



Composition and sources of sedimentary organic matter in the deep eastern Mediterranean Sea

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Received: 3 June 2015 – Published in Biogeosciences Discuss.: 3 July 2015

Revised: 20 November 2015 – Accepted: 28 November 2015 – Published: 16 December 2015

Abstract. Surface sediments collected from deep slopes and basins (1018–4087 m depth) of the oligotrophic eastern Mediterranean Sea have been analysed for bulk elemental and isotopic composition of organic carbon, total nitrogen and selected lipid biomarkers, jointly with grain size distribution and other geochemical proxies. The distribution and sources of sedimentary organic matter (OM) have been subsequently assessed and general environmental variables, such as water column depth and physical circulation patterns, have been examined as causative factors of deep-sea sediment characteristics. Lithogenic and biogenic carbonates are the dominant sedimentary fractions, accounting for up to 85.4 and 66.5 % of the total weight respectively. The low OC and TN contents in the surface sediments of the study area, which ranged from 0.15 to 1.15 % and 0.06 to 0.11 % respectively, reflect the oligotrophic character of the eastern Mediterranean Sea. Both bulk and molecular organic tracers reflect a mixed contribution from autochthonous and allochthonous sources for the sedimentary OM, as indicated by relatively degraded marine OM, terrestrial plant waxes and anthropogenic OM (e.g. degraded petroleum by-products) respectively. Wide regional variations have been observed amongst the studied proxies, which reflect the multiple factors controlling sedimentation in the deep eastern Mediterranean Sea. Our findings highlight the role of deep eastern Mediterranean basins as depocentres of organic-rich fine-grained sediments (mean $5.4 \pm 2.4 \mu\text{m}$), with OM accumula-

tion and burial being attributed to aggregation mechanisms and hydrodynamic sorting. A multi-proxy approach is applied aiming to investigate the biogeochemical composition of sediment samples, which sheds new light on the sources and transport mechanisms along with the impact of preservation vs. diagenetic processes on the composition of sedimentary OM in the deep basins of the oligotrophic eastern Mediterranean Sea.

1 Introduction

The burial of organic matter (OM) in marine sediments constitutes the main link between “active” pools of carbon in the oceans, atmosphere and landmasses and carbon pools that cycle on much longer, geological, timescales (Burdige, 2007). Therefore, investigating the processes that control the composition of sedimentary OM that is buried in deep-sea sediments is crucial for understanding carbon cycling on a global scale.

The deep sea receives inputs of organic particles from multiple sources, both autochthonous (e.g. biogenic particulate matter from primary production in ocean surface waters) and allochthonous (i.e. land-sourced OM from soils, plant debris, riverine phytoplankton and man-made compounds transported by runoff and atmospheric deposition into the marine domain) (Bouloubassi et al., 1997; Durrieu

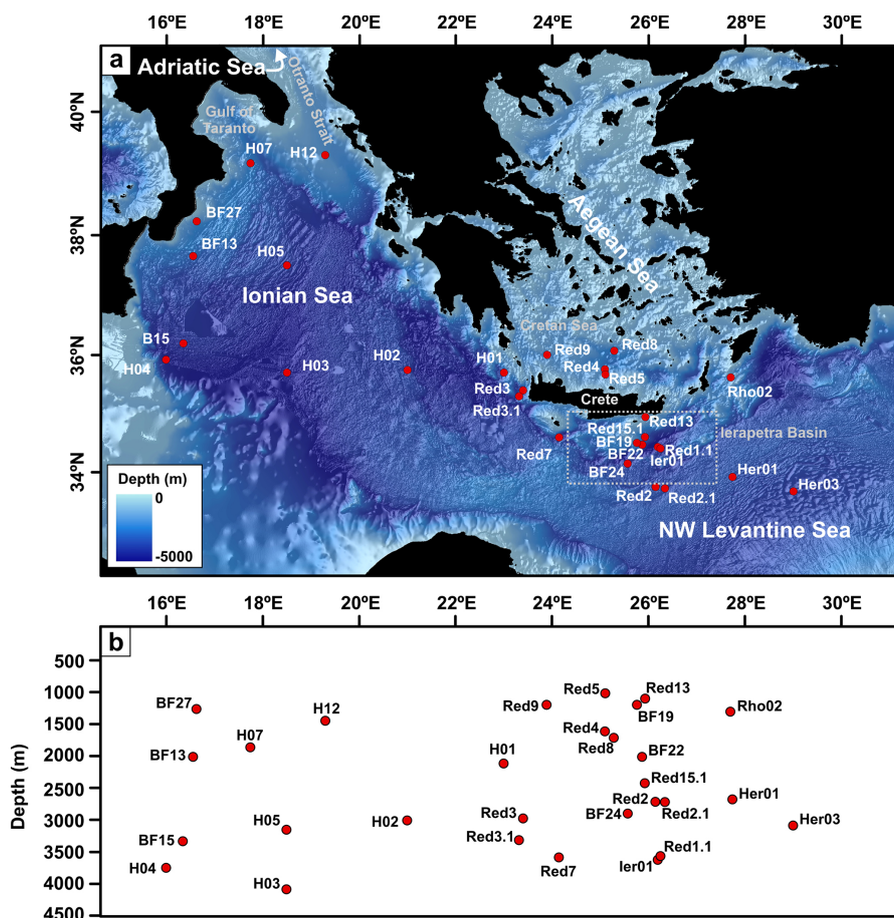


Figure 1. (a) Location of sampling sites across the open eastern Mediterranean Sea (see also Table 1). The map was produced using GEBCO Digital Atlas (IOC, IHO and BODC, 2003). (b) Plot of longitude vs. water depth of sampling stations.

de Madron et al., 2000; Kaiser et al., 2014). Consequently, sedimentary OM constitutes a heterogeneous and complex mixture of organic compounds with a wide range of chemical and physical properties (Mayer, 1994; Hedges and Oades, 1997; Hedges et al., 1997; Goñi et al., 1998). Therefore, the combined use of bulk geochemical indicators such as total nitrogen (TN) to organic carbon (OC) ratios, stable isotope of OC ($\delta^{13}\text{C}$) and molecular proxies such as lipid biomarkers can aid to gain knowledge on the origin, delivery and preservation of OM in marine sediments (Bouloubassi et al., 1997; Meyers, 1997; Goñi et al., 2003; Volkman, 2006).

The biogeochemical composition of sediments in deep basins of the oligotrophic eastern Mediterranean Sea (EMS), as well as the sources, transport and preservation of sedimentary OM, have been scarcely investigated so far. Previous studies have shown that the composition of surficial sediments is principally controlled by the geochemical characteristics of the source areas, the prevailing metoceanic conditions on the adjacent shelves, the contribution of atmospheric aerosols and the dominant regional circulation (e.g. Weldeab et al., 2002; Ehrmann et al., 2007; Hamann et al., 2008). Nev-

ertheless, the factors involved in the supply, distribution and fate of sedimentary OM are still poorly known.

In the present study, surface sediments collected from deep slopes and basins of the EMS have been analysed for physical and geochemical parameters such as grain size distribution, lithogenic, calcium carbonate (CaCO_3), opal, OC and TN contents, along with molar TN / OC ratios, stable isotopic ratios of OC ($\delta^{13}\text{C}$) and selected lipid biomarkers. Our main goal is to investigate the spatial distribution and main sources of sedimentary OM and to evaluate the impact of autochthonous vs. allochthonous contributions in the study area. We also examine whether and up to which point general environmental factors, such as water mass circulation patterns and water column depth, could explain the observed deep-sea sediment geochemical properties.

2 Oceanographic setting

The EMS is a land-locked sea with a complex topography including shelves, slopes, ridges, seamounts, trenches and four main basins: the Adriatic Sea, the Ionian Sea, the Aegean Sea

and the Levantine Sea (Fig. 1) (Amblàs et al., 2004; Medimap Group, 2007). The Ionian Sea to the west and the Levantine Sea to the east are longitudinally connected and cover most of the EMS area. They are also the deepest basins of the EMS, with the maximum depth (5267 m) located at the Hellenic Trench, south of the Cretan Arc. The Aegean Sea and the Adriatic Sea represent the northern extensions of the EMS. Both are relatively shallow, in particular the Adriatic Sea, which is dominated by a broad shelf and a slope sub-basin shallower than 1200 m. In the Aegean Sea, which has a particularly complex topography with tens of depressions, highs and islands, water depths up to 2500 m are found north of the island of Crete (Amblàs et al., 2004; Medimap Group, 2007). The southern Aegean Sea (Cretan Sea) is the sea area comprised between the Cyclades Archipelago to the north and the island of Crete to the south, which also includes the western Cretan Straits.

The general circulation pattern of the EMS is anti-estuarine, which results from interactions between basin, sub-basin and mesoscale flows (Bethoux, 1979). The EMS communicates with the western Mediterranean Sea through the Sicily Strait, with an inflow of low-salinity modified Atlantic water (MAW) at the upper 100–150 m of the water column (Rabitti et al., 1994; Malanotte-Rizzoli et al., 1997). MAW flows in an easterly direction getting progressively saltier and warmer until it transforms into Levantine intermediate water (LIW) into the Levantine Sea where it sinks to mid depths (Milliff and Robinson, 1992; Lascaratos et al., 1993).

The eastern Mediterranean deep water (EMDW) is a relatively well-oxygenated water mass, likely as a result of the formation and sinking of warm deep-water that ventilates the deepest levels in the EMS (Schlitzer et al., 1991; Roether and Well, 2001; Meador et al., 2010). Waters from the Adriatic Sea (Adriatic deep waters) have been considered as the main contributor of deep and bottom waters to the EMS (Malanotte-Rizzoli and Hecht, 1988). Nevertheless, the Aegean Sea constitutes a sporadically significant contributor to EMDW through the Cretan deep waters, as in the case of the eastern Mediterranean Transient anomaly in the 1990s (Lascaratos et al., 1999; Theocharis et al., 1999). Additionally, the Aegean Sea constitutes a possible secondary source of intermediate waters to the adjacent Ionian and Levantine seas, through outflows across the Cretan Arc straits (Robinson et al., 2001).

Key factors that control the exchanges through the Cretan Arc straits are the thermohaline properties of water masses and mesoscale variability. For example, the Ierapetra anticyclonic gyre, which is located off the southeast corner of Crete (Ierapetra Basin), exhibits a strong seasonal signal that is linked to variations of the outflow across the eastern Cretan Arc straits (Theocharis et al., 1993; Larnicol et al., 2002). Actually, the several permanent and/or recurrent eddies in each of the EMS sub-basins enhance exchanges between continental shelf and slope waters (Robinson et al., 1992;

Malanotte-Rizzoli et al., 1997; Millot and Taupier-Letage, 2005), which in turn influence primary productivity and the settling of OM to the deep-sea floor (Danovaro et al., 2010).

Thermohaline circulation and overall environmental conditions make the EMS one of the most ultra-oligotrophic environments of the world ocean (Psarra et al., 2000; Krom et al., 2005; Thingstad et al., 2005; Gogou et al., 2014). Annual primary production in the EMS averages between 121 and 145 g C m⁻² yr⁻¹ (Bosc et al., 2004; Gogou et al., 2014). However, the fraction of primary production exported below 2000 m of water depth averaged 0.3 % (Gogou et al., 2014). The low autochthonous contribution to OM fluxes in the deep open EMS is counterbalanced by allochthonous inputs resulting from long-range atmospheric transport and deposition by aerial dust (Jickells, 1995; Gogou et al., 1996; Tsapakis and Stephanou, 2005). The overall sedimentation rate in the deep areas of the open EMS is low (i.e. 2–5 cm kyr⁻¹) (Van Santvoort et al., 1996, 2002; Garcia-Orellana et al., 2009; Stavrakakis et al., 2013).

3 Materials and methods

3.1 Sampling

Short sediment cores were collected with a multicorer at 29 stations, ranging from 1018 to 4087 m water depth, during six oceanographic cruises in the Ionian Sea, the southern Aegean Sea (Cretan Sea) and the northwestern Levantine Sea from January 2007 to June 2012 (Fig. 1 and Table 1). Once onboard, multicores were visually described and sliced at 1 cm intervals. Sub-samples collected for grain size and elemental and stable isotopic composition were stored in sealed plastic bags at 4 °C, while those collected for the analysis of lipid biomarkers were stored in pre-combusted aluminium foils at –20 °C. Only the undisturbed top centimetre of each sediment core is considered in this study.

