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# Sedimentary Transport Influences on Diagenetic Processes at the Amazon Continental Shelf, Brazil

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## Authors' contributions

All authors participated of the samples collection, date and statistical analysis and wrote the first draft of the manuscript.

# Article Information

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# ABSTRACT

This research aimed to correlate the sedimentary transport with the diagenetic processes in the coastal zone and Amazon Continental Shelf (ACS). Physical and physical-chemical parameters, trace element contents (Cr, Pb, Ni, Zn and Hg), and O<sub>2</sub>, CO<sub>2</sub> and iron flux were determined in sediment and pore water. Sedimentary incubation (96 hours) and algorithms were applied to determine the variation of the activity coefficient ( $\Delta$ I) and ionic strength (F<sub>i</sub>) of the predominant chemical species, and to estimate the net production and mineralization of the organic matter ( $\Delta$ CO<sub>2T</sub>) in the system. There are not many studies applying incubation tests to identify the diagenetic processes, especially in fluvial-marines sediments. The results showed a strong zonation associated to the transport and deposition processes, influenced mainly by the grain-size and texture of sediment and fluvial streams. The distribution of trace elements followed the trend of the sedimentary pattern, with higher levels of metals in the degree of importance of each parameter under the distribution and mobility of trace elements, suggests that the mobility of Cr, Ni and Zn is controlled by depth, clay and organic compounds contents, and concentration of dissolved oxygen.

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The vertical flow of  $O_2$  and  $CO_2$  and the Fe<sup>2+</sup>/Fe<sup>3+</sup> ratio in the pore water suggest a predominance of organic matter oxidation in the sedimentary layer between 0.0 and 0.2 m, with partially anaerobic mineralization of the sediments below 0.4 m. Increases in trace element concentrations were observed in iron reduction zones, indicating processes of desorption of oxides and hydroxides of Fe and mineralization of organic matter. The extrapolation of the results of the incubation test to the studied system allowed to establish three hypotheses related to the diagenetic processes: 1) the flow of marine currents may be allowing the aerobic oxidation in the sandy sediments, with the nitrification route more accentuated than the ammonification route; 2) in the region of the coastal zone and inner continental shelf the routes of oxidation and reduction may be alternating according to the physical-chemical factors and seasonality; 3) in the coastal zone and inner shelf the net mineralization rate exceeded the net production rate of the organic matter ( $\Delta CO_{2T} > 0$ ).

Keywords: Pore water; mineralization; net productivity; sedimentary characterization; diagenetic processes; Amazon continental shelf.

# **1. INTRODUCTION**

The sedimentary transport regime analysis is much applied in association with geochemistry in quantitative studies, revealing areas of cumulative tendency or sedimentary dispersion. Studies that have demonstrated the sedimentation associated rate with microbiological activity, organic carbon content and seasonal events affect the concentration of dissolved chemical species directly in the sediments [1,2], especially in water [3,4], which justifies the influence of sedimentary transport on the diagenetic regime, with particular emphasis on quantifying the seasonal rates and spatial patterns of carbon remineralization.

Pore water can be interpreted as a physicalchemical and biological transition phase between what happens in the water and sediment compartments. This suggests a complexity of factors and processes acting jointly in the genesis and flow of chemical species, either by diffusion. advection. precipitation and/or dissolution of minerals. metallic adsorption/desorption, bioturbation, production or mineralization of organic matter [4-6]. The composition (clay and organic matter), physical (porosity) and physical-chemical properties (pH and alkalinity) of the sediments influence in the diffusion and transport of the chemical species in the pore water [5-7], either by vertical or horizontal movements.

