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Organic matter composition in the sediment of three Brazilian coastal lagoons – District of Macaé, Rio de Janeiro (Brazil)

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ABSTRACT

Freshwater lagoons comprise important coastal ecosystems and natural buffers between urbanized land areas and open ocean in the Rio de Janeiro State, Brazil. Studies of sediment and water chemistry, zooplankton and bacterial communities to assess the extent of anthropogenic disturbance are available. Here we contribute with an organic-geochemical approach supplemented by some microbiological aspects to complete the characterization of these lagoonal ecosystems. Bulk organic matter and extractable lipids (aliphatic hydrocarbons, alcohols and fatty acids, sterols) were investigated from two locations per lagoon: at the seaward site and landward ends – and at two depth intervals (0-3 and 3-6 cm) per site. Urbanized Imboacica Lagoon received increased anthropogenic input over the most recent years represented by the topmost 3 cm of sediment, whereas deeper sediment layers are less affected by human influence. Eutrophication or nutrient availability favored enhanced algal/cyanobacterial growth. In remote Cabiúnas and Comprida Lagoons pristine conditions are preserved. Organic matter from vascular plants dominates (chain length of free lipids up to C36), which is exceptionally well preserved by acidic lagoonal waters. Differentiation between landward and seaward sites in these two lagoons is less well established due to much smaller surface/volume to catchment ratios. No anthropogenic influences are yet detectable in sediments of Cabiúnas and Comprida Lagoons.

Key words: coastal lagoons, aqueous/terrigenous bioproduction, lipids, fatty acids, sterols, Brazil.

INTRODUCTION

In the densely populated Rio de Janeiro State (Brazil), freshwater lagoons comprise important coastal ecosystems, often forming natural buffer regions between urbanized land areas and the open

ocean. These lagoons are highly sensitive to anthropogenic influences due to heavy utilization for fishing, recreational activities and discharge of urban sewage. Several studies have shown the extent of anthropogenic disturbance in these natural ecosystems by means of investigating sediment and water chemistry (Furtado et al. 1997, Santos et al. 1998), as well as monitoring zooplankton (Kozłowski-Suzuki

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et al. 1998) and bacterial communities (Furtado and Esteves 1997, Biesboer et al. 1998). In addition studies in the Patos Lagoon (Brazil), the largest in South America, have demonstrated chemical processes concerning nutrients and trace metals on a larger scale but in a comparable environmental setting (Windom et al. 1999).

Organic material from autochthonous and allochthonous origin accumulates in the sediment of aquatic ecosystems. Here it supports bacterial activity that results in decomposition and nutrient recycling. By mediating Eh and pH, microorganisms also influence physicochemical properties of the water column and sediment pore water. For a complete characterization of lagoonal ecosystems, an approach using organic-geochemical techniques combined with microbiological data and general limnological studies is especially informative.

In this investigation sediments from three coastal lagoons in the district of Macaé, State of Rio de Janeiro, Brazil, each experiencing different degrees of exposure to anthropogenic pollution, were studied by organic-geochemical techniques. Bulk organic matter and extractable lipids were investigated from two locations per lagoon.

Biological sources for organic matter deposited in the sediment, i.e. terrigenous, aquatic, macrophytic, soil-derived or of bacterial origin were identified. In order to discern changes in organic matter composition that occurred over the last decades and also to account for naturally occurring diagenetic processes, two depth intervals (0-3 and 3-6 cm) were sampled at each site. Organic-geochemical results obtained from sediment samples are discussed and interpreted along with available ecological data.

MATERIALS AND METHODS

STUDY AREA

The lagoons are situated in the coastal area of the Macaé region, Brazil (22°30'S and 44°42'W) (Fig. 1) and were formed during the last Holocene transgression period (Perrin 1984). The regional climate is warm and humid; temperatures range from

18.7°C to 27.4°C with lowest precipitation in winter (43.8 mm) and highest in summer (185.8 mm) (FIDERJ 1977).

The lagoons are completely separated from the ocean by sand bars about 100 m in width. Imboacica Lagoon has a length of 5.3 km, the length of the two other lagoons approximate 1 km, max. depths are 2 m (Imboacica Lagoon), 4 m (Cabiúnas Lagoon), and 2.5 m (Comprida Lagoon). Water temperatures of the three lagoons vary around 20°C minimum and 29°C maximum and the values of pH, and salinity decrease from Imboacica Lagoon (pH 8.3; salinity maximum 17.9‰) to Comprida Lagoon (pH 5; salinity maximum 5‰) (Table I), reflecting the influence of marine water. Oxygen concentrations show comparable trends. Salinity in the Cabiúnas and Comprida Lagoons varies due to seawater sprays, high tide events, and artificial opening of the sand bar.

The Cabiúnas and Comprida Lagoons show a typical freshwater fauna (Kozłowsky-Suzuki et al. 1998). Mainly in Comprida Lagoon the water is colored by humic substances, resulting in lower values of transparency (Table I). These two lagoons situated in the National Park Restinga de Jurubatiba are surrounded by a typical dense sand dune vegetation (called Restinga) mainly represented by Theophrastaceae, Anacardiaceae, Myrtaceae, Clusiaceae, Ericaceae, Melastomataceae, and Bignoniaceae (Araújo and Henriques 1984).

All three water bodies are polymictic through the whole year with the littoral zone colonized by macrophytes like *Typha domingensis*, *Eleocharis fistulosa*, *Potamogeton stenostachys*, *Nymphaea ampla*, *Salvinia* sp., and *Utricularia* sp., and the macroalga *Chara*, showing high abundance in the pelagic zone of Imboacica Lagoon.

Eutrophic Imboacica Lagoon is situated in the urban area of Macaé. From 1969 to 1994 about 7.3% of the littoral zone was settled by farms and residences (Santos et al. 1998) that released their agricultural and domestic waste into Imboacica Lagoon. The daily input of nitrogen and phosphorus is about 70 kg and 7 kg respectively (Lopez-Ferreira

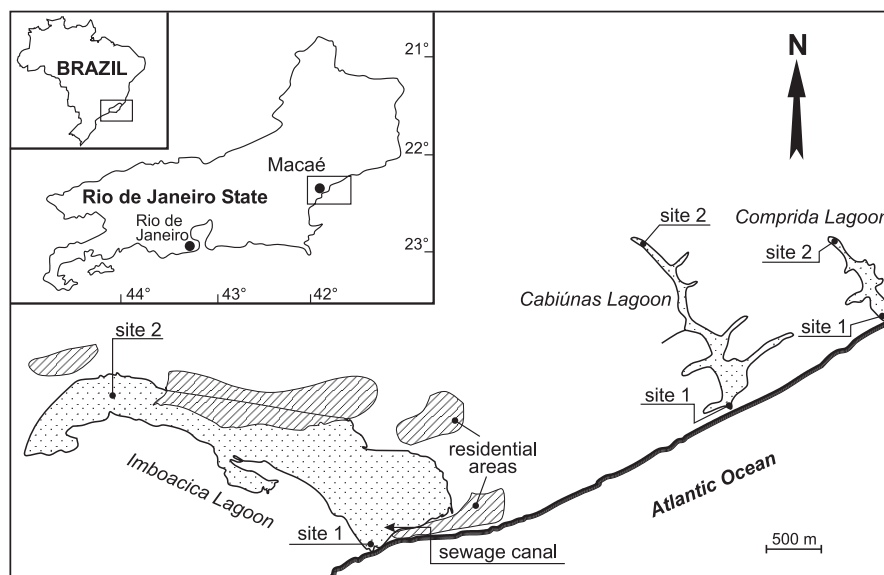


Fig. 1 – Location of study area and indication of sampling sites in three investigated coastal lagoons.