3.2 Analytical procedures

3.2.1 Particle size characterization

The grain size distribution of sediment samples was determined using a Coulter LS230 Laser Diffraction Particle Size Analyzer, which measures sizes between 0.04 and 2000 µm. Prior to analysis, freeze-dried samples were oxidized with a 10 % H₂O₂ (v/v) solution in order to remove OM. Each sample was then divided into two sub-samples, one of which was treated with 1M HCl to remove carbonates and thus obtain the grain size distribution of lithogenic (siliciclastic) particles. Subsequently, both bulk and lithogenic fractions were dispersed into 20 cm³ of a 5 % NaPO₅ (v/v) solution and mechanically shaken for 4 h and then introduced into the particle size analyzer after using a 2000 µm sieve to retain occasional coarse particles that might obstruct the flow circuit of the instrument.

The measured particle size spectrum is presented as % volume in a logarithmic scale, where volume is calculated from particle diameter, assuming spherical shapes. Results were recalculated to percentages of clay (< 4 µm), silt (4–63 µm) and sand (63 µm–2 mm).

3.2.2 Elemental and stable isotopic analysis of carbon and nitrogen

For the determination of total carbon (TC), TN, OC contents and stable isotopic composition of OC ($\delta^{13}\text{C}$) freeze-dried and ground sediments were analysed using a Flash 1112 EA elemental analyser interfaced to a Delta C Finnigan MAT isotope ratio mass spectrometer. Samples analysed for % OC and $\delta^{13}\text{C}$ were initially de-carbonated using repetitive additions of a 25 % HCl (*v/v*) solution, separated by 60 °C drying steps, until no effervescence was observed (Nieuwenhuize et al., 1994). Stable isotope data are reported using the conventional per thousand $\delta^{13}\text{C}$ notation relative to the Pee Dee Belemnite standard. Uncertainties for elemental composition were lower than 0.1 %, while uncertainty for $\delta^{13}\text{C}$ was lower than 0.05 ‰.

In consistency with published data in the Mediterranean Sea we assumed OM as twice the OC content (e.g. Heussner et al., 1996; Masqué et al., 2003). The inorganic carbon content was calculated from the difference between TC and OC measurements. Assuming all inorganic carbon is contained within calcium carbonate, CaCO_3 content was determined using the molecular ratio of 100/12.

Molar TN/OC ratios were also calculated. TN/OC is plotted in order to constrain the elemental ratios of N-depleted samples (i.e. $\text{TN/OC} \approx 0$ rather than $\text{OC/TN} \infty 0$) following Goñi et al. (2006) and to avoid the underestimation of the terrestrial-derived carbon fraction (Perdue and Koprivnjak, 2007).

3.2.3 Biogenic opal and lithogenic fraction analysis

The biogenic silica content was analysed using a two-step 2.5 h extraction with a 0.5 M Na_2CO_3 solution, separated by centrifugation of the leachates. Si and Al contents of both leachates were analysed with a Perkin-Elmer Optima 3200RL Inductive Coupled Plasma Optical Emission Spectrometer (ICP-OES), correcting the Si content of the first leachate by the Si/Al ratio of the second one. All values are reported as opal ($\text{SiO}_2 \cdot 0.4\text{H}_2\text{O}$), a parameter defined by 2.4 times the weight percentage of biogenic Si content determined for each sample (Mortlock and Froelich, 1989). The opal detection limit, associated with the detection limit of the ICP-OES system, is approximately 0.2 %.

The lithogenic fraction was estimated by subtracting the concentration of the major constituents from total dry weight (% lithogenic = $100 - [\% \text{OM} + \% \text{CaCO}_3 + \% \text{opal}]$). This fraction represents the residual component of particles such

as quartz, feldspars, clay minerals and aluminosilicates (Mortlock and Froelich, 1989).

3.2.4 Lipid biomarkers analysis and definitions of molecular indices

The analytical procedure followed for the determination of lipid biomarkers has been previously presented in detail (Gogou et al., 1998, 2000, 2007). Briefly, freeze-dried sediment samples were initially solvent-extracted three times by sonication with a dichloromethane : methanol mixture (4 : 1, *v/v*). Combined extracts were subsequently separated into different compound classes by column chromatography using silica gel that had been activated for 1 h at 150 °C. The following solvent systems were used to elute different compound classes: (1) *n*-hexane (fraction F₁, aliphatic hydrocarbons), (2) dichloromethane/*n*-hexane (fraction F₂, carbonyl compounds) and (3) ethyl acetate/*n*-hexane (fraction F₃, alcohols, sterols).

F₁ and F₃ fractions were analysed by gas chromatography–mass spectrometry (GC-MS) while F₂ fractions were analysed by gas chromatography using flame ionization detection (GC-FID). Hydroxyl-bearing compounds (fraction F₃) were derivatized to the corresponding trimethylsilyl ethers prior to GC-MS analysis using *N,O*-bis-(trimethylsilyl)-trifluoroacetamide (BSTFA) + 1 % trimethylchlorosilane (TMCS) for 1 h at 80 °C. Details regarding the GC instrumental parameters are presented elsewhere (Gogou et al., 2007; Parinos et al., 2013).

The individual lipids were identified by a combination of comparison of GC-retention times to authentic standards and comparison of their mass spectral data to those in the literature. Quantification was based on the GC-MS or GC-FID response and comparison of peak areas with those of known quantities of standards added prior to the extraction of the sediment samples ($[\text{}^2\text{H}_{50}]n$ -tetracosane for *n*-alkanes, *n*-hexatriacontane for long-chain alkenones, 5 α -androstan-3 β -ol for sterols and heneicosanol for *n*-alkanols). The signal of the unresolved complex mixture (UCM) of aliphatic hydrocarbons was defined by the chromatographic area (fraction F₁) between the solvent baseline and the curve defining the base of resolved peaks. UCM quantification was performed relatively to $[\text{}^2\text{H}_{50}]n$ -tetracosane using the average response factor of *n*-alkanes.

Procedural blanks processed in parallel to the samples were found to be free of contamination. Reproducibility of the analytical method based on multiple extractions of sediments was better than 6 % in all cases.

A range of selected lipid biomarkers are considered in this study, namely long-chain *n*-alkanes and *n*-alkanols, long-chain alkenones, long-chain diols and keto-ols and a suite of sterols, along with lipid biomarkers' indices, as proxies of organic matter sources and/or degradation. As OC can vary due to the supply of inorganic material (dilution effect) the con-

Table 1. Location, depth and collection date of sediment samples.

Sample code	Latitude (N)	Longitude (E)	Water depth (m)	Date of collection	Physiographic regions
Ionian Sea					
North					
H12	39.30	19.30	1450	Jan 2007	Otranto Valley
H07	39.17	17.75	1866	Jan 2007	Taranto Valley
Central					
BF27	38.22	16.63	1264	Jun 2009	Calabrian Slope
BF13	37.66	16.56	2012	Jun 2009	Calabrian Arc
BF15	36.20	16.35	3335	Jun 2009	Ionian Basin
H04	35.92	16.00	3750	Jan 2007	Ionian Basin
West					
H02	35.75	21.00	3008	Jan 2007	Ionian Basin
H05	37.50	18.50	3154	Jan 2007	Ionian Basin
H03	35.70	18.50	4087	Jan 2007	Ionian Basin
S Aegean Sea (Cretan Sea)					
Red5	35.68	25.10	1018	May 2010	Cretan Trough
Red9	36.00	23.89	1200	May 2011	Cretan Trough
Red4	35.76	25.10	1615	May 2010	Cretan Trough
Red8	36.07	25.28	1715	May 2011	Cretan Trough
W Cretan Straits					
H01	35.70	23.00	2117	Jan 2007	Kythira Strait
Red3	35.40	23.40	2976	May 2010	Antikythira Strait
Red3.1	35.30	23.32	3317	May 2010	Antikythira Strait
Red7	34.60	24.15	3589	May 2011	Ptolemy Strait
NW Levantine Sea					
Ierapetra Basin					
Red13	34.95	25.93	1101	Jun 2012	Cretan-Rhodes Ridge
BF19	34.51	25.76	1200	Jun 2009	Hellenic Trench
BF22	34.48	25.87	2015	Jun 2009	Hellenic Trench
Red15.1	34.61	25.92	2428	Jun 2012	Hellenic Trench
Red1.1	34.40	26.25	3568	Jun 2012	Hellenic Trench
Ier01	34.44	26.19	3626	Jan 2007	Hellenic Trench
Open Sea					
Rho02	35.62	27.70	1305	Jan 2007	Rhodes Strait
Her01	33.92	27.74	2680	Jan 2007	EM Ridge
Red2	33.74	26.15	2717	May 2010	EM Ridge
Red2.1	33.71	26.34	2720	May 2010	EM Ridge
BF24	34.15	25.57	2902	Jun 2009	Pliny Trench
Her03	33.67	29.00	3090	Jan 2007	Herodotus Basin

January 2007 samples were collected during the M71 (Leg 3) cruise onboard the R/V *Meteor* (University of Hamburg, Germany), June 2009 samples during the Biofun1 cruise onboard the R/V *Sarmiento de Gamboa* (CSIC-UB, Spain) and May 2010, 2011 and June 2012 samples during the ReDEco cruises onboard the R/V *Aegaeo* (HCMR, Greece).

concentrations of the reported lipid compounds were normalized to OC contents.

The sum of the concentrations of the most abundant high molecular weight *n*-alkanes and *n*-alkanols, which are major components of epicuticular higher plant waxes (Eglinton and Hamilton, 1967; Ohkouchi et al., 1997), are defined respectively as

$$\sum \text{TerNA} = \sum n\text{-C}_{27,29,31,33}, \quad (1)$$

$$\sum \text{TerN-OH} = \sum n\text{-C}_{24,26,28,30}. \quad (2)$$

The sum of the concentrations of the considered lipid biomarkers having a clear marine (algal) origin (see Sect. 5.1.2) was calculated as follows:

$$\sum \text{Mar} = \sum ({}_{28}\Delta^{5,22E} + {}_{30}\Delta^{22E} + \text{C}_{30}\text{diols \& keto-ols} + \text{alkenones}). \quad (3)$$

The abundance of the UCM of aliphatic hydrocarbons, a commonly observed persistent contaminant mixture in marine sediments consisting of branched alicyclic hydrocarbons (Gough and Rowland, 1990), is used as an indicator of the contribution from degraded petroleum products, i.e. chronic oil pollution in the study area (Wang et al., 1999).

The carbon preference indices of long-chain *n*-alkanes (CPI_{NA}) and *n*-alkanols ($\text{CPI}_{\text{N-OH}}$) have been used as indicators of terrestrial OM degradation with CPI values in fresh leaves being typically >4 (Collister et al., 1994). However, the abundance of non-degraded petroleum hydrocarbons could potentially bias (lower) CPI_{NA} values with increasing petroleum contribution, since *n*-alkane compounds of petroleum products present CPI_{NA} values ~ 1 (Wang et al., 1999). The indices were calculated respectively as

$$\text{CPI}_{\text{NA}} = \frac{\sum([n\text{-C}_{25}] - [n\text{-C}_{33}])}{\sum([n\text{-C}_{26}] - [n\text{-C}_{34}])}, \quad (4)$$

$$\text{CPI}_{\text{N-OH}} = \frac{\sum([n\text{-C}_{24}] - [n\text{-C}_{30}])}{\sum([n\text{-C}_{23}] - [n\text{-C}_{29}])}. \quad (5)$$

Finally, the abundance ratio of $\sum \text{TerNA}$ to $\sum \text{TerN-OH}$ is used as a proxy of the proportion of refractory vs. labile terrestrial components, since $\sum \text{TerNA}$ are more resistant to degradation than their alcohol counterparts (Eglinton and Hamilton, 1967; Ohkouchi et al., 1997). The ratio is defined as

$$\frac{[\text{NA}]}{[\text{N-OH}]} = \frac{[\sum \text{TerNA}]}{[\sum \text{TerN} - \text{OH}]}. \quad (6)$$

3.2.5 Data analysis and presentation

Statistical treatment of grain size data was carried out using the GRADISTAT v. 8.0 software (Blott and Pye, 2001). Median diameter (D_{50}), sorting and skewness were calculated geometrically (in metric units) following the approach of Folk and Ward (1957), which is most appropriate when data are non-normally distributed, as in the case of poly-modal sediments from the study area.