In the region between the Amazon and Pará mouths, and the Amazon Continental Shelf (ACS; Fig. 1), the sedimentary dynamics is the largest in the world, both in volume and dispersion area. The distributions of minerals in ACS sediments [8], as well as organic and inorganic components, including trace elements [9], are of unequaled proportions to any other fluvial-marines system on the planet. Fluvialmarines currents play an important role in this sedimentary dynamics, acting individually or together, and seasonally in the Amazonian system. The balance between the erosion, transport and sedimentation processes interferes with the volume of sediments and, consequently, with the concentration of chemical species in the fluvial-marine system. The Amazon River presents the highest suspended solids load transported in the world, about 1.2x10<sup>9</sup> metric tons/year of sedimentary debris to its lower reaches [10], and that once reaching the continental shelf will submit to a coastal dynamic, mainly by the influence of the North Brazilian Current (NBC) along the American continent. Sedimentary load representing ~3% of the global riverine particle flux, accumulates off the river mouth in Brazilian coastal waters as deltaic mud deposits having a northwest trending strike [11].

reactions Diagenetic and sediment-water exchange processes occurring in deposits of the ACS determining the eventual influence of the Amazon on a range of marine elemental cycles and characteristics of the resulting sediment record [12]. The diagenetic processes involve the genesis of chemical species with consequent alteration in these compositions in the environment. Among the processes involved in diagenesis, the net production and mineralization of organic matter have been highlighted in geochemical studies, as they interfere in the flow and balance of gases (CO<sub>2</sub>, O<sub>2</sub>, CH<sub>4</sub>, H<sub>2</sub>S and  $SO_4^{2-}$ ) in the water and sediment compartments, in the metallic mobilization, and in the aerobic and anaerobic benthic microbial productivity.

This research aimed to correlate the sedimentary transport with the diagenetic processes in the

area between the coastal zone and outer shelf of the ACS. For this, 1) the distribution and flow of trace elements (Cr, Pb, Ni, Zn and Hg); 2) the profile of the vertical flow of  $O_2$  and  $CO_2$  in the pore waters, relating them to the iron redox potential (Fe<sup>2+</sup> and Fe<sup>3+</sup>); and 3) net production and mineralization of organic matter, identifying the dominant pathway in the pore water system were determined and/or estimated.

# 2. STUDY AREA

The study has been conducted in the Amazon River, nearby coastal zone (CZ) and Amazon Continental Shelf – ACS (47°52'W-4°19'N and 51°04'W-2°16'S) between Orange Cape (State of Amapá) and Pará River mouth (State of Pará, Fig. 1). It is a dynamic region, influenced by the meeting of continental waters with the Atlantic Ocean on influence at the Equatorial North Current (ENC) and NBC. There is also interference from atmospheric forces as the Intertropical Convergence Zone (ICZ), generating trade winds and rainfall, which determining the climatic seasons and the hydrological pattern of Amazonian rivers [8]. This complex system of water circulation influences sediment transport and deposition, acting in a selective way to separate the sedimentary fractions and the content of organic material by zonation in the coastal zone and ACS. The climate of the region is "Am" hot and constantly humid (monsoon climate) for Marajó Bay, and "Af" equatorial for Amazon and Pará mouths, with mean temperature of 28 °C and rainfall exceeding 2000 mm/year, especially between the months of December and February. The Amazon coastal zone includes diverse ecosystems as mangroves, dunes, coastal forests, freshwater coastal, estuaries and river deltas with great aquatic and terrestrial biodiversity associated. Descriptions of the study area and sedimentary patterns can be found in many researchers [8,9,11,13].



Fig. 1. Map of the ACS and coastal zone between orange cape (OC) and Pará River mouth. Details of the shelf – coastal zone vector 1 – 4 (transect)

Legend: NC= North Cape; MC= Maguari Cape; AR=Amazon River; PR= Pará River; isobaths in meters

# **3. MATERIALS AND METHODS**

Physical-chemical, water column hydrography and sedimentary load measurements were made following the regime of seasonality in the region. The bathymetric lines (isobaths) showed in the Fig. 1 were obtained from probe eco-bathymetry and for consultations to nautical charts. A tendency line (exponential fitting curve) with the respective differential equations was determined for the bathymetric profile, with orientation axis from the coastal zone in direction to outer shelf. The sediment samplings, however, were performed during the months of lowest flow (May and June) in the Amazon and Pará rivers and at the ACS. Samples of bottom sediments (0.0 - 0.5 m) were obtained between the 5 and 100 m isobaths using Van-Veen sampler (A=682 cm<sup>2</sup>) to the bottom sediments, and box-core sampler (A=177 cm<sup>2</sup>) to the vertical samples. The sediment samples were stored in plastic bags and kept in freezer at -5°C until the analysis. Vertical samples were used to determine the pH, salinity (dS/m) and dissolved oxygen (DO µmol/L) in the pore water.