TABLE I

Selected morphological and limnological parameters of Imboacica, Cabiúnas and Comprida Lagoons.

Lagoon		Imboacica	Cabiúnas	Comprida
Area (km ²) ¹		3.26	0.34	0.13
Depth (m) ¹	mean	1.1	2.4	1.6
	max.	2.0	4.0	2.5
Length (km) ¹	max.	5.3	0.9	1.0
Width (km) ¹	max.	1.3	0.2	0.5
Water temperature (°C) ²	min.	18.9	21.0	22.0
	max.	29.9	28.5	28.9
Secchi depth (m) ²	min.	0.7	0.8	0.3
	max.	1.7	2.8	0.7
pH ²	min.	7.2	6.0	3.9
	max.	9.4	7.4	6.1
Conductivity (mS cm ⁻¹) ²	min.	2.2	0.3	0.1
	max.	16.1	14.3	9.6
Salinity (‰) ²	min.	0	0	0
	max.	17.9	12.0	5.0
Oxygen (% saturation) ²	min.	58.6	24.0	43.7
	max.	163.0	116.0	99.0

¹According to Panosso et al. (1998). ²According to Petrucio (1998). Data from May 1993 to November 1995.

1998). Another anthropogenic impact is the artificial opening of the sand bar, causing the drainage of the lagoon to the ocean (last opening for Imboacica and Cabiúnas Lagoons occurred in February 1998). After the drainage the water level is very low and large areas of the lagoon sediment are uncovered.

SAMPLING AND ANALYSIS

Sampling was carried out in September 1998 using a core sampler as described in Ambül and Bühner (1975). At each location sediment cores were taken at the most seaward end (site 1: close to the sand-bar separating lagoon and ocean) and at the most landward position (site 2: macrophytic littoral) of each lagoon and sliced each in 0-3 cm and 3-6 cm subsamples. At both positions selected sediments are representative for the two contrasting sedimentation regimes within the lagoons. Accumulation rates for sediment could not be calculated because absolute age determination via e.g. ^{210}Pb -dating was not available.

Sediments from site 1 consist of coarse sand and gravel, and sediments from site 2 are sandy to silty and organic rich with frequent coarse plant detritus. After removal of coarse pebble (diameter > 10 mm) and plant debris, 12 samples were oven-dried at 45°C. Following homogenization, elemental analysis was carried out on a LECO CS225 with determination of total carbon (TC), total sulfur (TS) and, after acidic removal of carbonates, of total organic carbon (TOC) by combustion and IR-detection of evolved gases. Total inorganic carbon (TIC) was calculated by difference between TC and TOC and calcite carbonate concentrations obtained by multiplication of TIC with a stoichiometry factor of 8.33. Total nitrogen (TN) values were determined on a HERAEUS elemental analyzer on decarbonated sediment (Table II). Due to low concentrations of organic matter total nitrogen was below detection limit in all seaward sites.

Due to the heterogeneity and limited amounts of sediment material the extent of analytical processing had to be restricted, and the initial weight of sediment for extraction varied depending on the

different densities (Table III). Dried samples were extracted with dichloromethane (DCM) using ASE (Accelerated Solvent Extraction) at 75°C and 50 bar for 20 min to obtain the complete free lipid fraction including polar compounds. This approach is comparable to methods applied to modern sediments by Hinrichs et al. (2000), Lutz et al. (2000), Budge and Parrish (1998), and Ficken et al. (2000). Excess solvent was removed by evaporation under reduced pressure and the organic extract was again taken up in small amounts of DCM. BSTFA was added to silylate alcohols and convert carboxylic acids to corresponding esters prior to gas chromatography.

Gas chromatography was performed on a HP 5890 instrument equipped with a HP5 column (length 30 m, 0.25 mm ID, 0.25 μm film thickness) coated with 5% phenylsiloxane. Helium was used as carrier gas with a constant flow of 1.5 ml/min. Injection was done at 40°C in on-column mode and signals recorded by flame ionization detection. Oven temperature was programmed from 40°C to 140°C at 8.5°C/min, followed by at 3.5°C/min to 320°C and kept isothermal for 60 min. Compound identification was achieved by adding external standards and comparing retention times with standard mixtures. Elution order of mono-unsaturated fatty acids was determined by comparison with literature data (e.g. Volkman et al. 1989). Four samples were analyzed by gas chromatography/mass spectrometry using a Single Quadrupole instrument (HP 5890/5889MS Engine). EI-mass spectra were acquired at 70 eV in full scan mode recording from 50 to 650 amu. GC conditions were identical to the ones used for GC-FID.

RESULTS AND DISCUSSION

ELEMENTAL ANALYSES

In all three lagoons samples from the area near the sand bar (site 1: seaward site) have a total carbon content (TC) of less than 1% of dry weight (dw) and a sulfur content of < 0.1% of dw, i.e. close to the detection limit (Table II). This low abundance of organic matter is in accordance with the coarse

TABLE II

Elemental analysis data and organic carbon/nitrogen ratio in sediment samples from Imboacica, Cabiúnas and Comprida Lagoons (1: seaward site; 2: landward site).

Lagoon	Depth interval [cm]	TC[%]	TOC[%]	TIC[%]	TS[%]	TN[%]	TOC/N [molar ratio]
		average n=3	average n=3	average n=3	average n=3	average n=2	average n=2
Imboacica 1 seaward	0-3	0.29	n.d.	n.d.	<0.05	n.d.	n.d.
	3-6	0.46	n.d.	n.d.	<0.05	n.d.	n.d.
Imboacica 2 landward	0-3	6.98	5.93	1.05	0.4	0.43	16.09
	3-6	5.44	5.41	0.03	0.31	0.36	17.53
Cabiúnas 1 seaward	0-3	0.05	n.d.	n.d.	<0.05	n.d.	n.d.
	3-6	0.06	n.d.	n.d.	<0.05	n.d.	n.d.
Cabiúnas 2 landward	0-3	0.70	0.50	0.20	<0.05	n.d.	n.d.
	3-6	1.25	1.25	0	0.06	0.07	20.83
Comprida 1 seaward	0-3	0.11	n.d.	n.d.	<0.05	n.d.	n.d.
	3-6	0.11	n.d.	n.d.	<0.05	n.d.	n.d.
Comprida 2 landward	0-3	21.5	21.98	0	0.34	1.28	20.03
	3-6	34.1	34.88	0	0.48	2.13	19.11

TABLE III

Biomarkers detected in all studied samples (1: seaward site; 2: landward site).