D_{50} was calculated as the average equivalent diameter, which is the diameter where 50% of the sediment sample has a larger equivalent diameter and the other 50% has a smaller equivalent diameter. Sorting (expressed by the standard deviation) indicates the fluctuation in the degree of kinetic energy and the depositional regime on grain size characteristics. Skewness measures the degree of asymmetry onto

particle distribution. The skewness for a normal distribution is zero, and any symmetric data should have skewness near zero. Positive values indicate skewness towards the finer grain sizes (skewed left) while negative values indicate skewness towards the coarser grain sizes (skewed right). The results of grain size distribution analysis were hierarchically clustered (using IBM SPSS Statistics 18.0) according to the above statistical parameters (autoscaled prior to cluster analysis) in order to determine the similarity of samples within each station measuring the squared Euclidean distance.

Principal component analysis (PCA) was performed on standardized grain size and elemental composition data (% clay and sorting of lithogenic and bulk fractions, lithogenic, CaCO_3 , OC and TN contents), on standardized bulk organic matter signatures (molar TN/OC ratios and $\delta^{13}\text{C}$) and on standardized indices and mass-normalized concentrations of lipid biomarkers. A subroutine, the Varimax rotation, was applied to the first three factors in order to maximize or minimize loadings within each factor and thus simplify the visual interpretation of PCA projections. Correlation analysis was also performed using the same variables. In certain occasions during correlation analysis, the singular stations that were clearly out of the trend were excluded in order to strengthen the evident trends of the examined variables. These stations are explained in detail in the discussion section (see Sect. 5.2).

The spatial distribution of the various geochemical parameters' contents, bulk OM signatures and selected lipid biomarkers' concentrations/indices considered in this study were visualized using Ocean Data View (Schlitzer, 2011).

4 Results

4.1 Grain size characteristics

The grain size composition (% clay, silt and sand) and the sedimentary parameters (D_{50} , sorting and skewness) are presented in Table 2, while the statistical dendrogram, type-averaged grain size spectrum and spatial distributions of grain size types are presented in Fig. 2.

Silt- and clay-sized particles dominate the bulk sediment, accounting for up to 76.7 and 57.1% of the total weight respectively (Table 2). The lowest values ($<40\%$) for the silt fraction are found in the upper slope of the western Cretan straits (station Red3) and the northwestern Levantine Sea (station BF19), while the highest values ($>65\%$) correspond to the Ionian Sea (stations H12 and H03). The lowest clay contents ($<20\%$) are also found in the upper slope of the western Cretan Straits (station Red3) but also in the north-eastern Ionian Sea (station H12), while maximum values ($>55\%$) are recorded at the northwestern Levantine Sea (station Red1.1 in Ierapetra Basin) and the western Cretan Straits (station H01). Sand contents show large variations, i.e. from 0 to 47.7% (station Red3 in the upper slope of western Cre-

Table 2. Bulk composition and sedimentological parameters of the investigated surface sediments.

Sample code	Clay _{bulk} (%, < 4 μm)	Silt _{bulk} (%, 4–63 μm)	Sand _{bulk} (%, 63 μm–2 mm)	Sorting _{bulk}	Sorting _{litho}	Skewness _{bulk}	D _{50 bulk} (μm)	Lithogenic (%)	CaCO ₃ (%)	Opal (%)	OC (%)	TN/OC	δ ¹³ C (‰)	OC _{mar} (%)	OC _{terr} (%)
Ionian Sea															
North															
H12	18.1	76.7	5.24	3.47	3.31	-0.13	14.6	71.2	28.5	0.03	0.15	0.08	-24.2	42.3	57.7
H07	41.2	58.8	0.00	3.01	4.12	-0.07	5.34	75.7	22.0		1.15	0.08	-24.4	40.2	59.8
Central															
BF27	52.8	46.1	1.04	3.38	2.83	-0.04	3.97	80.8	17.8	0.16	0.64	0.12	-23.0	60.0	40.0
BF13	50.8	48.1	1.08	3.41	2.92	-0.05	4.17	76.7	22.0	0.21	0.58	0.12	-22.9	62.7	37.3
BF15	39.1	57.0	3.98	4.01	2.91	0.03	5.88	77.9	21.6		0.28	0.14	-22.5	68.8	31.2
H04	41.6	58.4	0.04	4.69	3.53	-0.23	5.52	85.4	13.2	0.04	0.65	0.13	-23.6	50.9	49.1
West															
H02	53.3	46.7	0.00	2.99	6.16	-0.07	3.92	47.7	51.3	0.04	0.45	0.10	-22.5	68.6	31.4
H05	41.6	58.4	0.00	2.98	3.48	-0.01	5.34	60.7	38.2		0.57	0.11	-23.7	50.2	49.8
H03	34.5	65.5	0.08	3.11	3.49	-0.04	7.03	77.6	21.0	0.11	0.63	0.10	-24.6	35.8	64.2
S Aegean Sea (Cretan Sea)															
Red5	54.0	46.0	0.00	3.32	3.61	-0.01	3.88	32.5	66.5	0.24	0.39	0.10	-22.4	69.2	30.8
Red9	50.1	49.9	0.00	3.24	3.28	-0.01	4.23	46.6	52.2	0.15	0.55	0.12	-22.6	67.4	32.6
Red4	52.6	47.4	0.00	3.26	3.58	-0.06	4.00	39.3	59.8	0.09	0.42	0.12	-22.2	73.0	27.0
Red8	53.1	46.9	0.00	3.12	3.36	-0.03	3.97	44.0	55.3	0.08	0.33	0.15	-22.0	75.5	24.5
W Cretan Straits															
H01	55.1	44.9	0.00	4.74	4.77	-0.11	3.65	42.8	56.6		0.29	0.12	-22.6	66.5	33.5
Red3	14.5	37.9	47.7	5.23	4.54	-0.45	56.6	59.2	39.9	0.15	0.37	0.08	-18.3		
Red3.1	37.7	61.5	0.88	4.32	3.82	-0.07	6.74	55.7	43.0	0.12	0.58	0.09	-22.9	62.4	37.6
Red7	36.0	63.5	0.59	3.55	3.57	-0.01	6.37	61.4	37.5	0.07	0.51	0.12	-22.7	64.7	35.3
NW Levantine Sea															
Ierapetra Basin															
Red13	50.1	49.3	0.66	4.84	3.07	-0.15	4.23	59.1	39.8	0.22	0.46	0.11	-22.3	71.7	28.9
BF19	51.1	38.5	10.4	4.98	3.43	0.27	4.12	39.4	59.9		0.34	0.14	-22.6	67.3	32.7
BF22	47.5	44.0	8.53	4.71	3.34	0.20	4.56	57.7	41.7		0.26	0.15	-21.2	87.7	12.3
Red15.1	47.5	51.8	0.73	4.60	2.77	-0.03	4.59	51.8	46.9	0.80	0.61	0.08	-23.3	56.1	43.9
Red1.1	57.1	42.8	0.15	3.73	2.61	-0.13	3.53	51.8	47.1	0.05	0.54	0.10	-21.5	83.8	16.2
Ier01	49.1	50.4	0.00	3.84	2.67	0.03	4.35	55.6	43.3	0.06	0.52	0.12	-21.7	80.2	19.8
Open Sea															
Rho02	43.0	55.9	1.11	4.43	3.25	-0.14	5.24	58.4	40.6	0.16	0.47	0.12	-22.7	65.2	34.8
Her01	48.2	51.8	0.00	3.88	4.89	0.04	4.44	35.8	63.4	0.13	0.31	0.15	-22.7	65.5	34.5
Red2	32.4	53.2	14.4	4.95	3.85	-0.01	11.1	45.0	53.9	0.07	0.50	0.09	-24.0	46.2	53.8
Red2.1	48.7	51.1	0.21	4.30	3.92	0.11	4.42	44.2	55.0	0.10	0.37	0.11	-22.4	70.5	29.5
BF24	43.1	45.0	11.9	5.25	3.51	0.20	5.34	55.0	44.3	0.11	0.29	0.16	-22.3	70.6	29.4
Her03	39.8	60.2	0.00	4.05	3.20	-0.19	5.81	48.2	50.7	0.11	0.49	0.11	-22.4	70.2	29.8

Empty cells are not determined.

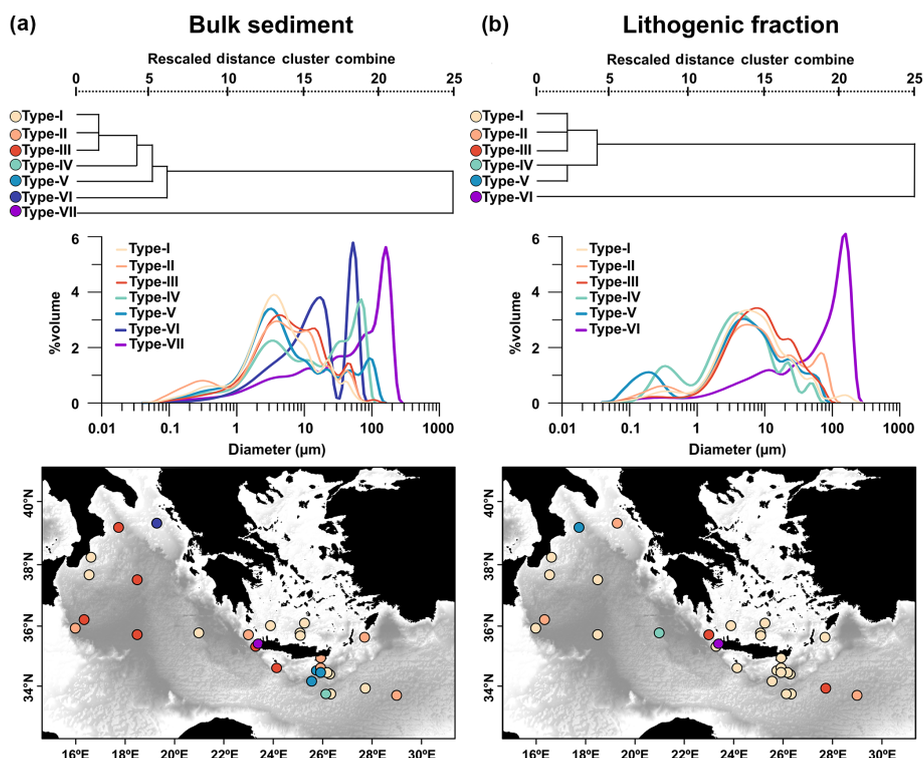


Figure 2. Statistical dendrogram of type-averaged grain size profiles and geographical distribution of grain size compositional types for (a) the lithogenic fraction and (b) the bulk fraction of the investigated sediments.

tan Straits), with values less than 2 % in most of the stations (Table 2). Relatively high values (> 10 %) are also obtained in the northwestern Levantine Sea (stations Red2, BF19 and BF24). D_{50} values range between 3.5 and 56.6 μm (Table 2).

Sorting of bulk sediment ranges from 3.0 to 5.2 (Table 2). Most of the northwestern Levantine Sea and western Cretan Straits' stations are very poorly sorted and all stations within the southern Aegean Sea and most of the Ionian Sea are poorly sorted (Table 2 and Fig. 2a). Skewness values for the investigated samples range from -0.45 to 0.27 (Table 2), varying from a very clear negative skewness in the upper slope of the western Cretan Straits (station Red3), to positive skewness in the northwestern Levantine Sea (stations Red2.1, BF19, BF22 and BF24).

The hierarchical cluster analysis of all bulk sediment samples resulted into seven grain size types (Fig. 2a). Most of the samples group into cluster types I ($n = 11$), II ($n = 6$) and III ($n = 6$), with grain size profiles almost symmetrical and poorly sorted, and a dominance of clay and silt fractions. Type V includes three very poorly sorted and positive skewed samples from the northwestern Levantine Sea, which consist mainly of clay and silt fractions. Types IV, VI and VII include only one sample each (Red2, H12 and Red3 respectively). Samples Red2 and Red3, dominated by coarse silt fractions (D_{50} 11.1 and 56.61 μm respectively), are both very poorly sorted but with different types of skewed distributions (symmetrical and negatively skewed respectively). Finally, sample

H12, composed mostly of fine silt, is poorly sorted and slight positive skewness.