At chemistry laboratories at Pará Federal University (UFPA) and National Institute of Research in Amazonian (INPA), the sediment samples used for geochemical analysis were fractionated, oven dried (45±0.5°C) and homogenized, sieved and pulverized until very fine sand fraction (125 - 63vfs µm). The pH of the samples was neutralized with 2M NH<sub>4</sub>OH solution and 30 mL of Mil-Q water followed by titration of the chloride, to ensure no interference of the CI- analyzes. For the incubation test, fresh samples were homogenized and stored in polyethylene bottles with a sealed acrylic lid. DO and CO<sub>2</sub> were determined with probes and analytical methods (titration) from the extraction of the pore water, and profiles of oxygen (DO  $\mu$ mol/L) and total CO<sub>2</sub> ( $\Sigma$ CO<sub>2</sub> mmol/L) were established in the vertical layer of the sediments sampled for the vector 1 - 4 (Fig. 1).

Grain-size analysis was determined by the gravimetric method [14]. Organic matter (OM) was determined by hot acidic extraction with excess hydrogen peroxide (30%) at 100°C [15]. The organic carbon content (OC) was determined by the Walkley-Black method [16], where the organic carbon of the sample is oxidized to  $CO_2$  with potassium dichromate [K<sub>2</sub>Cr<sub>2</sub>O<sub>7</sub>] and concentrated sulfuric acid [H<sub>2</sub>SO<sub>4</sub>]. The chromium of the extractive solution is reduced to Cr<sup>3+</sup>, and the excess of potassium

dichromate is titrated by the ammonium iron (II) sulfate [(NH<sub>4</sub>)<sub>2</sub> Fe(SO<sub>4</sub>)<sub>2</sub>.6H<sub>2</sub>O]. The calculation is done by applying equation 1 [14]. Free CO<sub>2</sub> (CO<sub>2L</sub> mmol/L) and total CO<sub>2</sub> (CO<sub>2T</sub> mmol/L) were determinates with probe and also obtained analytically by titrimetric method 4500-CO<sub>2</sub> A and The titration was performed D [17]. potentiometrically with NaOH to bring the sample to pH 8.3 and HCl to pH 4.3. From the results, the carbonate and bicarbonate alkalinities also were calculated. Inorganic carbon (IC) was calculated from the amount of total CO2 per molar transformation [15].

$$OC = 0.06V(40 - Va \times f)$$
 (1)

Where: OC (g/kg); V = volume of potassium dichromate and  $V_a$  = volume of ammonium iron(II) sulfate consumed in the reaction; f = 40/ volume of ammonium iron(II) sulfate used in the blank titration; 0.06 = correction factor.

Trace metals Cr, Pb, Ni and Zn (mg/kg) were extracted using mixture [HNO<sub>3</sub> + HCI]:HCIO<sub>4</sub>:HF (Merck 2:1:1) in an open system and determined in a flame atomic absorption spectrometry Shimadzu AA 6800 (Standard Method 3111B modified) [17]. Total Hg (ng/g) was extracted adding dry sediment and  $V_2O_5$  (1:1) and after a solution of HCI:HNO<sub>3</sub>: H<sub>2</sub>SO<sub>4</sub> (Merck 3:1:6) in a test tube closed, and determined in cold-vapor atomic absorption (Standard Method 3112B and 3500-HgB adapted) [17]. For analytical quality control, the recovery levels of acid digestion and determination of metals were tested using certified reference material (SRM 2710 nearshore seawater, National Research Council of Canada) analvzed in triplicate. The determinations of the total and reduced iron (Fe<sup>2+</sup>) fractions were obtained from the methods described by Jackson [14], Silva [15]. By mass difference the Fe<sup>3+</sup> fraction was calculated and Fe<sup>2+</sup>/Fe<sup>3+</sup> ratio was established. The oxidation potential of the solid-reactive phase of the iron in the sediments was estimated by leaching wet sediment in 6N HCl for 15 minutes at 22°C (10:1 mg sediment/ ml HCl) and analyzing the leachate for total Fe and Fe<sup>2+</sup>.