Lagoon	Depth interval [cm]	Initial weight for extraction [g]	Extraction yield [ppm]	Extraction yield [mgExt/gTC]	TAR: even FA [24-34]/[14-18]	FA [18:1 ω 7/18:0]	FA [16:1 ω 7/16:0]	n-alkane ratio 27/[27+29+31]
Imboacica 1 seaward	0-3	50.5	13	4.4	0.001	0.8	0.73	n.d.
	3-6	45.7	11	2.4	0.28	0.74	0.70	n.d.
Imboacica 2 landward	0-3	16.4	5526	80	27.2	0.46	0.34	26.7
	3-6	11.2	5815	107	50.2	0.41	0.23	n.d.
Cabiúnas 1 seaward	0-3	49.4	8	16	0.27	0.58	0.44	n.d.
	3-6	49.1	10	17	0.23	0.71	0.53	n.d.
Cabiúnas 2 landward	0-3	43.4	113	16	6.87	0.45	0.20	14.5
	3-6	46.8	226	18	5.72	0.32	0.24	n.d.
Comprida 1 seaward	0-3	47.0	15	13	1.38	0.43	0.44	n.d.
	3-6	51.3	14	13	2.85	0.36	0.37	n.d.
Comprida 2 landward	0-3	7.6	12440	58	9.05	0.84	0.31	14.8
	3-6	3.8	21719	64	7.04	1.20	0.44	n.d.

grain lithology of these samples. Samples from the macrophytic littoral (site 2: landward site) of each lagoon possess TC contents between 0.7 and 34.1%. Total organic carbon content (TOC) could only be determined for samples with > 1% TC (Table II)

due to restriction in sample amount. Only the surface sediment layer of the landward site at Imboacica Lagoon contained carbonate carbon – the calculated inorganic carbon content (TIC) shows a value of 1.05% corresponding to app. 9% calcium carbon-

ate by weight. The remaining samples are virtually carbonate-free as indicated by lack of CO₂ generation upon decarbonatization. The TC-values measured in triplicate were therefore taken as more accurately reflecting total organic matter abundance and used for normalization of extraction yields. The low abundance of carbonate is in accordance with the freshwater character of the lagoons and the higher acidic environment reported for Cabiúnas and Comprida Lagoons (Petrucio 1998). Enhanced alkalinity and pH-values (7.2-9.4) in Imboacica Lagoon during May 1993 to November 1995 (Table I) may account for the increased TIC-content noted exclusively for the uppermost sample and thus indicate anthropogenic influences.

TOC-values of the landward site at Comprida Lagoon are slightly higher than corresponding TC-values (Table II). The deviation however lies within the analytical error of the LECO elemental analyzer and may also be attributed to sample inhomogeneity. TC-concentrations exhibit significant differences between the lagoons (Fig. 2). Samples from Comprida Lagoon at the landward site, which is covered with dense macrophyte vegetation, show very high TC-values (0-3 cm = 21.5% and 3-6 cm = 34.1%) and TOC-values identical within analytical error limits, i.e. the sediments are carbonate-free. Even so the general trend of high organic carbon content is not affected by the discrepancy between TC and TOC. Increase of TC- and TOC-values with depth is typical for the undisturbed lagoons (Comprida and Cabiúnas) and can be explained by a better preservation with increasing sediment depth induced by stronger reducing conditions or by intensified dilution of superficial sediments by soil derived detritus. The lowest TOC-values at the landward site of Cabiúnas Lagoon reflected the lower macrophyte density. At this area the sediment is composed basically of sandy particles (Gonçalves et al. 1998).

In contrast, Imboacica Lagoon shows an increase of TC for the surficial layer of sediment (Table II), accompanied by a slight increase in TOC-values. As already noted for TIC, this can be attributed to anthropogenic eutrophication in Imboacica Lagoon.

In Comprida Lagoon sources for enrichment of organic matter are autochthonous (primary producers such as phytoplankton and macrophytes) and allochthonous (surrounding vegetation of Restinga). Comprida Lagoon shows the lowest pH and oxygen saturation (Table I) and water is colored dark by humic substances, which may inhibit microbial decomposition of accumulated organic matter (Qualls and Haines 1990). Imboacica and Comprida Lagoons show higher TC-values and sulfur contents in excess of 0.1% (Table II) at the landward site only. Sulfur concentrations of 0.31 to 0.48% in organic matter rich depositional setting are particularly low, even for freshwater environments known to be sulfate depleted. This might indicate low activity of sulfate reducing bacteria and either a limitation of iron or sulfate in the pore water. Consequently a low potential for pyrite fixation prevailed in the lagoonal sediments.

Total nitrogen content determined for the five organic rich samples from landward sites in the three lagoons range between 0.07 to 2.13%. A positive correlation between total nitrogen and TOC is observed with C/N-ratios varying between 16.1 to 20.8 (Table II). This clearly indicates a significant contribution of N-depleted terrigenous organic matter, although strong microbial activity will affect N-cycling in the lagoonal sediments (Meyers and Ishiwatari 1993, Tyson 1995). Pedrosa et al. (1999) described an obvious dominance of the N₂-fixing cyanobacteria *Cylindrospermopsis raciborskii* in N-depleted Cima Lake, Brazil. In Imboacica Lagoon blooms of cyanobacteria dominated by few species – e.g. *Cylindrospermopsis raciborskii* – and one bacillariophyceae (diatom) were observed periodically in summer when the water level decreases and sewage concentrates in the lagoon. Nitrogen fixation by these diazotrophic cyanobacteria might increase the N-content of the sediment. Enhanced nitrogen availability due to sewage input potentially triggered diatom blooms and subsequently nitrogen-rich algal biomass was deposited. A comparable ecological effect could not be observed for the two other Macaé lagoons. The lower C/N-ratio in upper

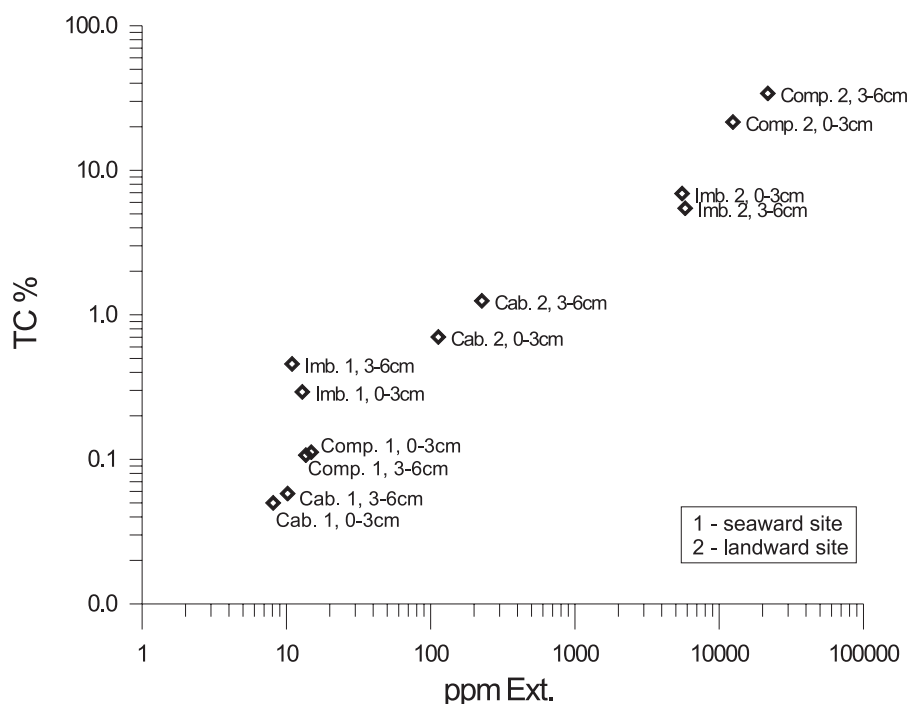


Fig. 2 – Crossplot of total carbon (% dw) versus dichloromethane-extractable organic matter (ppm).

and partly the lower Imboacica Lagoon landward site sediments (Table II) is thus a combined effect of higher macrophyte productivity, nitrogen fixing cyanobacteria and sewage induced diatom blooms. All of these processes are related to anthropogenically-induced eutrophication.