As in the bulk sediment, silt- and clay-sized particles dominate the lithogenic fraction, accounting for up to 73.5 and 50.8 % of the total weight respectively. The hierarchical cluster analysis of the lithogenic fraction identified six grain size types (clusters) (Fig. 2b). A majority of samples are highly similar (types I–V), with an average composition of 35.2 ± 5.6 % clay, 63.5 ± 5.3 % silt and 1.3 ± 2.3 % sand, a D_{50} of 6.6 ± 1.3 μm and a bimodal or trimodal, symmetrical, poorly sorted grain size distribution. Sample Red3 from the upper slope of western Cretan Straits is an exception, belonging to type VI. The composition of its lithogenic fraction is 10.3 % clay, 34.6 % silt and 55.1 % sand, with a D_{50} of 78.1 μm and a bimodal, very poorly sorted and negatively skewed grain size distribution (Fig. 2b and Table 2).

4.2 Bulk geochemical sediment composition

The spatial variability of lithogenics, CaCO_3 , OC and TN contents within the study area is presented in Fig. 3a–c.

The lithogenics content in the analysed surface sediments range between 32.5 and 85.4 % (Fig. 3a). Higher percentages (> 70 %) are found in stations of the Ionian Sea (with the exception of station H02), while the lowest percentages (< 40 %) are found in the southern Aegean (stations Red4 and Red5) and northwestern Levantine seas (stations

Her01 and BF19) (Table 2). The CaCO₃ contents also show a wide range of values throughout the study area, from 13.2 to 66.5 % (Fig. 3b and Table 2). Stations in the western Ionian Sea (H04, BF27, H03, BF15, BF13 and H07) have the lowest CaCO₃ contents (< 22 %), whereas most stations in the southern Aegean and northwestern Levantine seas and in the western Cretan Straits have elevated CaCO₃ contents (> 40 %).

Opal contents are very low, ranging from below detection limits to a maximum of 0.24 % in the southern Aegean and northwestern Levantine seas (stations Red5 and Red13) (Table 2). Since opal contents are very close to the detection limits, those values can be considered as negligible. Therefore inorganic geochemical fraction of the investigated deep EMS sediments consists only of lithogenic (terrigenous) and carbonate components.

OC contents in the studied samples range from 0.15 to 1.15 %, with an average value of 0.47 % (Table 2). The lowest values are recorded in the northeastern Ionian Sea, south of Otranto Strait (station H12), while the highest values are found off the Gulf of Taranto (station H07), followed by stations in the Ionian Sea (stations BF27, H04 and H03). TN contents range from 0.01 to 0.11 % with an average value of 0.06 %. TN display a pattern similar to OC, also with the highest values recorded off the Gulf of Taranto (station H07) and the lowest south of Otranto Strait (station H12), both in the northern Ionian Sea.

4.3 Elemental and stable isotopic composition of sedimentary OM

The spatial distribution of molar TN/OC ratios and $\delta^{13}\text{C}$ within the study area is presented in Fig. 3d–e.

Molar TN/OC ratios of the sedimentary OM range from 0.08 to 0.16 (Fig. 3d and Table 2). The highest molar TN/OC ratios values (> 0.14) are recorded in the southern Aegean (station Red8) and northwestern Levantine (stations Her01, BF22 and BF24) seas, whereas the lowest molar TN/OC ratios (< 0.09) are recorded in stations from the northwestern Levantine Sea (station Red15.1), the western Cretan Straits (stations Red3 and Red3.1) and the northern Ionian Sea (stations H07 and H12). However, there is no clear spatial trend (Fig. 3d).

The spatial distribution of $\delta^{13}\text{C}$ values shows that relatively lower values are more common in the Ionian Sea than in the northwestern Levantine Sea, the southern Aegean Sea and the south Ionian Sea (Fig. 3e). $\delta^{13}\text{C}$ values range from –18.3 to –24.6 ‰ (Fig. 3e and Table 2), with stations from the Ionian Sea (stations H03, H07 and H12) yielding relatively depleted $\delta^{13}\text{C}$ values (< –24 ‰) and stations from the western Cretan Straits (station Red3) and the northwestern Levantine Sea (Ierapetra Basin, stations BF22, Red1.1 and Ier01) having relatively enriched $\delta^{13}\text{C}$ values (> –22 ‰).

4.4 Lipid biomarkers

The analysed sedimentary aliphatic hydrocarbons comprise of a series of resolved compounds, mainly *n*-alkanes, and a UCM (Parinos et al., 2013). The UCM dominate amongst aliphatic hydrocarbons in concentrations ranging between 0.50 and 6.64 mg g OC⁻¹ (Fig. 4a and Table 3). Maximum concentrations (> 5 mg g OC⁻¹) are recorded in the northwestern Levantine Sea (station Red2.1) followed by the deep central Ionian Sea (station H03). The lowest UCM values (< 1.0 mg g OC⁻¹) are obtained in the northern Ionian Sea (station H12) and in the western Cretan Straits (station H01), while rather low values are also recorded in the southern Aegean Sea and west of Crete.

The molecular profile of the *n*-alkanes is dominated by long-chain homologues ($C_n \geq 24$), maximizing at *n*-C₃₁, with elevated CPI_{NA} values (4.9 ± 1.6) (Table 3). $\sum\text{TerNA}$ range between 40.8 and 483 $\mu\text{g g OC}^{-1}$, with an average value of 172 $\mu\text{g g OC}^{-1}$ (Fig. 4b and Table 3). The station with the highest concentration (station Red2.1) is found in the northwestern Levantine Sea, while the stations with the lowest ones (< 110 $\mu\text{g g OC}^{-1}$) are located in the northern Ionian Sea (stations H07 and H12) and in the southern Aegean Sea (station Red9). Furthermore, relatively elevated $\sum\text{TerNA}$ concentrations (> 210 $\mu\text{g g OC}^{-1}$) are recorded at the deep station of the western Cretan Straits' (station Red3.1) and the deep central Ionian Sea (stations H03 and H05).

The aliphatic alcohol fraction is dominated by a series of *n*-alkanols ranging from *n*-C₂₂ to *n*-C₃₀, with maxima at *n*-C₂₆ and elevated CPI_{N-OH} values (4.5 ± 0.8) (Table 3). $\sum\text{TerN-OH}$ range from 13.4 to 105 $\mu\text{g g OC}^{-1}$, with an average of 40.4 $\mu\text{g g OC}^{-1}$, displaying similar distribution with $\sum\text{TerNA}$ (Fig. 4c and Table 3). The [NA]/[N-OH] ratios for the analysed sediments range between 2.9 and 6.9, with an average of 4.3 ± 0.9 (Table 3).

Long-chain di- and tri-unsaturated C₃₇ and C₃₈ methyl ketones and C₃₈ ethyl ketones, commonly referred to as long-chain alkenones, are present in all samples with total concentrations ranging from 3.41 to 30.5 $\mu\text{g g OC}^{-1}$, 13.0 $\mu\text{g g OC}^{-1}$ on average. The major C₂₇–C₃₀ sterols considered in this study, i.e. cholesterol (cholest-5-en-3 β -ol; 27 Δ^5), brassicasterol (24-methylcholesta-5,22-dien-3 β -ol; 28 $\Delta^{5,22}$), β -sitosterol (24-ethylcholesta-5-en-3 β -ol; 29 Δ^5) and dinosterol (4 α ,23,24-trimethyl-5 α (H)-cholest-22(E)-en-3 β -ol -30 Δ^{22}), have total concentrations ranging between 10.3 and 62.4 $\mu\text{g g OC}^{-1}$, averaging 31.7 $\mu\text{g g OC}^{-1}$. Long-chain C₃₀ *n*-alkan-1,15-diols and the corresponding C₃₀ keto-ols are also found in concentrations ranging from 7.30 to 35.8 $\mu\text{g g OC}^{-1}$, with an average of 20.3 $\mu\text{g g OC}^{-1}$ (Table 3).

$\sum\text{Mar}$ range between 18.2 and 72.6 $\mu\text{g g OC}^{-1}$, 43.6 $\mu\text{g g OC}^{-1}$ on average (Fig. 4d and Table 3), displaying a generally increasing eastward trend with maximum concentrations (> 55 $\mu\text{g g OC}^{-1}$) recorded in the deep north-

Table 3. Concentrations (OC-normalized) and indices of the considered lipid biomarkers.

Station	$\sum \text{TerNA}^a$ ($\mu\text{g g}^{-1}$ OC)	$\sum \text{TerN-OH}$ ($\mu\text{g g}^{-1}$ OC)	$\sum \text{Mar}$ ($\mu\text{g g}^{-1}$ OC)	$\sum \text{Alkenones}^b$ ($\mu\text{g g}^{-1}$ OC)	$\sum \text{C}_{30}$ diols & keto-ols ^c ($\mu\text{g g}^{-1}$ OC)	$\sum \text{Sterols}^d$ ($\mu\text{g g}^{-1}$ OC)	$27\Delta^5$ ($\mu\text{g g}^{-1}$ OC)	$28\Delta^{5,22}$ ($\mu\text{g g}^{-1}$ OC)	$29\Delta^5$ ($\mu\text{g g}^{-1}$ OC)	$30\Delta^{22}$ ($\mu\text{g g}^{-1}$ OC)	CPI_{NA}^e	$\text{CPI}_{\text{N-OH}}$	$[\text{NAI}/\text{N-OH}]$
Ionian Sea													
North													
H12	54.9	19.0	29.4	6.75	14.7	62.4	29.7	3.22	24.8	4.72	4.14	4.00	2.90
H07	40.8	13.4	34.5	5.97	14.6	35.6	6.77	5.52	14.9	8.37	4.39	4.14	3.03
Central													
BPF27													
BPF13													
BPF15													
H04	169	52.1	54.1	12.3	21.5	59.6	10.6	8.57	28.6	11.7	4.04	5.95	3.24
West													
H02	176	35.2	34.2	8.14	19.7	17.1	3.67	2.36	7.06	3.96	3.57	3.73	5.00
H05	211	42.8	47.2	12.5	25.0	24.2	4.12	3.43	10.4	6.28	4.07	3.63	4.93
H03	214	37.0	35.7	8.77	17.4	25.2	6.04	3.65	9.62	5.92	2.80	4.08	5.78
S Aegean Sea (Cretean Sea)													
Red5	206	50.7	45.3	20.1	15.9	40.1	10.8	4.74	20.0	4.51	5.07	4.89	4.06
Red9	101	14.7	22.2	9.85	7.30	17.7	3.77	2.66	8.83	2.42	7.25	5.12	6.88
Red4	196	48.4	41.0	17.4	14.8	38.6	11.5	4.63	18.3	4.15	5.36	5.21	4.05
Red8	127	29.4	35.5	16.8	12.3	26.3	5.62	3.23	14.2	3.20	6.40	4.97	4.32
W Cretean Straits													
H01	127	26.2	18.2	3.41	11.3	10.3	2.22	1.35	4.55	2.22	6.64	6.13	4.84
Red3	120	30.4	70.1	21.2	28.1	34.4	7.85	9.33	5.79	11.4	2.90	4.27	3.93
Red3.1	218	50.0	29.0	10.8	11.8	20.7	3.87	3.15	10.4	3.30	3.63	4.02	4.37
Red7	184	43.7	59.2	30.5	19.4	28.6	6.17	3.61	13.1	5.69	3.99	4.63	4.21
NW Levantine Sea													
Teraeetra Basin													
Red13	156	36.8	38.8	14.1	16.1	34.5	10.2	4.97	15.8	3.60	7.95	5.82	4.24
BPF19													
BPF22													
Red15.1	126	28.3	32.6	11.5	14.1	21.3	4.20	3.21	10.1	3.88	5.26	5.03	4.44
Red1.1	166	38.6	36.9	14.1	14.7	23.9	5.14	3.67	10.7	4.39	6.07	4.79	4.30
Te01	206	44.4	59.0	11.1	35.0	29.8	4.94	4.69	11.9	8.30	6.37	3.68	4.63
Open Sea													
Rho02	166	49.9	51.2	5.92	33.2	30.2	5.56	4.90	12.6	7.17	6.88	4.68	3.33
Her01	195	52.2	66.3	14.2	35.8	46.5	11.3	7.59	18.9	8.71	4.43	3.70	3.74
Red2													
Red2.1	483	105	72.6	23.6	35.5	47.8	7.86	6.54	26.4	6.97	1.93	4.23	4.59
BPF4													
Her03	157	39.7	45.7	7.95	29.5	22.2	4.57	3.05	9.39	5.23	3.69	3.49	3.96

Empty cells are not determined; ^a reported by Parinos et al. (2013); ^b sum of the concentrations of C37:3M, C37:2M, C36:2PFAME, C38:3Et, C38:3M, C38:2Et, C38:2M (the corresponding unsaturated homologues are indicated with the number of their carbon atoms (*n*) and the number of double bonds (*x*) – C_n:_x); M: methyl; Et: Ethyl; FAME: Fatty acid methyl ester; ^c sum of the concentrations of long-chain C₃₀-*n*-alken-1,15-diol and C₃₀ keto-ols; ^d sum of the major C₂₇–C₃₀ sterols considered in this study, i.e. cholesterol (cholest-5-en-3 β -ol, 27 Δ^5), brassicasterol (24-methylcholest-5,22-dien-3 β -ol, 28 $\Delta^5,22$), β -sitosterol (24-ethylcholest-5-en-3 β -ol, 29 Δ^5) and dinosterol (4 $\alpha,23,24$ -trimethyl-5 α (H)-cholest-22(E)-en-3 β -ol, 30 Δ^{22}).

