core samples and the majority of the Azorean lakes surface sediments clustered together, due to the presence of S-GDGTs. Most of the marine samples (Monterey, Vena del Gesso sediments, Black Sea SPM, Mediterranean sapropels and sediment core and Arabian Sea surface sediments) clustered in one quadrant. Overall, the PCA shows a clear division between the marine and lacustrine sediments based on ether lipid abundances, with a distinct difference between the Messel oil shale sediments and the other lacustrine samples.

#### 3.3. Evaluation of ether lipids as indicators of lacustrine anoxia

#### 3.3.1. S-GDGTs and S-GDDs

Neither S-GDGTs nor S-GDDs were detected in any of the marine sediments, but they were present in the majority of the lacustrine sediments (Table 1). Previously, based on their presence in terrestrial sulfidic settings, Liu et al. (2016a) hypothesized that S-GDGTs are indicative of sulfidic conditions. This led Liu et al. (2016a) to suggest that S-GDGT-producing organisms may represent novel archaeal species which mediate the cycling of sulfur and terrestrial organic matter in coastal environments. However, the detection of S-GDGTs in the Azorean lakes, Messel oil shale sediments and Lake Chala sediments casts doubt on this hypothesis. For example, the Messel core section

investigated in this study corresponds to the Middle Messel-Formation, which was formed under anoxic-non-sulfidic conditions (Bauersachs et al., 2014). Deep Azorean lakes are monomictic and show seasonally induced water column stratification with bottom water anoxia during summer months (lakes Funda, Negra, Azul and Verde) (Fig. S3). Lakes Empadadas and São Jorge are lakes which are usually oxic throughout the year and Lake Cubres represents a coastal lagoon with one side more exposed to the Atlantic Ocean (sample SS6, West side; Table S1). On average,  $S_{tot}$  was < 0.5 % (between 0.2 % in surface sediments of Lake Azul to 0.5 % in Lake Lomba), except for Lake Cubres (2.3 % on average), possibly due to the periodic inundations of the lake with seawater. These low sulfur contents make it unlikely that the bottom waters were euxinic. Some H<sub>2</sub>S can be produced in the water column of these lakes during the biological processes of sulphate reducing bacteria where available methane acts as electron-donor, as reported for other Azorean lakes by Tassi et al. (2018). Finally, the Lake Chala water column below 40 - 60 m is permanently stratified and anoxic (Wolff et al., 2011; Buckles et al., 2014), however with no evidence that anoxic bottom waters are sulfidic (e.g., low % of sulfur in sediment trap material, up to 1 %; Wolff et al., 2014). The same trap samples also show a positive correlation between sulfur and TOC ( $R^2 = 0.80$ ), indicating that most sulfur is bound to organic matter (Wolff et al., 2014) making it highly unlikely that the remaining sulfur content was high enough to induce bottom water euxinia.

Interestingly, in Lakes Negra (SS1 and SS2) and Funda (SS1), which are deep lakes that have seasonally anoxic bottom waters, S-GDGTs and S-GDDs were not detected. This suggests that the archaeal producers of S-GDGTs and S-GDDs are found in lakes with anoxic bottom waters, but that anoxia does not guarantee their presence, and thus additional factors play a role in determining their production. In these deep, eutrophic lakes this could partly be due to a sharp gradient of physico-chemical properties in the water column (pH, dissolved oxygen, turbidity and nutrients; https://www.azores.gov.pt/Gra/srrn-drotrh/menus/principa l/Monitorizacao/) that limits the niche for archaeal producer(s) of S-GDGTs and S-GDDs.

The absence of S-GDGTs and S-GDDs in nearly all marine sediments (Table 1) supports the previous hypothesis that they are produced by freshwater archaea (Liu et al., 2016a). However, we detected S-GDGTs in two beds of Vena del Gesso (VII and XI; Table 1), sediments deposited in an ancient evaporitic marine basin, with likely hypersaline conditions (Vai and Ricci Lucchi, 1977; Hsü et al., 1977). This fits with reported occurrence of S-GDGTs in salt marshes (Liu et al., 2016b) suggesting that S-GDGT producing archaea could be present in small saline lakes and evaporitic basins. Overall, our results suggest that S-GDGTs and S-GDDs are likely indicative of anoxic conditions (both intermittent and permanent) in lacustrine environments. Further studies are needed to determine their archaeal producer(s) and their biosynthetic pathway.