LIPID CONCENTRATION

Organic solvent yields of lipid extraction and TC-normalized extraction yield (mgExt/gTC) are very low for the seaward site, contrasting with the high values found at the landward site from Imboacica and Comprida Lagoons (Table III). As observed for TC, the lowest values of ppm Ext were measured at the seaward site of Cabiúnas Lagoon, reflecting the sediment coarseness and the lower macrophyte density. A relation between lipid yields and TC- or TOC-values is noted for all samples (Fig. 2) and extraction yields of the landward site from all three lagoons increase with depth from 0-3 to 3-6 cm. Increasing TC-normalized extraction yields

with depth indicate enhanced early diagenetic transformation of polymeric biolipids into free organic lipids in the sediment (Table III). Extraction yields normalized to TC for Imboacica Lagoon at the landward site are higher than for Comprida Lagoon (Table III). This points towards a higher relative proportion of autochthonous algal organic material in Imboacica Lagoon than in Comprida Lagoon. TC-normalized extraction yields for Cabiúnas Lagoon at the landward site exhibit low ratios that do not differ from seaward site values. This either indicates strong in situ degradation of sedimented organic matter in the macrophyte littoral region, if oxidizing conditions prevail, or accumulation of predominantly refractory terrigenous organic matter at this site.

Bulk results confirm the contrast in depositional setting between the seaward and landward parts of the lagoons. In the latter organic matter accumulation prevails in the vicinity of littoral macrophyte growth areas. In seaward sites planktonic

algal biomass accumulation predominates. More brackish and eutrophic conditions with intensified algal and bacterial productivity and biomass accumulation were noted only in the uppermost 3 cm of Imboacica Lagoon.

LIPID COMPOSITION

Molecular composition of lipids from the seaward sites

Free lipid extracts of these samples are clearly dominated by short-chain saturated and unsaturated nC14-nC18-carboxylic acids (Fig. 3a-c), mainly produced by planktonic algae and/or cyanobacteria (Han and Calvin 1969, Gelpi et al. 1970, Volkman 1986, Volkman et al. 1989). A minor contribution of short-chain fatty acids will also always result from vascular plants. Especially the extract of the surface sediment (0-3 cm) of Imboacica Lagoon shows an extremely high abundance of n-C16:1 ω 7 and n-C16:0 fatty acids (Fig. 3a) accompanied by abundant n-C14:0 fatty acid. Higher proportions of n-C16:1 ω 7 are known from diatoms but can also be enhanced in cyanobacteria (e.g. Volkman et al. 1998, Wakeham 1995, Napolitano 1998). Conversely, a low diversity of isoprenoidal and long-chain components occurs. This composition might be caused by mass blooms of cyanobacteria and diatoms with high tolerance towards drastically changing water chemistry in Imboacica Lagoon (very low water level, increase of sewage concentration). These changes are a consequence of sewage-induced eutrophication (note vicinity of sewage discharge channel to the seaward site shown in Fig. 1). Alternatively, the artificial opening of the sand bar to stimulate water exchange between lagoon and open ocean might stimulate ingress and growth of brackish algae species. The n-C16:1 ω 9 unsaturated fatty acid almost exclusively reported for heterotrophic marine bacteria (Bertone et al. 1996) was only detected in Cabiúnas Lagoon, although in minor abundance. In this small and ecologically pristine lagoon the lipid distribution is not preferentially controlled by algal/bacterial mass blooms, usually

leading to a dominance of the n-C16:1 ω 7 isomer. A more marine influenced microbial composition in this particular lagoon is favored by artificial opening of the lagoon towards the open ocean.

Water temperature affects the ratio of di-unsaturated vs. mono-unsaturated short-chain fatty acids (Kawamura and Ishiwatari 1981), but this factor does not play an important role in Macaé coastal lagoons. In Cabiúnas Lagoon seasonal variation of water temperature is low – even in winter water is about 20°C (Furtado et al. 2001). Di-unsaturated acids were only detected as trace amounts in landward sites, 0-3 cm, of Comprida and Cabiúnas Lagoons. The proportion of mono-unsaturated (n-C16:1 ω 7) to saturated n-C16-fatty acids consequently indicates changes in algal/bacteria species composition and the intensity of biolipid degradation (Cranwell 1974, 1976, Kawamura and Ishiwatari 1985, Meyers 1997) or even a specific diatom input (Mudge et al. 1998). The Imboacica Lagoon samples (0-6 cm) reveal an averaged value of 0.72 while the two other lagoons show lower averaged values of 0.49 and 0.41 (Table III). In Imboacica Lagoon samples the more labile unsaturated fatty acid may be enhanced due to an almost exclusive origin of autochthonous algal lipids. In the other lagoons a higher proportion of short chain acids could be derived from hydrolysis of very long-chain (C44-C49) esters stemming from terrestrial plant waxes (Kolattukudy 1976). Plant waxes upon longer transportation have undergone more severe diagenetic reactions, and ester hydrolyzation will yield exclusively saturated acids.

The degree of unsaturation of n-C18-fatty acids (C18:1 ω 7/C18:0) for the 0-3 cm samples varies between values of 0.8 to 0.43 for the three lagoons. This ratio reflects the intensity of bacterial input because C18:1 ω 7 is a characteristic compound in bacteria (e.g. Volkman et al. 1998, Mudge et al. 1998). Values decrease from Imboacica Lagoon over Cabiúnas Lagoon to Comprida Lagoon giving evidence for the change in bacterial contribution to the lagoon's organic matter (Table III). The occurrence of C18:1 ω 9 fatty acid in all lagoons can reflect

several different sources: green algae, bacteria, zooplankton, or higher plants (Wakeham 1995, Mudge et al. 1998). With regards to the lipid distribution in the seaward samples, an algal or bacterial origin is obvious, but for Comprida Lagoon additional contribution from higher plants is suggested (Fig. 3c).