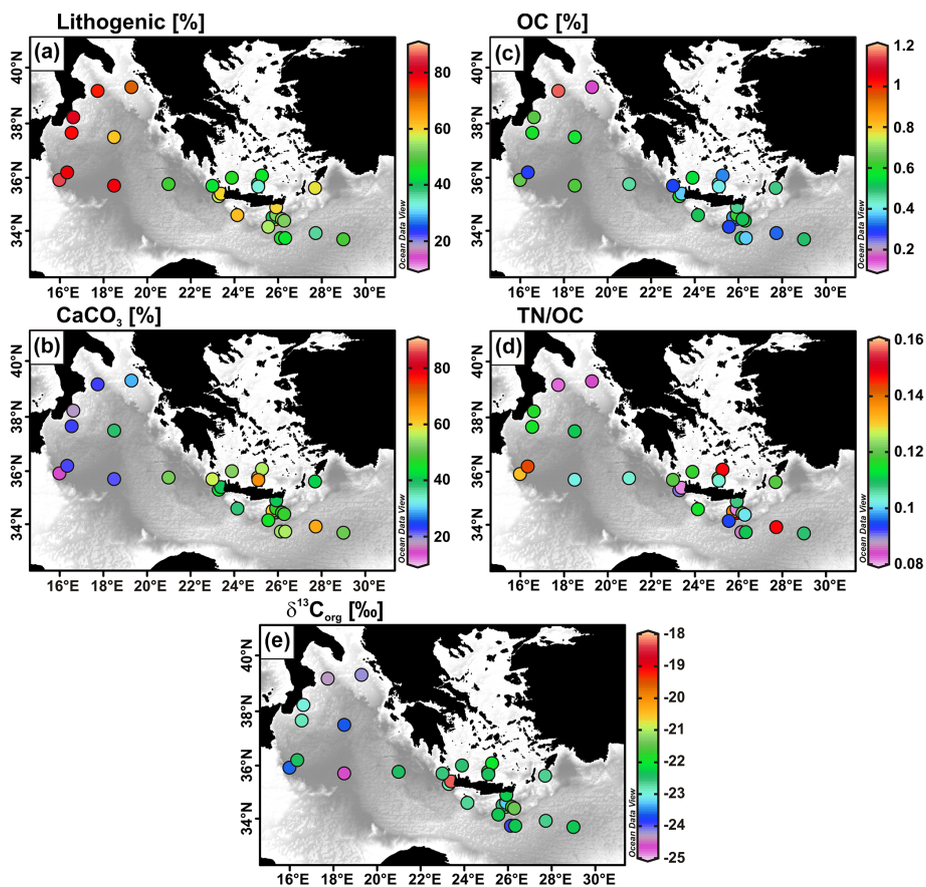


Figure 3. Spatial distribution of (a) lithogenics, (b) CaCO₃ contents, (c) OC contents, (d) molar TN / OC ratios and (e) $\delta^{13}\text{C}$ values in surface sediments of the deep eastern Mediterranean Sea.

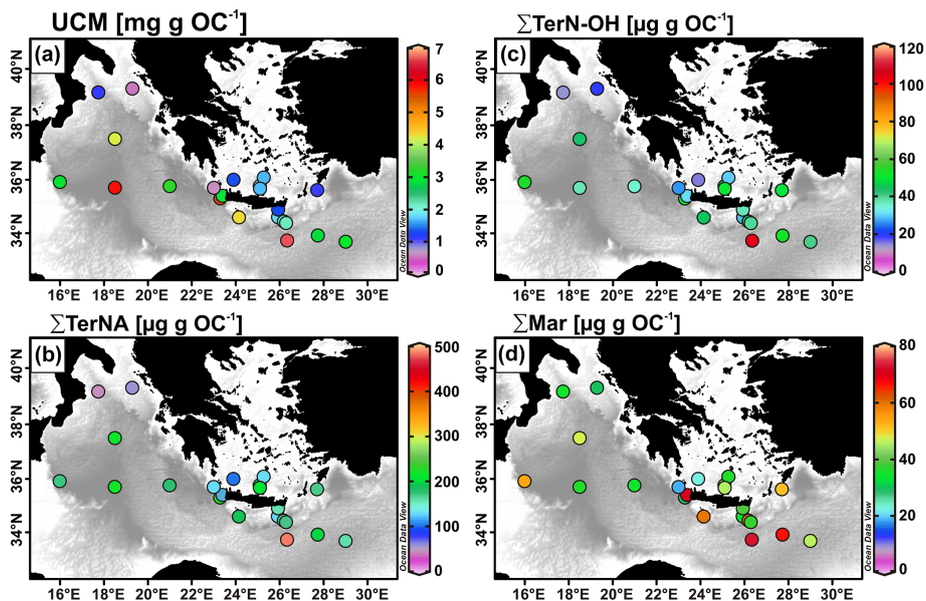


Figure 4. Spatial distributions of the OC-normalized concentrations of (a) unresolved complex mixture (UCM), (b) ΣTerNA , (c) $\Sigma\text{TerN-OH}$ and (d) ΣMar in surface sediments of the deep eastern Mediterranean Sea. Abbreviations of lipid biomarkers are defined in the text.

western Levantine Sea (stations Red2.1, Her01, Ier01 and Red7). Elevated ΣMar values are also recorded at stations Red3 in the upper slope of the western Cretan Straits as well as stations H04 and H05 in the deep Ionian Sea. The lowest values ($< 40 \mu\text{g g OC}^{-1}$) are obtained in the southern Aegean Sea (stations Red8, Red9), the northern Ionian Sea (station H12) and the western Cretan Straits (station H01).

4.5 Multivariate analysis of geochemical parameters

Three main principal components (PCs) are identified from PCA, accounting for 64.3 % of the variation within the data set (23.8, 22.8 and 17.7 % for PC1, PC2 and PC3 respectively). PC1 is characterized by positive loadings for water depth, ΣTerNA , $\Sigma\text{TerN-OH}$, ΣMar , UCM and negative loadings for CPI_{NA} and $\text{CPI}_{\text{N-OH}}$. The highest positive loadings on PC2 are associated to % CaCO_3 , % clay in the bulk sediment and TN / OC values, while negative loadings are associated to % lithogenics. Finally, the geochemical parameters with high positive loadings on PC3 are % OC, % TN and % clay of the lithogenic fraction, while those with negative loadings are $\delta^{13}\text{C}$ and sorting of bulk sediment (Fig. 5a).

Factor scores on each PC display significant variability amongst the studied stations, both within the same area and from one area to another (Fig. 5b). High positive factor scores on PC1 are observed both in stations to the west (Ionian Sea) and east (western Cretan Straits and northwestern Levantine Sea). For PC2, an eastward increasing contribution of positive factor score values seems to exist, with the highest ones located in the southern Aegean Sea and the northwestern Levantine Sea. In contrast, the prevalence on PC3 is recorded in stations of the Ionian Sea and in parts of the northwestern Levantine Sea (Ierapetra Basin).

The contents of CaCO_3 show an increasing north–south and west–east gradient (Figs. 3b and 5). In the southern Aegean Sea, the northwestern Levantine Sea and the western Cretan Straits' stations, CaCO_3 contents are positively correlated to % clay of the bulk sediment ($r = 0.48$, $p < 0.05$) and to alkenone concentrations ($r = 0.62$, $p < 0.05$). Lithogenic contents are higher in the north and west (Ionian Sea) while being significantly positively correlated to OC and TN contents ($r = 0.65$ and $r = 0.72$, $p < 0.05$ respectively excluding stations BF15, H07 and H12 of the Ionian Sea). Furthermore, OC and TN contents of stations deeper than 2100 m show a significant positive correlation to water depth ($r = 0.54$ and 0.70 respectively $p < 0.05$). However, this is not highlighted by the PCA. A significant positive correlation is also observed for OC and TN contents in the analysed samples ($r = 0.87$, $p < 0.0001$) (Fig. 6a).

Surface sediments of the Ionian Sea show a significant ($p < 0.05$) positive correlation of OC and TN contents to % clay ($r = 0.80$ and $r = 0.73$) and a negative correlation to % silt ($r = -0.75$ and -0.65) and D_{50} of the lithogenic fraction ($r = -0.79$ and -0.83). $\delta^{13}\text{C}$ values (excluding station Red3 of the western Cretan Straits) are significantly ($p < 0.05$) and

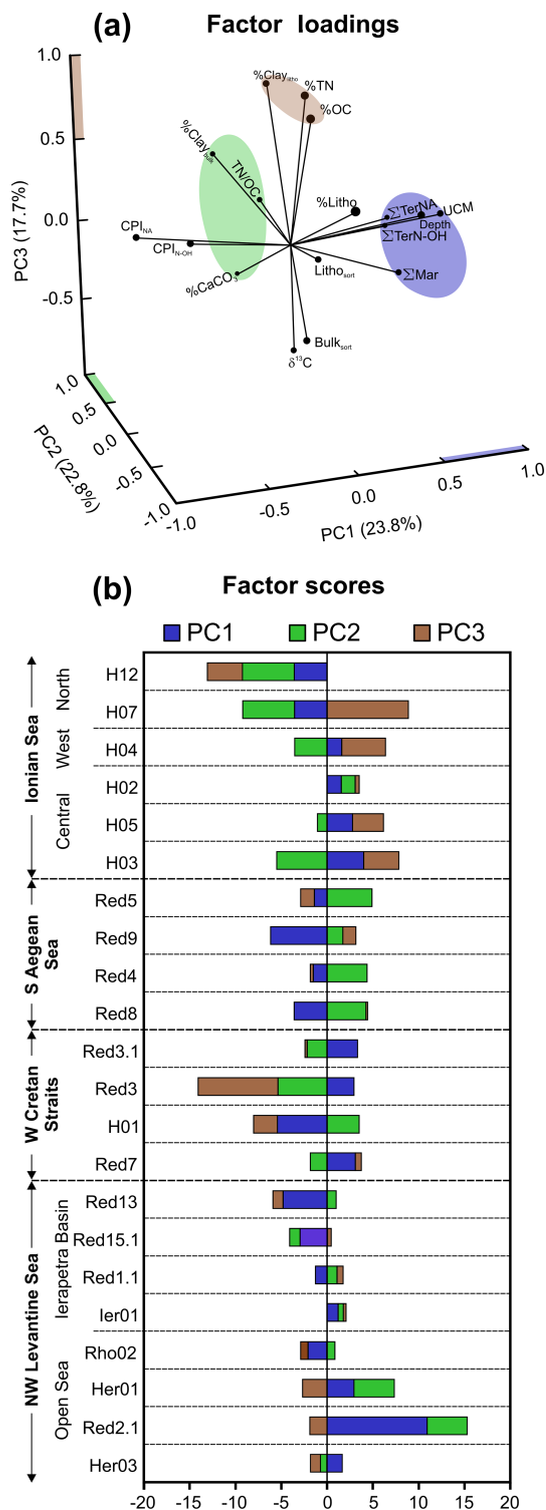


Figure 5. (a) Scatter plot of the factor loadings of the three principal components obtained in the principal component analysis; (b) plot of the scores found at each station. Abbreviations of lipid biomarkers are defined in the text.