#### 3.3.2. Unsaturated GDGTs

Unsaturated GDGTs (uns-GDGTs) were reported in a suite of marine environments by Zhu et al. (2014) and based on their abundances and distributions, the authors suggested that they constitute an important archaeal lipid group in sediments deposited in euxinic basins and seep sites. Uns-GDGTs were also reported in a lacustrine environment (Fayetteville Green Lake surface sediment) by Liu et al. (2016a). Even though unsaturated diphytanyl glycerol diethers, such as archaeol, with multiple unsaturations (1-6) have been previously reported in Halorubrum lacusprofundi (a psychrotroph originally isolated from hypersaline Deep Lake in Antarctica; Gibson et al., 2005) and various halo(alkali) philic euryarchaeal strains isolated from hypersaline lakes (Bale et al., 2019b), uns-GDGTs are much more uncommon. For unsaturated diether lipids it was found that their proportion increases with increasing NaCl concentration in cultured halophilic archaea- which indicates that their biosynthesis represents a phenotypic adaptation (Maestrojuan et al., 1992; Hafenbradl et al., 1993; Dawson et al., 2012; Bale et al., 2019b). In this study, uns-GDGTs seem to be restricted to Lake Chala and almost all

Azorean lakes (Table 1), while respective *uns*-GDDs were not detected. The presence of *uns*-GDGTs in the Azorean lakes coincides with the detection of S-GDGTs (with the exception of Lakes Negra and Funda, Fig. 3), and hence their presence in these settings may also be associated with low-oxygen conditions that occur during summer months. However, their absence in older samples, such as in Messel oil shales, suggests poor preservation potential for geological studies. Furthermore, their reported presence in marine systems (Zhu et al., 2014) introduces ambiguity to their association with lacustrine settings.

#### 3.3.3. GMGTs and GMDs

GMGTs, glycerol mono-alkyl glycerol tetraethers, also called H-GDGTs, were first identified in a hyperthermophilic methanogen and hypothesized to be an adaptation to heat stress in order to maintain membrane stability at high temperatures (Morii et al., 1998). Subsequently, GMGTs have been identified by Schouten et al. (2008a, b) in Aciduliprofundum boonei (obligate thermoacidophilic euryarchaeote) and a variety of marine and lacustrine sediments. More recently, GMGT-0 was reported in a sediment core from the New Jersey Shelf, USA (last 90 Myrs) (de Bar et al., 2019) and diverse GMGTs were detected in Pyrococcus furiosus (piezosensitive, hyperthermophilic archaeon) by Tourte et al. (2020). Naafs et al. (2018) reported GMGTs in globally distributed peatland soils. GMGTs were also detected in two peat cores from NE China (Tang et al., 2021). GMDs (glycerol monoalkene diol diethers) were reported in sediments deposited under hydrothermal conditions suggesting that (hyper)thermophilic Archaea may constitute a biological source (Bauersachs and Schwark, 2016). Liu et al. (2016b) reported GMD-0 in oil contaminated surface sediment from Guaymas Basin (Mexico) while de Bar et al. (2019) found it throughout a sediment core (New Jersey Shelf, USA) spanning last 90 Myrs.

In this study we detected GMGTs in lacustrine, marine and hypersaline environments, in agreement with their reported occurrence in a range of different lacustrine, marine (Schouten et al., 2008a, b; Liu et al., 2012b), and terrestrial settings (Naafs et al., 2018). However, GMDs were only detected in Messel oil shale sediments. Bauersachs and Schwark (2016) proposed that GMDs occur in hydrothermally affected environments, conditions which are not applicable for the Messel paleolake environment. It is more likely that, as was found for GMGTs (Schouten et al., 2008b), they can be synthesized by non-thermophilic archaea at low temperatures as well. More studies are needed to assess how widespread GMDs are in a variety of natural settings and what is their connection to respective GMGTs.