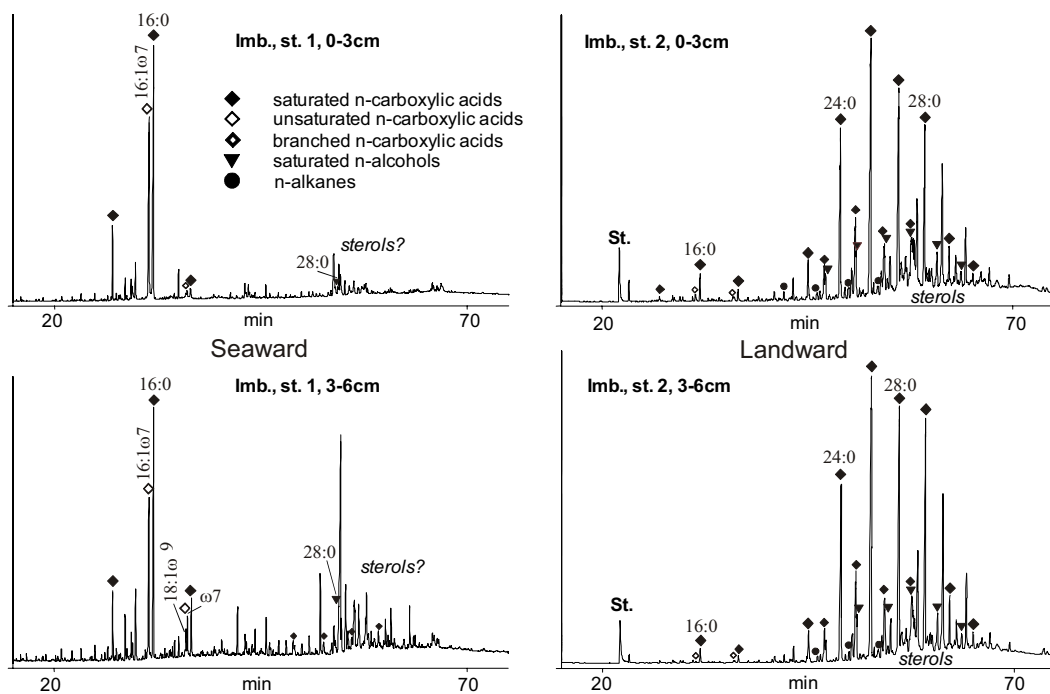
A parameter commonly used for the distinction between terrigenous and aqueous organic compounds is the TARFA (terrigenous/aqueous ratio of fatty acids) (Bourbonniere et al. 1991, Silliman et al. 1996, Meyers and Ishiwatari 1993). This ratio expresses the proportion of even numbered long-chain (C24-C34; in this study) to short-chain (C14-C18) saturated fatty acids. Low ratios were determined for the samples from Imboacica Lagoon: TARFA = 0.001 for the 0-3 cm and TARFA = 0.28 for the depth interval 3-6 cm (Fig. 4a). This very strong predominance of short-chain fatty acids reflects the almost exclusive origin from algal precursor biomolecules, which is in agreement with: i) the sampling site position very close to the sea, ii) the enhanced nutrient availability for algae resulting from man-made eutrophication in the entire Imboacica Lagoon, and iii) the location of the sampling point close to a sewage discharge channel. High concentrations of n-C16 fatty acids were reported from a comparable setting of the Rio Formosa Lagoon, Portugal (Mudge et al. 1998).

In both lagoons still remaining under natural conditions, the molecular diversity of carboxylic acids is significantly greater, with higher values of TARFA for the surface sample (0-3 cm) of Cabiúnas Lagoon (0.27) and especially Comprida Lagoon (1.38) (Table III, Fig. 4a). Whereas Cabiúnas Lagoon is thus almost comparable to the setting noted for Imboacica Lagoon (3-6 cm) before human activity strongly altered TARFA, Comprida Lagoon is dominated by terrigenous biomass even in the seaward part. The low proportion of algal biomass is probably due to low nutrient availability and low pH resulting from high humic and fulvic acid concentrations. Additionally, the low overall size and size/catchment ratio for Comprida Lagoon favor terrigenous organic matter accumulation. The increas-

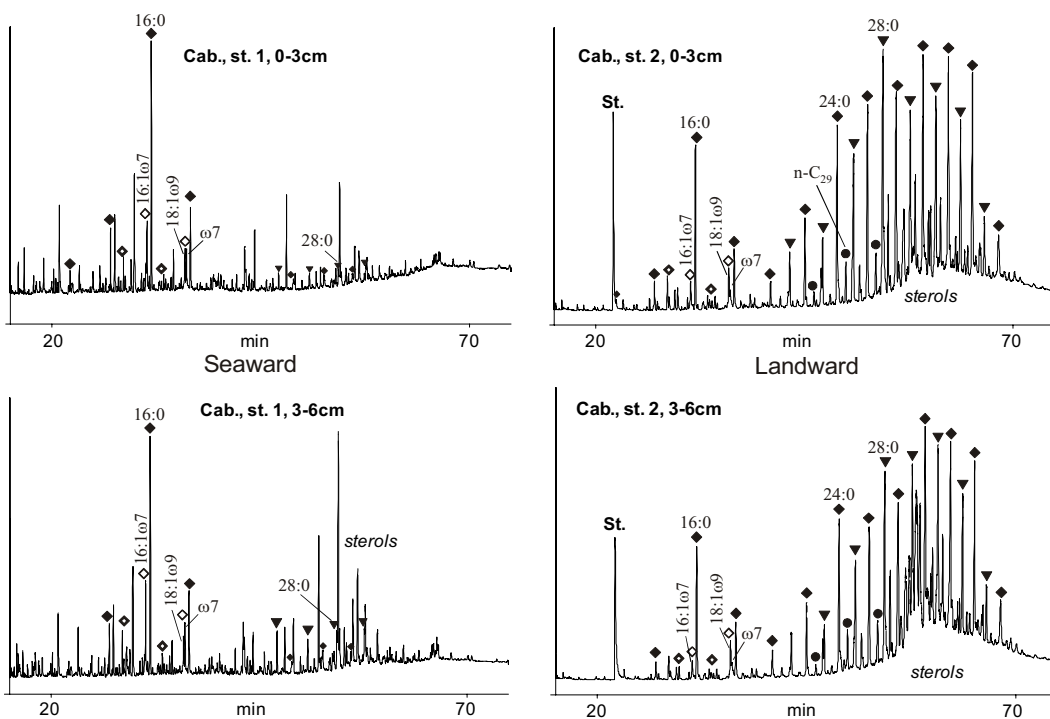
ing values for TARFA of deeper samples can be explained either with preferential degradation and/or condensation/incorporation into polymeric organic matter of short-chain components (Meyers and Ishiwatari 1993) or with a change from limnic to more terrigenous input over time. The relatively high abundance of branched carboxylic acids with chain length C15 to C19 in the free lipids of Comprida and Cabiúnas Lagoons at samples from the seaward site indicates contribution from bacterial species different from those occurred in Imboacica Lagoon, most likely from cyanobacteria.

In comparison to Imboacica Lagoon the smaller surface to volume ratio of Cabiúnas and Comprida Lagoons will have an effect on the transport and the distribution of organic matter within the lagoons. Small size of lagoonal accumulation area in comparison to a large catchment area leads to strong terrigenous character even in the central regions. Another prominent group of bacterial biomarkers, the hopanoic carboxylic acids or alcohols could not be detected by either GC-FID or GC/MS analysis. This might be an indication of a low diversity in the bacterial community or a preferential incorporation of hopanoic acids and alcohols into macromolecular organic matter. Di-, tri- and tetracyclic diterpenoid acids (e.g. agathic acid, abietic acid, pimaric acid), common among land-derived compounds, also could not be detected.

After the carboxylic acids, n-alcohols are the next dominant class of biolipids at samples from the seaward site of the three lagoons (Fig. 3a-c), followed by sterols and n-alkanes. No short-chain homologues are observed for the n-alcohols and n-alkanes in any samples from the seaward site. This absence emphasizes the predominance of terrigenous input, which clearly decreases from Comprida Lagoon to Cabiúnas Lagoon to Imboacica Lagoon (see Fig. 3a-c). The latter location is virtually free of terrigenous n-alcohols and n-alkanes. The terrestrial plant wax origin of long-chain alcohols (Kolattukudy 1976) is further indicated by a pronounced even over odd predominance for the homologous series. In the case of long-chain n-alkanes the plant



(a)



(b)