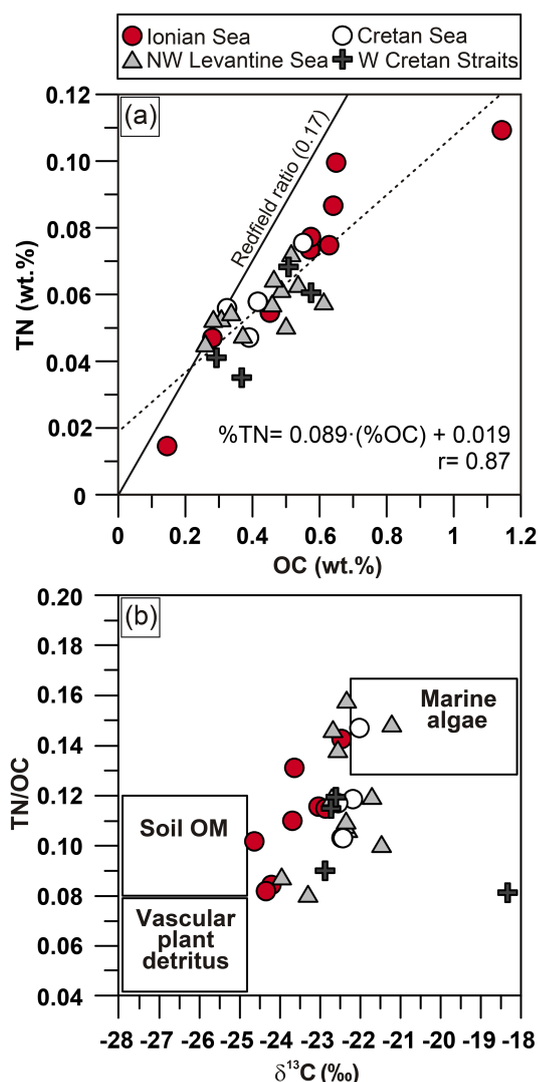


Figure 6. (a) Co-plot of the weight percent content of nitrogen (TN) vs. organic carbon (OC) for the deep-sea surface sediments analysed. The linear fit of the data is shown (dotted line) along with the Redfield ratio N_{16}/C_{106} associated with the fresh marine phytoplankton (0.17 wt/wt). (b) Plot of molar TN/OC ratios vs. $\delta^{13}C$, in surface sediments of the deep eastern Mediterranean Sea. The compositional ranges of organic matter sources illustrated in plots (boxes) derive from previously published studies (see Sect. 5.1).

positively correlated to $CaCO_3$ contents ($r = 0.53$), % clay of the bulk sediment ($r = 0.65$) and molar TN/OC ratios ($r = 0.53$) and negatively correlated to % OC ($r = -0.46$), % silt ($r = -0.66$) and D_{50} ($r = -0.59$) of the bulk sediment.

Terrestrial lipid biomarkers concentrations ($\sum TerNA$ and $\sum TerN-OH$) display a significant positive correlation amongst them ($r = 0.95$, $p < 0.0001$), as well as to % clay (both with $r = -0.55$, $p < 0.05$) and D_{50} ($r = 0.62$ and 0.58 respectively $p < 0.01$) of the bulk lithogenic fraction. Moreover, $\sum TerNA$ and $\sum TerN-OH$ show a significant positive correlation ($p < 0.05$) to β -Sitosterol ($_{29}\Delta^5$) ($r = 0.71$ and

0.56 respectively excluding stations H07 and H12 of the Ionian Sea). $\sum TerNA$ and $\sum TerN-OH$ (not normalized to OC contents) are also significantly correlated to OC ($r = 0.81$ and 0.76 respectively $p < 0.001$), again excluding stations H07 and H12 of the Ionian Sea. $\sum Mar$ (not normalized to OC contents) displays a significant positive correlation with OC ($r = 0.7$, $p < 0.001$), while a significant positive correlation ($r > 0.65$, $p < 0.005$ in all cases) is evident amongst the concentration of cholesterol ($_{27}\Delta^5$) and marine algal markers ($_{28}\Delta^{5,22E}$, $_{30}\Delta^{22E}$, C_{30} diols&keto-ols and alkenones; see Sect. 5.2.1).

5 Discussion

5.1 Sources of sedimentary material in the deep eastern Mediterranean Sea

Clearly, the surface sediments of the deep EMS mostly consist of lithogenics and carbonates, with low OC contents while opal is nearly absent (Table 2). The range of lithogenics, carbonates and opal contents recorded in the investigated samples are similar to those previously reported for the eastern Mediterranean Sea (Emelyanov and Shimkus, 1986; Bethoux, 1989; Cros, 1995; Kemp et al., 1999; Rutten et al., 2000; Struck et al., 2001). OC contents reach values slightly above 1 % and are also comparable to those found in the eastern Mediterranean Sea (0.56–1.51 %, Danovaro et al., 1993; 0.23–0.99 %, Bianchi et al., 2003; 0.30–0.82 %, Gogou et al., 2000; 0.25–1.73 %, Polymenakou et al., 2006) and relatively lower than those found in the western Mediterranean Sea (0.80–1.60 %, Kaiser et al., 2014; 0.47–1.53 %, Masqué et al., 2003; 0.23–1.85 %, Roussiez et al., 2006). Values found are comparable to the typical hemipelagic sediments found in continental slopes (Rullkötter, 2006) and slightly higher than those in deep basin areas (Seiter et al., 2004).

5.1.1 Lithogenics and carbonate

The grain size of the lithogenic fraction found in the studied sediments is very similar to that of Saharan dust particles, which mainly consist of clayey silts and silty clays with diameters ranging from 0.5 to 60 μm ($D_{50} \sim 5 \mu m$) and two main modes at 3–4 and 60 μm (Ratmeyer et al., 1999; Guerzoni and Molinaroli, 2005 and references therein). This strongly suggests that the Sahara is the main source of lithogenics to the deep EMS. This is further supported by observations pointing to Saharan dust transported by southerlies blowing over the great North African desert as the main lithogenic input to the EMS (Guerzoni et al., 1999; Weldeab et al., 2002). The Saharan dust spreads rather uniformly across the EMS (Rutten et al., 2000; Jickells et al., 2005). Furthermore, to a lesser extent volcanic ash deposition into the EMS represents another external source of fine-grained particles (< 5–50 μm) (Kelepertsis et al., 2003). Mount Etna, located on the island of Sicily, generates volcanic ash plumes

that are transported by the wind reaching as far as Greece and Libya (Olgun et al., 2013 and references therein).

Riverine inputs have a rather minor influence onto the open EMS sedimentation as they are small and localized (Weldeab et al., 2002; Statham and Hart, 2005). The relatively higher lithogenic contents found in most of the Ionian Sea stations (Fig. 3a and Table 2) points to fluvial inputs reaching the area from the Adriatic Sea. The main source of riverine inputs is the Po River, opening into the northernmost end of the Adriatic Sea, although inputs from smaller rivers draining the Apennines could be also relevant (Weldeab et al., 2002). In the Ionian Sea, river-sourced particles are carried by both surface and deep currents flowing southwards along the Italian Peninsula as part of the overall anticlockwise circulation in the Adriatic Sea (Orlic et al., 1992). It should be noted that dense water formation takes place seasonally in the Adriatic Sea, which triggers episodes of fast-flowing, sediment-loaded dense near-bottom currents that cascade into the deeper Meso Adriatic depression before passing through the Otranto Strait, subsequently spreading into the Ionian Sea where their particle load settles to the bottom (e.g. Zoccolotti and Salusti, 1987; Manca et al., 2002; Canals et al., 2009).

The grain size variability of the carbonate particles recorded in the studied sediment samples is indicative of calcareous skeletons of primary producers. While the abundance of particles $< 8 \mu\text{m}$ is attributable to coccoliths, which is the most abundant primary producer in the EMS (Emelyanov and Shimkus, 1986), coarser carbonate particles mostly correspond to shells and fragments of calcareous dinoflagellates and planktonic foraminifers, in agreement with Ziveri et al. (2000) and Frenz et al. (2005). Although part of the carbonate fraction might also have a terrestrial provenance as transported, for instance, with Saharan dust (Chester et al., 1977; Correggiari et al., 1989; Rutten et al., 2000), this does not seem to be the case with the investigated samples. Like other aeolian particles, aeolian carbonates are typically better sorted than those formed in situ, as shown by a well-sorted unimodal distribution due to gravitational settling during atmospheric transport (Skonieczny et al., 2013). The predominance of very poorly sorted grain size distributions within the bulk sediment samples (Fig. 2a) and the highly variable CaCO_3 contents (Fig. 3b) in the investigated samples suggests that even within such a highly oligotrophic environment biogenic carbonates are the main source of CaCO_3 in the deep EMS.

5.1.2 Sources of sedimentary organic matter

Bulk geochemical proxies such as elemental (TN / OC) and stable isotopic ratios of OC ($\delta^{13}\text{C}$) have been widely used to assess the sources of OM in marine sediments (Meyers, 1994; Goñi et al., 2003; Hu et al., 2006). Sedimentary molar TN / OC ratios and $\delta^{13}\text{C}$ values determined in this study are consistent with values previously reported for surface sedi-

ments of the deep EMS (Tesi et al., 2007b; Meyers and Arnaboldi, 2008; Carlier et al., 2010; Goudeau et al., 2013).

Marine-derived OM is characterized by high TN contents yielding TN / OC ratio values > 0.12 , while vascular plants are N-depleted yielding TN / OC ratio values < 0.08 (Redfield et al., 1963; Hedges and Oades, 1997). $\delta^{13}\text{C}$ values in marine algae from low- to mid-latitude temperate seas vary from -18 to -22‰ (Goericke and Fry, 1994; Meyers, 1994; Harmelin-Vivien et al., 2008), whereas most terrestrial OM inputs from C3 plants show depleted $\delta^{13}\text{C}$ values ranging from -25 to -28‰ (Hedges et al., 1997).

In order to constrain the origin of sedimentary OM and assess the spatial variability in its marine-to-terrestrial blend molar TN / OC ratios were plotted against $\delta^{13}\text{C}$ values. Plots show that the elemental and isotopic composition of sedimentary OM fall out of the typical compositional ranges of the potential sources (Fig. 6b). Excluding station Red3, located in the upper slope of western Cretan Straits, an overall positive relationship for molar TN / OC ratios and $\delta^{13}\text{C}$ values ($r = 0.53$, $p < 0.05$) becomes apparent (Fig. 6b), thus indicating that the composition of the OM in the studied sediment samples could be explained as a mixture of terrigenous (low TN / OC and $\delta^{13}\text{C}$) and marine-derived (high TN / OC and $\delta^{13}\text{C}$) materials.

Additionally, the relative contribution of the marine vs. terrestrial sources of OC over the study area has been evaluated by means of a simple $\delta^{13}\text{C}$ -based binary mixing model (Table 2), where a marine $\delta^{13}\text{C}$ value of -20.4‰ and a terrestrial $\delta^{13}\text{C}$ value of -27.0‰ (Tesi et al., 2007a) are assumed, considering that the contribution of C4 vascular plants ($\delta^{13}\text{C}$ from -12 to -15‰) can be considered negligible throughout the study area (e.g. Collatz et al., 1998).