# 3.4. The relationship between GDGTs and GDDs

It has been postulated that GDDs are either degradation products of GDGTs or that they are biosynthesized by archaea, either as membrane components or as GDGT intermediates (Liu et al., 2012a; Yang et al., 2014; Meador et al., 2014; Coffinet et al., 2015; de Bar et al., 2019). We investigated the relationship between GDGTs and GDDs by first comparing the relative ring distributions of GDDs to those of GDGTs by calculating ring indices (RI; He et al. 2012) whereby we modified the RI equation to include the S-GDGTs and S-GDDs (section 2.6). The degree of cyclization of GDDs (Fig. S4) irrespective of the sediment age. The strongest correlation was in the Lake Chala sediment core (n = 168) and the Azorean lake surface sediments (n = 18) (both  $r^2 = 0.97$ , Fig. S4). Similar distribution of rings in GDGTs and GDDs argues for a similar source and thus potentially a diagenetic conversion of GDGTs into GDDs.

We further examined this relationship by comparing the abundance (peak area/g sediment) of S-GDGTs with those of S-GDDs. We found strong positive correlations in the Messel oil shale sediment core (S-GDGT-1 vs S-GDD-1:  $r^2 = 0.96$ , n = 14 and S-GDGT-2 vs S-GDD-2:  $r^2 = 0.92$ , n = 14; Fig. 5) and in the Azorean lake surface sediments (S-GDGT-1 vs S-GDD-1:  $r^2 = 0.79$ , n = 14; Fig. 5), potentially suggesting a diagenetic origin of GDDs. However, S-GDDs in many samples were



Fig. 5. Correlation graphs showing S-GDGT-1 vs S-GDD-1 and S-GDGT-2 vs S-GDD-2 (peak area per g sediment) in Messel oil shale sediment core (a), Azorean lake's surface sediments (b) and Lake Chala sediment core (c).

below detection level, leading to a poor correlation in the Azorean lakes (S-GDGT-2 vs S-GDD-2:  $r^2 = 0.29$ , n = 4, Fig. 5) and in the Lake Chala sediment core (S-GDGT-1 vs S-GDD-1,  $r^2 = 0.30$ , n = 112 and S-GDGT-2 vs S-GDD-2:  $r^2 = 0.32$ , n = 93; Fig. 5). Furthermore, GDD equivalents of *uns*-GDGTs as well as GMGTs were rarely found (Table 1). Finally, we did not observe an increase in GDDs versus GDGTs in the sediment cores of Lake Chala and Messel oil shale (Fig. S5) with depth as would be expected from a diagenetic conversion of GDGTs into GDDs and which has been observed previously (Yang et al., 2014). Therefore, it seems unlikely that GDDs are a direct diagenetic product of GDGTs. Overall, the observations from our study support the theory of a joint cellular origin for GDGTs and GDDs.

# 4. Conclusions

In this study we report a suite of isoprenoidal GDGTs and GDDs in modern and ancient lacustrine settings. Regular GDGTs and GDDs are the most abundant of all investigated isoprenoid lipids, while emphasis was on examining the potential of GMGTs, *uns*-GDGTs and newly reported S-GDGTs and their respective GDDs, as anoxia-related markers. GMDs were only detected in Messel oil shale, indicating they could be synthesized by non-thermophilic archaea much the same as GMGTs and show the same preservation potential in the geologic record. *Uns*-GDGTs were present in recent lacustrine settings (Azorean lakes and Lake Chala), both in surface sediments and in the core, while *uns*-GDDs could not be confidently identified. S-GDGTs and newly reported S-GDDs were detected in all lacustrine sediments which coupled with reports from previous studies on bottom water anoxia in these lakes, makes them potential indicators of low-oxygen or anoxic conditions. They were not detected in marine sediments and SPM with the exception of Vena del Gesso marls, which represent an example of an ancient hypersaline lagoonal system, indicating ability of S-producers to live in more saline conditions in an enclosed environment. Although their archaeal precursor(s) is/are not determined as of yet, we propose that the S-GDGTs and S-GDDs are produced in anoxic bottom waters of predominantly freshwater lakes, and in other stagnant water systems such as saline lagoons with established marine connection.

# **Declaration of Competing Interest**

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

# Data availability

Data will be made available on request.

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# Appendix A. Supplementary data

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