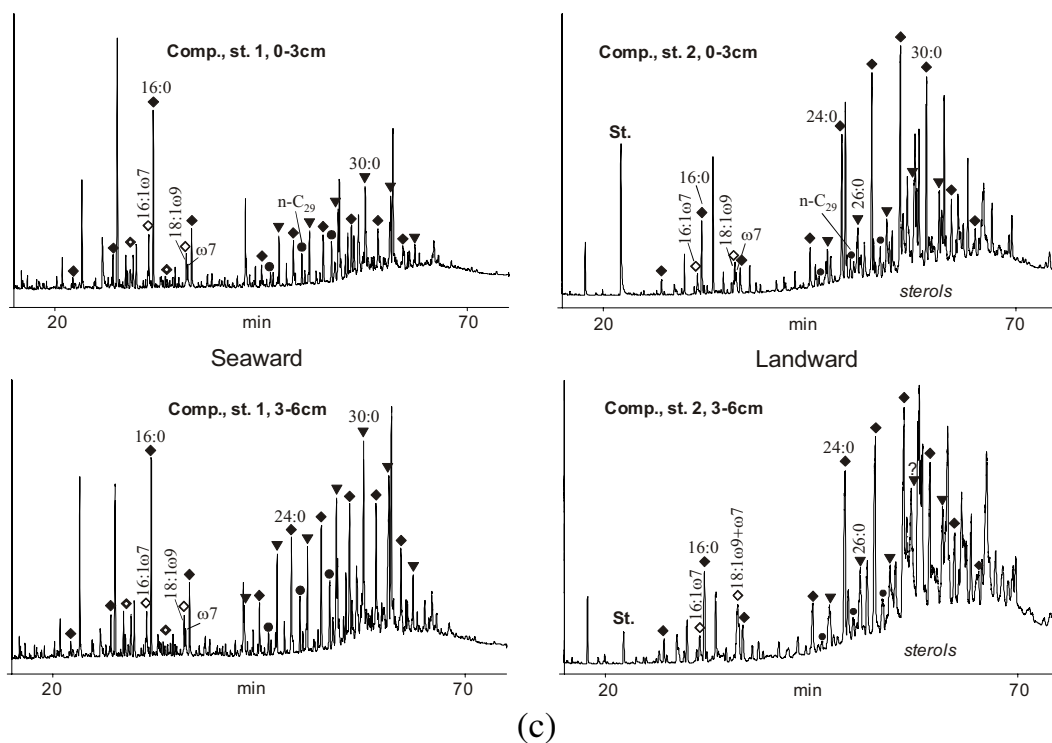


Fig. 3 – a) GC-FID trace of Imboacica Lagoon sediment total extract, derivatized with BSTFA. b) GC-FID trace of Cabiúnas Lagoon sediment total extract, derivatized with BSTFA. c) GC-FID trace of Comprida Lagoon sediment total extract, derivatized with BSTFA.

wax pattern is dominated by odd-numbered analogues (C27, C29, C31 in fig. 3a-c) (Eglinton and Hamilton 1967, Cranwell 1973, Farrimond and Flanagan 1996). This interpretation is in accordance with results obtained from bulk elemental analysis and carboxylic biomarker analyses. Sterols and a second series of unknown compounds (tentatively identified as unsaturated hydroxy-fatty acids) are present in samples from the seaward site but due to lack of mass spectrometric analyses are not discussed further.

Molecular composition of lipids from the landward sites

The gas chromatograms obtained for samples from site 2 (landward) and site 1 (seaward) revealed striking differences in the amounts of short-chain vs. long-chain lipids (Fig. 3a-c, Fig. 4a,b). The sam-

ples from all landward sites are clearly dominated by long-chain saturated carboxylic acids and alcohols reflecting a source of higher plant and aquatic macrophytes (e.g. the emergent *Typha* and submerged *Potamogeton*) in this part of the lagoons and from eroded soil material. The abundant occurrence of very long-chain fatty acids with carbon chain length up to C36 (Cabiúnas Lagoon) is nevertheless unusual and might be connected with different herbs and grasses. The partial predominance of *Typha*, and *Potamogeton* can be responsible only for the enhanced abundance of C20 and C22 fatty acids (Fig. 3b) (Cranwell 1984, Ficken et al. 2000) but probably not for the significant occurrence of C36 compounds. In addition, long-chain n-alkanes mainly with a strong odd over even predominance in n-C25 to n-C35 range and some sterols are present.

Short-chain carboxylic acids again have to be

assumed to derive from algal/bacterial lipids and to a minor degree from hydrolysis of wax esters. The ratio of C16-unsaturated to saturated acids in samples from landward sites in general is lower, which can be explained by higher relative contribution of plant wax acids in relation to algal/bacterial lipids. Also the 18:1 ω 7/18:0-FA ratio is lower in samples from the landward site, except for samples from Comprida Lagoon where the enhanced value for the 18:1 ω 7/18:0-FA ratio of 0.84 and 1.20 might indicate additional sources than bacteria (Table III). In Comprida Lagoon the high abundance of both C18:1 ω 9- and C18:1 ω 7-fatty acids could be attributed to higher plant input (Cranwell 1982, Mudge et al. 1998), although C18:1 ω 9 is also reported a prominent marker for zooplankton contribution (Wakeham 1995). The preference of the unsaturated acids is probably due to a specific plant precursor (e.g. an aquatic macrophyte). Alternatively, an origin not preferentially from epicuticular wax coatings but from intracuticular waxes or different plant constituents (essential oils, seed coatings, etc.) might be possible. The predominance of C18:1 ω 9 over C18:1 ω 7 in all landward sites is concordant with this observations (Fig. 3a-c). Nevertheless, it has to be considered that C18:1 ω 9 can also originate from phytoplanktonic sources (e.g. Napolitano 1998), which agrees with their relative abundance in the seaward sites (Fig. 3b).

Extracts of Imboacica Lagoon show the most variable compound distributions. Compared to samples from the seaward site, which have almost exclusively n-C16:1 ω 7- and n-C16:0-fatty acids, extracts of Imboacica Lagoon from the landward site have the lowest concentration of short-chain acids among the three lagoons. In contrast, Cabiúnas and Comprida Lagoons still show relatively high concentrations of C16- and C18-fatty acids, although the terrigenous long-chain components dominate. Calculation of the TARFA confirms these observations with values varying from 5.7-9.0 for Cabiúnas and Comprida Lagoons to 27 and 50 for Imboacica Lagoon (Fig. 4b, Table III).

Another characteristic of Imboacica Lagoon is

the very high abundance of the long-chain fatty acids compared to alcohols. The opposite is noted for Cabiúnas Lagoon with Comprida Lagoon being intermittent (Fig. 3a-c). Two possible mechanisms may account for the high acid to alcohol ratio in Imboacica Lagoon. First, oxidation of primary n-alcohols will lead to the corresponding n-fatty acids. This oxidation process may be completely abiotic or microbially mediated (George and Jardine 1994). The oxidative degradation of lipids will ultimately also decrease the relative proportion of C14- to C18-compounds, which is observed for Imboacica Lagoon samples. This stronger oxidative regime inferred from carboxylic acid distribution is in part substantiated by in situ measurements of oxygen saturation (Petrucio 1998). Another route that could potentially lead to selective accumulation of carboxylic acids may be the preferred deposition of intracuticular carboxylic acids in fine-grained Imboacica Lagoon sediments showing almost no intact plant fragments. In this sediment type, quite distinguishable from Comprida and Cabiúnas Lagoons, a higher proportion of dissolved free lipids is likely to accumulate.

Long-chain alcohols found in samples from Cabiúnas Lagoons could potentially also be attributed to freshwater algae (Volkman et al. 1999). However, such long-chain algal lipids show another distribution pattern and require almost uni-specific input from *Eustigmatophyceae*, not reported in Cabiúnas Lagoon (Mello and Suzuki 1998). Volkman et al. (1999) reported C22 as the main straight-chain alcohol and high amounts of mono-unsaturated long-chain alcohols, and neither was detected in the lagoonal sediment extracts.