As evident in Table 2, sediments from the Ionian Sea are characterized by elevated contributions of terrestrial OC (OC_{terr}), reaching up to 64.2%. In contrast, W Cretan Straits, Cretan Sea and NW Levantine Sea stations show low OC_{terr} contributions and marine OC (OC_{mar}) peaks of 87.7% (see also Sect. 5.2). The elevated OC_{terr} values recorded in the Ionian Sea suggest that terrigenous OM entering the Adriatic Sea escapes towards the deep Ionian basins. Indeed, during energetic dense shelf water cascading events, lateral flux prevails over the vertical flux in the southern Adriatic Sea and the contribution of soil-derived OC increases up to $\sim 60 \%$ (Turchetto et al., 2007; Tesi et al., 2008). Despite the relatively high OC_{terr} found preserved in the deep Ionian Sea, values are lower than those recorded in areas impacted by extreme events such as storms that enhance the export of organic matter from the shelf to the deep environment, for example in the western Mediterranean (up to 78% of OC_{terr}) (Pedrosa-Pàmies et al., 2013) and the East China Sea (up to 90%) (Selvaraj et al., 2015).

Lipid biomarkers have often been used as molecular proxies to identify specific biological precursors of sedimentary OM (Meyers, 1997; Volkman, 2006). The concentrations of the sedimentary lipid biomarkers determined in this study are

fairly comparable to those previously reported in areas devoid of significant fluvial influence both in the eastern and western Mediterranean basins (Grimalt and Albaigés, 1990; Gogou et al., 2000; Gogou and Stephanou, 2004; Kaiser et al., 2014).

The patterns of long-chain *n*-alkanes and *n*-alkanols with elevated CPI_{NA} and CPI_{N-OH} values (Sect. 4.4; Table 3) respectively indicate the presence of allochthonous natural (terrestrial) inputs from epicuticular higher plant waxes (Eglinton and Hamilton, 1967). Saharan dust is probably the main vector for the transport of small charcoal-like fragments of burnt vegetation, leaf wax-derived lipids adsorbed on clays and cuticular fragments into the open EMS given the relatively minor direct influence of riverine inputs (Gogou et al., 1996; Eglinton et al., 2002).

Lipid biomarkers preserved in the surface sediments of the study area also highlight the contribution from autochthonous marine OM derived from in situ phytoplankton production. More specifically, the abundance of brassicasterol ($_{28}\Delta^{5,22E}$) reveals the presence of diatoms and prymnesiophytes, while dinosterol ($_{30}\Delta^{22E}$) is a major compound in dinoflagellates (Volkman, 1986). The presence of long-chain alkenones reflects the productivity from algal species of the *Prymnesiophyte* class, e.g. *Emiliania huxleyi* (Marlowe et al., 1984), which constitutes the dominant primary producers across the Mediterranean Sea (Ziveri et al., 2000; Triantaphyllou, 2004). Regarding the long-chain C_{30} *n*-alkan-1,15-diols and the corresponding C_{30} keto-ols, although their major sources remain unknown, microalgae of the genus *Nannochloropsis* (class *Eustigmatophyceae*) are potential sources, while C_{30} keto-ols might result from oxidation of the corresponding C_{30} diols (Volkman, 1986; Volkman et al., 1999; Rampen et al., 2012).

In addition, while the abundance of cholesterol ($_{27}\Delta^5$) highlights the existence of marine consumer organisms such as zooplankton and benthic animals (Grice et al., 1998), β -sitosterol ($_{29}\Delta^5$) may derive from both terrigenous and marine sources (Volkman, 1986). In the study area, however, the positive correlation between $\sum TerNA$ (and $\sum TerN-OH$) to β -sitosterol argues for a dominant terrestrial origin for this compound.

Aside from natural sources, the abundance of UCM indicates a contribution of anthropogenic OM resulting in chronic oil pollution of the investigated sediments (Parinos et al., 2013). UCM levels recorded in the deep EMS are comparable to those reported for surface sediments in unpolluted coastal and/or open-sea areas and are at least one order of magnitude lower than those reported for coastal areas subjected to enhanced anthropogenic inputs (Gogou et al., 2000; Parinos et al., 2013; Kaiser et al., 2014; Romero et al., 2015; and references therein). Two main pathways have been identified for the introduction of petroleum hydrocarbons into the deep EMS, which are direct discharges from merchant shipping and oil transportation (UNEP, 2010) and atmospheric

transport and deposition (Gogou et al., 1996; Castro-Jiménez et al., 2012; Parinos et al., 2013).

5.2 Regional variability and oceanographic control on the geochemical composition of deep eastern Mediterranean Sea surface sediments

The PCA provides a robust overview of the variables and processes controlling the geochemical composition of the investigated deep-sea surface sediments (Fig. 5).

The significant positive loadings of $\sum TerNA$, $\sum TerN-OH$, $\sum Mar$, UCM and depth on PC1 are indicative of a considerable contribution from both natural (marine and terrestrial) and anthropogenic (degraded petroleum products) OM preserved in deep-sea surface sediments of the EMS. Moreover, the negative PC1 loadings of CPI_{NA} and CPI_{N-OH} ratios vs. depth indicate that the terrestrial OM is relatively altered with increasing water column depth. Although the negative PC1 loading of CPI_{NA} ratio could be also indicative of an enhanced contribution from non-degraded petroleum inputs, the patterns of aliphatic hydrocarbons for the investigated sediment samples indicate no important bias associated to non-degraded petroleum products on CPI_{NA} ratio values (Parinos et al., 2013). Overall, PC1 represents the degradation processes and fate of the sedimentary OM in the study area.

The second PC separates samples with high carbonate contents, molar TN / OC ratios and enhanced contribution of clay-sized particles from those with high lithogenic contents. Therefore, samples with positive loadings on PC2 are enriched in fine marine carbonate particles, while those with negative loadings are enriched in lithogenic particles (see Sect. 5.2.2).

Finally, PC3 separates samples with high contents of OC, TN and clays from those with high values of $\delta^{13}C$. Consequently, positive loadings on PC3 are associated to sediments with an enhanced contribution of OC-rich fine particles, thus pointing to hydrodynamic processes that control grain size sorting and remobilization/deposition of sedimentary material with different OC contents (see Sect. 5.2.2).

5.2.1 Processes modulating the biogeochemical signal of the sedimentary organic matter

The low OC and TN contents in the surface sediments of the study area reflect the oligotrophic character of the EMS (e.g. Krom et al., 2003). In the studied sediments some processes may have further pushed TN to OC ratios towards low values (Fig. 6a). These include the preferential degradation of N-rich proteinaceous components of algal OM during early diagenesis (Meyers et al., 1996; Meyers, 1997; Hopmans et al., 2004) and the enrichment of OC relative to TN due to the input of petroleum residues (Friligos et al., 1998). Furthermore, a significant contribution of inorganic N, presumably as NH_4^+ adsorbed on clays (Müller, 1977; Meyers, 1997), is

inferred from the positive intercept on the N axis at around 0.02 % (Fig. 6a).

Although isotopic fractionation specifically associated with early diagenesis is negligible and the isotopic composition of sedimentary OM is fairly conservative (e.g. Di Leonardo et al., 2009 and references therein), $\delta^{13}\text{C}$ values can be potentially shifted by microbial rearrangements (Lehmann et al., 2002) and inputs of anthropogenic OM. $\delta^{13}\text{C}$ values for crude oil and petroleum products are in the order of -28.5 and -28.9% respectively (Rumolo et al., 2011 and references therein). In the study area, the positive PC1 loadings of depth and UCM indicate an enhanced contribution of degraded petroleum products with increasing water depth. This points to an enrichment of the sediments in degraded petroleum hydrocarbons in the deep Ionian Sea and western Cretan Straits' stations H03 and Red3.1 where such an isotopic shift is observed (Figs. 3e and 7), in addition to maximum concentrations of UCM along with relatively low molar TN / OC ratios (< 0.11) (Table 2, Figs. 3d and 4a).

Lipid biomarkers provide further information on the natural sources of sedimentary OM. The significant positive correlation of $\sum\text{TerNA}$ and $\sum\text{TerN-OH}$ concentrations (not normalized to OC) to OC contents suggests a close association of terrestrial OM to OC, while the significant positive correlation of $\sum\text{TerNA}$ and $\sum\text{TerN-OH}$ to % clay suggests that the transport and accumulation of terrestrial OM is associated to fine particles (see Sect. 5.2.2). The relatively uniform spatial distribution of CPI_{NA} and $\text{CPI}_{\text{N-OH}}$ ratio values (Fig. 8) together with the negative PC1 loadings of CPI_{NA} and $\text{CPI}_{\text{N-OH}}$ ratios vs. depth (Fig. 5), are overall indicative of the reworking of the terrestrial OM accumulated in deep EMS basins. Moreover, the elevated values of the $[\text{NA}] / [\text{N-OH}]$ ratio in the study area (Fig. 8; Table 3) are indicative of the enhanced degradation of $\sum\text{TerN-OH}$ relatively to $\sum\text{TerNA}$. The above are rather consistent with the long-range atmospheric transport and long residence time in the water column and into the sediments of the terrestrial OM and the fact that $\sum\text{TerNA}$ are more resistant to degradation than their alcohol counterparts (Gogou and Stephanou, 2004). Moreover, the relatively elevated retention time of terrestrial OC in the inner shelf of the Adriatic Sea, before being conveyed to the Ionian Sea by dense shelf water cascading events, allows for its significant microbial degradation or marine dilution (Tesi et al., 2008).

While a significant positive correlation is observed for $\sum\text{Mar}$ concentrations (not normalized to OC contents) and OC, indicating that the latter exerts an important control on the distribution of algal markers' concentrations in the study area, no significant correlation is observed between $\sum\text{Mar}$ and grain size. These correlations together with the deviation trend observed for molar TN / OC ratios from the classical Redfield ratio (16/106) with increasing water depth, which is more evident for the deep Ionian Sea stations (Fig. 6a), probably reflect the preferential degradation processes during the transport and deposition of marine labile sedimentary

OM, which probably also masks the association of marine OM to fine particles. The observations above, jointly with the presence of cholesterol ($_{27}\Delta^5$) and its significant positive correlation with the concentrations of marine algal markers, are altogether indicative of the re-working of algal OM by zooplankton and benthic invertebrates (Volkman et al., 1990; Gogou and Stephanou, 2004).

5.2.2 Sediment transport and deposition processes

The second and third PCs of the PCA highlight the main processes that affect sediment dispersal and deposition in the study area. These relate to pelagic settling of marine skeletons from surface waters (corresponding to PC2) and hydrodynamic sorting of organic-rich fine sediment by bottom currents (corresponding to PC3).

Particulate matter exported from the upper layers of the water column in the EMS is primarily composed of biogenic particles and atmospheric dust, which while settling to the seabed are able to transfer OC, other nutrient elements and OC-associated organic pollutants (e.g. Stavrakakis et al., 2000, 2013; Theodosi et al., 2013). In the deep EMS, the distribution of pelagic carbonates (second PC) seems to be mainly influenced by planktonic contributions. In the study area, the phytoplankton biomass and primary production are relatively higher in regions of cyclonic water mass circulation. The Rhodes cold-core gyre, situated in the southeast of the Rhodes Island (NW Levantine Sea), is the most prominent dynamic feature in the EMS and is the main source area of the LIW. In this cyclonic gyre, which is enhanced during winter period, dense water masses from deeper layers tend to upwell at its centre, feeding the upper layers with nutrient-rich waters (Salihoğlu et al., 1990). Therefore, this gyre plays an important role in the productivity of the Levantine Sea.