The river hydrographic states were used to identify seasonal sampling times, and the respective seasonality. Sampling of the coastal zone and ACS were designed to include a range of major environmental conditions corresponding to changes in likely sedimentation patterns, Amazon River and Para River flows, trade wind stress, and influence of the Equatorial North Current and North Brazil Current (NBC). The study of the diagenetic processes was concentrated at the sampling sites inserted in the vector 1 - 4 (Fig. 1), starting at the deepest local (site 1), near the isobath 100 meters, and ending at the coastal zone (site 4), already under influence of the river currents. The sedimentary transport regime was identified by the analysis of the tendency of space-time clustering of the sedimentary fractions. To estimate diffusive sedimentary flux, the incubation method described by Mackin and Swider [18] was used. In order to estimate potential diffuse exchange rates and the rates of net production and mineralization of organic matter, samples stored in the sealed vials were shaken vigorously for 5 minutes and then allowed to stand at 20±0.5°C and light/dark phases controlled for 4 days (96h). Pore water samples were removed of the sealed vials at incubation times  $t_0=0h$  and  $t_7=96h$  with a syringe to estimate  $CO_{2L}$  and  $CO_{2T}$  contents, according to the analytical protocols described by Silva [15], APHA [17]. Preliminary tests showed that after four days is more difficult to estimate the diffuse sedimentary flux for the adopted method as well as for the conditions at the ACS, especially in the coastal zone with high OM levels. This occurs because the DO levels after four days are very low almost imperceptible. Thus, a maximum time of measure of four days or 96 hours was established ( $t_0$ =0h and  $t_7$ =96h in equation 4). Diffusive fluxes of the sediments were calculated from flux (vertical sampling from box core) by dividing the slope of a least squares line fit of the total mass change of a solute at time 't' in overlying water versus elapsed incubation time by the area of the flux core. For this study the close system method was adopted for incubation tests (to see item 4.4 Sedimentary incubation). Pore water samples were collected and analyzed for DO and CO2T according to methods already decrypted. The DO and CO<sub>2</sub> curves of consumption/ production in function of the time ( $\Delta t$ ) are determined, suggesting the speed of the reaction of the diffuse flux (slow moderate - fast). Exchange liquid flux by diffusion at the water-sediment interface can be estimated from tables according to temperature and pressure in the environment (pressure of gas diffusion). The net production is obtained from the determination of the CO<sub>2</sub> rates in function of the time ( $\Delta t$ ). Using the algorithms of the software Carbmar 1 and 2, and Alcagran (Basic© and TBasic© 1994) and applying equations 2 -7, the net production rate for  $\Delta(CO_{2T})_b < 0 \Leftrightarrow$  $\Delta(O_2)_b > 0$ ; net mineralization rate for  $\Delta(CO_{2T})_b > 0$  $0 \Leftrightarrow \Delta(O_2)_b < 0$ ; and net remineralization reaction rate of the organic matter in the orientation of the vector 1 - 4 (Fig. 1) were calculated.

$$I = 0.5 \times \sum Z_i^2[i] \qquad (eq. 2)$$

$$Logf_i = -A \times Z_i^2 \times \left[ \left( \frac{I^{1/2}}{1 + I^{1/2}} \right) - 0.3I \right] \qquad (eq.3)$$

$$\Delta(CO_{2T})_a = 0.5 \times (FCO_{2T}(t_0) + FCO_{2T}(t_7)) \qquad (eq.4)$$

$$\Delta(CO_{2T})_b = \Delta(CO_{2T}) - \Delta(CO_{2T})_a \qquad (eq.5)$$

$$C_{eq} = C + \frac{Mn}{6} + \frac{(Cr + Mo + V)}{5} + \frac{(Ni + Cu)}{15} \qquad (eq.6)$$

$$\sum \Delta M(t) = [C(t) - Cc(t - \Delta t)] \times V' \qquad (eq.7)$$

Where: [eq.2] I= is the activity coefficient of the ionic functions; i = major ionic species with corresponding electrical charges (Z); [eq.3] f = ionic strength for A= 0.5 [19,20]; [eq.4]  $(CO_{2T})_a$  exchange liquid flux by diffusion at the watersediment interface in the interval *t*, and FCO<sub>2T</sub> in  $t_0$  and  $t_7$  represent the flux of CO<sub>2</sub> in each moment *t*; [eq.5]  $(CO_{2T})_b$ = net production or mineralization rate of  $CO_{2T}$ ; [eq.6]  $C_{eq}$ = represents the equivalent carbon with the element contents in %; [eq.7] C(t) = solute concentration at time t; Cc = concentration of solute corrected in the previous sample as a function of time variation (*t* -  $\Delta t$ ); and V = volume of water overlying the core [18].