STEROLS

The distribution of several ubiquitous steroids can provide supporting information concerning the source of sedimentary organic matter (Huang and Meinschein 1979, Cranwell et al. 1987). The evaluation of organic matter sources to lagoonal sediments can be performed using a ternary diagram displaying relative proportion of C27, C28 and C29

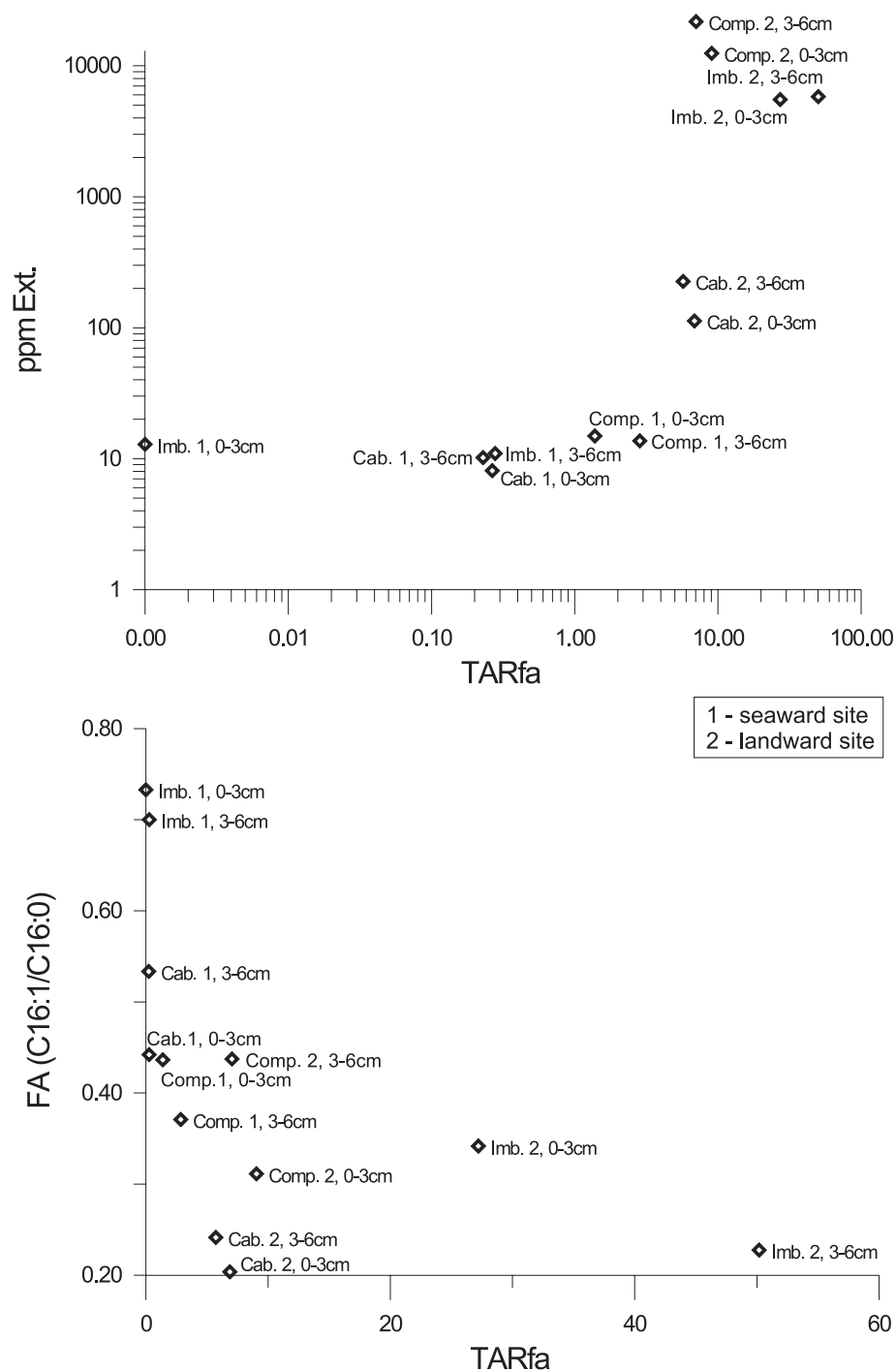


Fig. 4 – a) Crossplot of terrestrial over aquatic ratio of fatty acids [TARFA] versus extractable organic matter (ppm); b) Crossplot of terrestrial over aquatic ratio of fatty acids [TARFA] versus ratio of mono-unsaturated/saturated C16-fatty acids [FA(C16:1 ω 7/C16:0)].

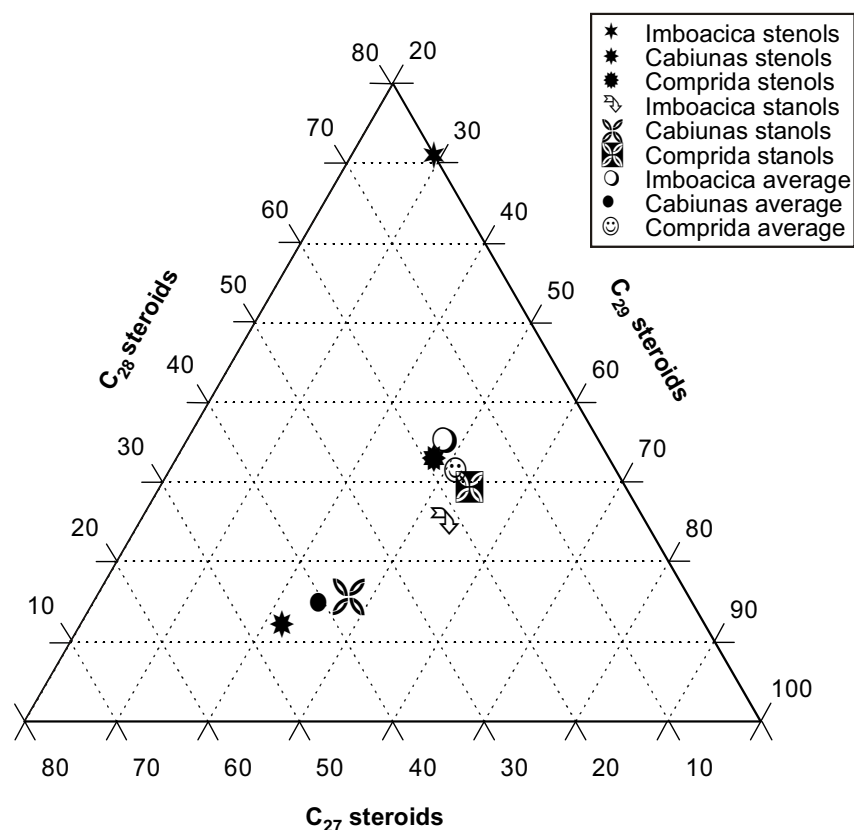


Fig. 5 – Ternary diagram showing distribution of saturated and mono-unsaturated C27 to C29 steroids.

steroids (Fig. 5). For the surficial samples from the landward site from all three lagoons, the relative proportions of mono-unsaturated Δ^5 steroids cholesterol, campesterol, sitosterol and saturated counterparts cholesterol, campestanol and stigmasterol were quantified from mass fragmentograms using m/z 215 key ion (Fig. 6). As a major C29-steroid stigmasterol [24 β -ethyl-cholest(5,22)dien-3 β ol] occurs in significant concentrations in all samples. If this C29 stanol were included in the calculations, all samples would acquire an even more terrestrial source signature.