The third PC separated samples with high OC, TN and clay contents, which is indicative of a close OM–mineral association. This is in agreement with the high OC contents found in the fine-grained sediment samples from the deeper stations representing an essentially quiet environment (Figs. 2, 3 and 7). This is in contrast to the lower OC contents observed in coarser samples ($D_{50} > 14\ \mu\text{m}$) from shallower depths where currents up to $20\ \text{cm s}^{-1}$ occur commonly (Kontoyiannis et al., 2005; Ursella et al., 2014). Fine-grained particles have high capacity for OM adsorption due to their large specific surface area and thus enhanced OM contents relative to coarse-grained particles (Mayer, 1994; Hedges and Keil, 1995). Physical processes such as hydrodynamic sorting remobilize and transport sedimentary material with different OC contents, with the OC-rich finest ones easily reaching the deep EMS. A similar situation has been reported in active sedimentary environments such as the submarine canyons in the western Mediterranean Sea (Pedrosa-Pàmies et al., 2013), the northwestern Gulf of Mexico (Goñi et al., 1998) and the Peru Margin (Bergamaschi et al., 1997). However, none of these studies reported such OM–mineral

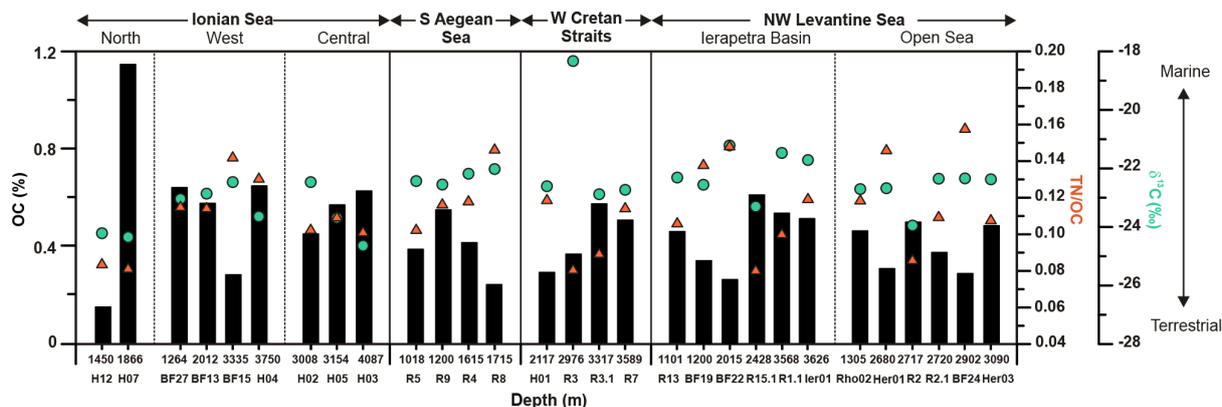


Figure 7. Distribution of OC content and bulk sediment tracers (TN / OC and $\delta^{13}\text{C}$) in surface sediments of the deep eastern Mediterranean Sea. The histogram reflects a spatially variable mixture of marine and terrestrial sources for sedimentary OM. Orange triangles and green circles correspond to molar TN / OC ratios and $\delta^{13}\text{C}$ values respectively.

associations at depths beyond 4000 m. This indicates that this organic–mineral interaction is maintained from the shallow to high depths, which constitute the final sedimentary sink. In short, the deep EMS behaves as a depocentre for OM-rich fine particles.

Moreover, in the EMS these bulk sediments show a poorer sorting than the lithogenic fraction (Fig. 2). This could be related to the presence of coarse biogenic carbonate particles in bulk samples or the effective hydrodynamic sorting linked to the prevailing depositional conditions in such deep low-energy environments (Friedman, 1969).

The poor sorting and positive skewness found in grain size types I, II, IV and V of the bulk sediment samples (Fig. 2a) in the southern Aegean Sea and the northwestern Levantine Sea are explained by the prevalence of pelagic biogenic sedimentation, as shown by the high positive score values observed on PC2 (Fig. 5b). Accordingly, high percentages of CaCO_3 have been measured in stations of the southern Aegean Sea, the northwestern Levantine Sea and western Cretan Straits, which correlate to % clay fraction of the bulk sediment samples and the concentrations of alkenones (Table 3). This links to the formation of clay–carbonate concretions that have been reported in particularly large amounts in the southern Aegean Sea (Emelyanov and Shimkus, 1986). Vertical mixing and upward transport of nutrients in the eddies and gyres, such as the Rhodes gyre, may trigger primary production and subsequently the sinking and dominance of pelagic biogenic particles over particles from other sources in the northwestern Levantine Sea (Siokou-Frangou et al., 1999). The % OC_{mar} and the $\sum\text{Mar}$ distribution are indicative in that respect as they show a general eastward increasing trend with peak concentrations in deep basins of the northwestern Levantine Sea.

Surface sediments in the Ierapetra Basin (stations Red15, Red1.1 and Ier01) also show positive scores on PC3 (Fig. 5b), which point to an influence of hydrodynamic sort-

ing processes. The relatively high OC content in these stations (Fig. 7), along with the elevated values of the associated natural and anthropogenic lipid concentrations (Table 3), suggests that the Hellenic Trench is a sink of OC associated to fine particles transferred by the active outflows of the Cretan Straits, besides the pelagic sedimentation related to the well-known semi-permanent Ierapetra anticyclone (Larnicol et al., 2002; Taupier-Letage, 2008).

Sediments with grain size types I–III of the bulk sediment samples (Fig. 2a) in the Ionian Sea show lower CaCO_3 contents due to the dilution by lithogenic components, with station H02 being the only exception. High positive score values on PC3 (Fig. 5b), and the significant positive correlation for OC and TN contents to % clay of the lithogenic fraction, further suggest a significant influence of hydrodynamic sorting processes, which largely determine a differential distribution of OM in surface sediments according to grain size. In this area, the Otranto Strait may act as a preferential conduit by funnelling fine OC-rich particles from the Adriatic Sea towards the deep basins of the adjacent Ionian Sea, which may eventually reach the Levantine Sea (Bensi et al., 2013). The enhanced contributions of OC_{terr} (Table 2) and terrestrial biomarkers concentrations (Figs. 4b–c and Table 3) of the Ionian samples off the southern mouth of the Adriatic Sea probably also reflect the preferential degradation of labile marine OM in deep Ionian Sea basins due to slower sedimentation rates and longer residence time of OM. Low OC content (Fig. 3c), poor sorting and very negatively skewed high % sand of grain size type VI (Fig. 2a) at the shallower station H12 just south of Otranto Strait support winnowing of fine OC-rich particles due to episodic events of high current speed exiting the Adriatic Sea (Bignami et al., 1991; Gacic et al., 1996; Poulos et al., 1999).

Finally, station Red3 from the upper slope of western Cretan Straits representing grain size type VII of the bulk sediment samples (Fig. 2a) shows the highest contents of sand

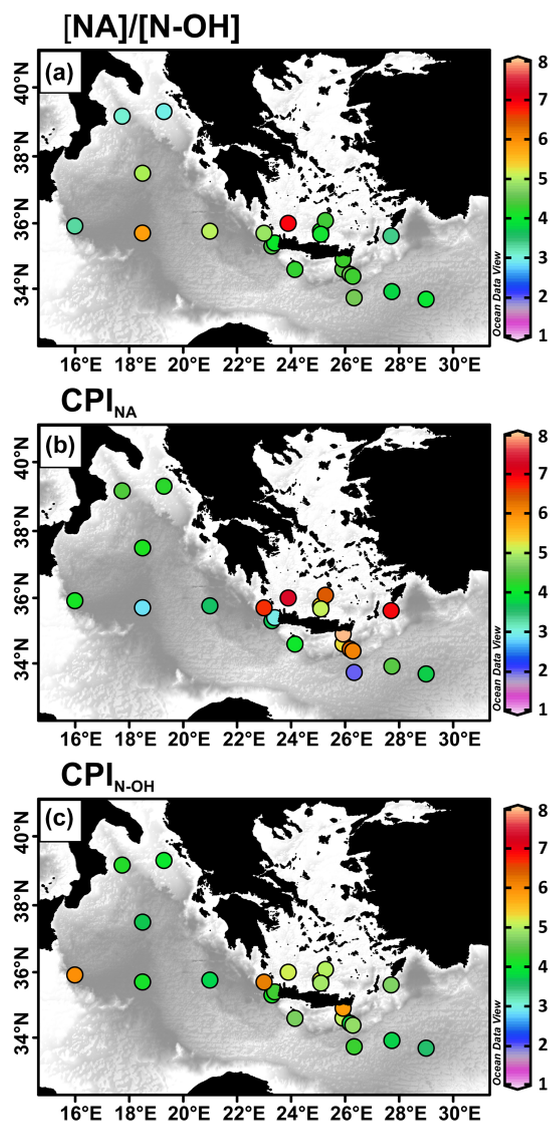


Figure 8. Spatial distribution of lipid biomarker indices (a) $[NA]/[N-OH]$, (b) CPI_{NA} and (c) CPI_{N-OH} . Abbreviations of lipid biomarker indices are defined in the text.

(47.7%), which is poorly sorted. This is in agreement with the occurrence of the topographically restricted deep outflow of the western Cretan Straits. These straits are characterized by maximum outflow speeds during winter and minimum speeds during fall (Kontoyiannis et al., 2005). The turbulent, fluctuating outflow current should normally trigger sediment resuspension and induce selective transport, thus leaving coarse OC-poor particles in the upper slope of the western Cretan Straits (negative factor scores of PC2 and PC3) and carrying fine OC-rich particles to the lower slope. A similar pattern has been also observed in other submarine canyon settings of the Mediterranean Sea, such as the Cap de Creus Canyon (Sanchez-Vidal et al., 2008) and the Blanes Canyon (Pedrosa-Pàmies et al., 2013). The top $\delta^{13}C$ values found in

the upper slope (Fig. 7 and Tables 2–3) indicates high contribution of marine OC, which, however, is not supported by the lipid biomarkers results. Winnowing of fine particles loaded with terrestrial OC, thus shifting the isotopic signal of the remaining coarse particles towards high and more marine values seems to be the most plausible explanation.

6 Conclusions

Surface sediments collected from deep basins of the oligotrophic EMS were investigated using a multi-proxy approach that involved elemental composition, grain size, stable isotopes and selected lipid biomarkers' analyses resulting in a robust database to determine sediment sources, the degradation and preservation state of OM and processes that affect sediment dispersal and deposition. The PCA analysis helped to identify the main controlling factors of the observed geochemical variability in the investigated sediments. Such factors are sediment sources in terms of allochthonous vs. autochthonous, a highly variable physiography, the thermohaline structure and the regional and local circulation, leading to hydrodynamic sorting and regulating particle settling/deposition and OM preservation state.

Surface sediments of the investigated part of the EMS mostly consist of airborne lithogenic particles and biogenic carbonate particles, the latter deriving from primary production into surface waters. Sedimentary OM appears in rather low contents (0.15–1.15 % OC), with bulk and molecular organic tracers reflecting a mixed contribution from both natural (autochthonous and allochthonous) and anthropogenic sources. Samples from locations in the Ionian Sea and the western Cretan Straits that are under the direct influence of the Adriatic dense waters outflow through the Otranto Strait and of currents exiting the southern Aegean Sea respectively are appreciably sorted. Current regime impacted not only grain size but also OC loadings within each subregion of the study area, with winnowing of fine OC-rich particles to the deepest EMS. In contrast, coarse OC-poor particles tend to occur in upper slope settings. While OC associated to fine particles was relatively non-degraded terrestrial OM, marine OM was found to be mostly degraded and reworked during transport processes and before reaching the deep seafloor.

The spatial variability in the yields of sedimentary OC and lipid biomarkers presented in this study highlights the heterogeneous nature of the particle load exported to the deep basins of the eastern Mediterranean Sea. Such variability must be taken into account during the development of quantitative carbon budgets for this area.

Acknowledgements. This research has been supported by the EU-funded project PERSEUS (GA 287600), the EU–Greek co-funded project KRIPIS (MIS 451724; NSRF) and REDECO (CTM2008-04973-E/MAR), BIOFUN (CTM2007-28739-E) and MEDECOS (ΣAE013/8-2010ΣE01380001) projects. Researchers from GRC Geociències Marines benefited from a *Grups de Recerca Consolidats* grant 2014 SGR 1068 by the *Generalitat de Catalunya* autonomous government. We sincerely thank chief scientist K. C. Emeis (R/V *Meteor*) and the officers and crews of R/V *Meteor*, R/V *Aegaeo* and R/V *Sarmiento de Gamboa* for their precious help during the cruises. Elemental and isotopic analyses have been performed at the scientific-technical services of the University of Barcelona. R. Pedrosa-Pàmies is supported by a predoctoral FPU grant and A. Sanchez-Vidal by a Ramon y Cajal contract. Juergen Mobius is acknowledged for his constructive comments on an earlier version of the manuscript. We would like to thank the editor, T. Tesi and three anonymous referees for their constructive comments that helped us significantly improve the quality of the manuscript during the revision process.

Edited by: G. Herndl

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