The extreme position of Imboacica Lagoon with an obvious dominance of C28 and C29 higher plant steroids and no detectable amounts of C27 algal steroids confirms the results obtained from n -carboxylic acid analysis. Cabiúnas and Comprida

Lagoons sediments are more enriched in the algal/zooplankton derived cholesterol (C27) and therefore plot more towards the center of the ternary diagram, in an estuarine/lagoonal position. The elevated amount of C27 and reduced contribution of C29 steroids for samples from Cabiúnas Lagoon is in agreement with an enhanced contribution from phytoplanktonic algae as compared to samples from Comprida Lagoon. Sterol discriminant analysis places all lagoonal samples in the estuarine/lagoon/bay field and is not particularly well suited for deduction of predominant organic matter sources.

OTHER SOURCES

The fact that no methane production was observed for any lagoon sediments (Furtado et al. 2002) is confirmed by biomarker analysis on those samples.

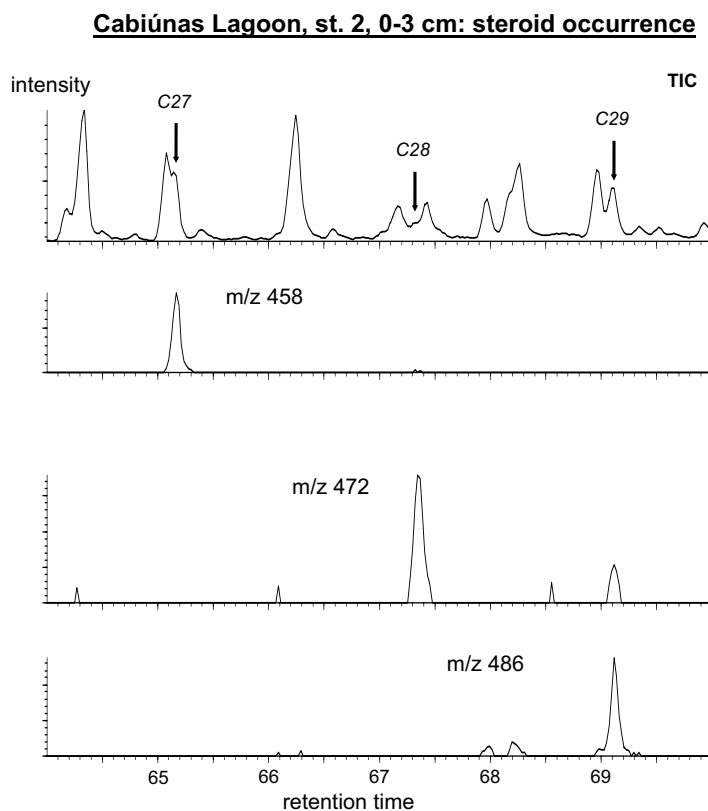


Fig. 6 – Total reconstructed ion current and selected mass fragmentograms indicating steroid distribution.

No significant amounts of biomarkers from methanogens (e.g. specific isoprenoids like pentamethylicosanones, biphytanes, squalene) or methylotrophic bacteria (3α -methylhopanes) could be detected. Additionally, hopanoids, typical for bacteria and cyanophytes, are absent in all samples from the landward site as revealed by GC/MS analysis. The only indication for bacterial activity is the abundance of C16 and C18 mono-unsaturated ω 7 fatty acids and higher concentrations of branched isomers of short-chain fatty acids (C15, C16, C17) in samples from the seaward site. It follows that in three Brazilian lagoons studied here a special bacterial community prevails that differs from microbial assemblages common in other lakes. The relatively low abundance and productivity of pelagic bacte-

ria were between 1.0 to 2.3×10^9 cells/l and 1.1 to $33.7 \mu\text{g C/l}$ in Cabiúnas Lagoon (Furtado et al. 2001). It remains to be confirmed by future studies if this specific bacterial association is a general feature of lagoonal environments.

Biomarkers like the fecal marker coprostanol (Nguyen et al. 1995) or PAH (polyaromatic hydrocarbons resulted from by incomplete combustion of fossil fuels) indicative for anthropogenic input into the lagoons could not be identified in significant concentrations. One reason for this absence might be fast intensive bacterial reworking of sewage material in the lagoon area where the discharge takes place, resulting in a lower fixation of anthropogenic pollution markers.

CONCLUSIONS

Organic biomarker analysis investigation has identified sources of organic matter, evaluated its diagenetic transformation pathways and detected human disturbance in three Brazilian lagoons.

Imboacica Lagoon, located in an urbanized area, has been receiving increased anthropogenic input over recent years. This impact is preferentially archived in the topmost 3 cm of sediment, whereas the deeper sediment layer shows less evidence of human influence. Eutrophication is a key process controlling composition of lagoonal biological communities particularly by favoring algal (diatoms) and cyanobacterial growth. One other important process is the artificial opening of the sand barrier between lagoon and ocean that controls water chemistry and exchange of marine and lagoonal fauna. Hence, physicochemical properties like alkalinity are affected as well, as manifested in lithological change towards more carbonate precipitation in the uppermost 3 centimeters of Imboacica Lagoon sediment.

In more isolated Cabiúnas and Comprida Lagoons pristine natural ecosystems are preserved. Terrigenous organic matter dominates which is exceptionally well preserved. This is related to the physicochemical properties of lagoonal waters, in particular their high acidity. Differentiation between landward and seaward sites in these lagoons is less pronounced because of much smaller surface/volume to catchment ratios. No anthropogenic influences are yet detectable in sediments of Cabiúnas and Comprida Lagoons.

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RESUMO

As lagoas costeiras são importantes ecossistemas costeiros no Estado do Rio de Janeiro, localizadas entre áreas urbanizadas da costa e o oceano aberto. Vários estudos enfocando o sedimento, a água, a comunidade zooplancônica e comunidades bacterianas demonstraram a extensão da influência antrópica sobre estes ecossistemas aquáticos. Neste estudo foi investigada a composição da matéria orgânica presente no sedimento, assim como, alguns parâmetros microbiológicos, com o intuito de caracterizar três lagoas costeiras. Matéria orgânica e lipídeos (hidrocarbonetos alifáticos, álcoois e ácidos graxos, esteróis) foram analisados em dois pontos de amostragem em cada lagoa (região continental e região marinha), em dois intervalos da profundidade (0-3 e 3-6 cm). A lagoa Imboacica apresentou maior influência antrópica nos últimos anos, representada na fração superficial do sedimento (0-3 cm), visto que na camada mais profunda (3-6 cm) observou-se reduzida influência humana. A eutrofização da lagoa ou disponibilidade de nutrientes favoreceram o crescimento acentuado de algas e cianobactérias. Nas lagoas Cabiúnas e Comprida, afastadas do perímetro urbano, as características naturais mostraram-se preservadas. A matéria orgânica oriunda de plantas vasculares é dominante, apresentando-se excepcionalmente bem preservada em função das características ácidas da água. A discreta diferenciação entre a região continental e a marinha, nestas duas lagoas, está diretamente relacionada à reduzida razão superfície/volume destes ecossistemas aquáticos. Nenhuma influência antropogênica foi observada no sedimento das lagoas Cabiúnas e Comprida.

Palavras-chave: lagoas costeiras, produção aquática/terrestre, lipídios, ácidos graxos, esteróis, Brasil.

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