#### 4. RESULTS AND DISCUSSION

#### 4.1 Sedimentary Characterization

Table 1 presents some relevant characteristics of the sediment and pore water determined at the ACS. The granulometric analysis revealed the presence of mud sediments with high percentage of fine silt and clay in the coastal zone, with variation of 80.6 - 90.6% (mean 84.8±3.6%). In the continental shelf area, not making a distinction between the influence of the river and marine currents, the values had high variation, from 5.8 to 81.4% (mean 53.6±29.1%). Isolating the areas by influence of the currents, we have that the ACS inner shelf (sites 3 and 5) had concentrations of fine sediments between 70.7 -81.4% (average 76.0±5.3%); and on the outer shelf (sites 1 and 2) from 5.8 to 56.1% (mean 31.2±25.5%). The coastal zone and the inner shelf presented enrichment of the bottom sediments with fine organic material originating from Amazon River sedimentary load. A similar pattern of change in particle size was observed by Gibbs [13], Garrels and Thompson [21], along the Amazon Shelf between 10 and 50 meter isobars above the North Cape (State of Amapá). According to the authors, the transition in the sedimentary pattern extended from the interior delta of the Amazon River towards the inner shelf with extensively reworked by intense tidal currents and waves. The bathymetry suggests a behavior in a little negative exponential curve. In other words, the increase in the depth of the coastal zone towards the ocean floor was slightly mild, with two very evident areas of reduction in the inclination of the curve: first one at 40 meters and the second one near 80 meters deep (Fig. 2A). The trend of accumulation of fine sediments in the fluvial-marine environment followed a distribution pattern, which can be explained by a variation of Gaussian the curve. The sedimentation - deposition and physicalarea reworking patterns indicated an of maximum thickness of the thin sediment laver with approximately 62 cm high and 49 km away from the coastal zone (Fig. 2A). Similar patterns were observed by [11,12]. Aller et al. [12] observed a layer of fine sediment between 50 and 60 cm thick accumulated about 65 km from the coastline, above the North Cape between 1.5° and 3.5° north latitude, which the authors identified as a result of physical reworking of sediments from a circulation pattern influenced by NBC. According to the authors, vertical sediment-water zonation in particle mobility and concentration are therefore typically present at the region, with the relative zonal thicknesses determined seasonally by the relative dominance of physical processes and sediment supply at each site.

The organic matter (OM) and organic carbon (OC) contents in the study area varied, respectively: in ACS (sites 1, 2, 3 and 5) OM= 0.6 - 2.8% (mean 1.3±0.4%) and OC= 0.3 -1.8% (mean 0.8±0.3%); only on the outside of the shelf, under the influence of marine currents, OM= 0.6 - 2.8% (mean 1.5±0.5%) and OC= 0.3 -1.8% (mean 0.8±0.3%); in the inner shelf, OM= 1.0 - 1.6% (mean 1.2±0.2%) and OC= 0.6 -0.9% (mean 0.7±0.1%). As expected, the coastal zone presented the highest individual OM and OC contents in the sediments, varying between OM= 2.2 - 4.9% (mean 3.3±0.7%) and OC= 0.5 - 2.7% (mean 1.5±0.7%). Most OM in water and sediment occurs as OC that has functional groups that form stable complexes with trace metals, due to its capacity of cations adsorption. Thus the presence of OC tends to increase the dissolved fraction of metals [22]. The decomposition of OM can affect both hydrodynamic processes and geochemical redox cycles, providing driving forces for metal mobilization [23]. The mobilization and distribution of trace metals in water-sediment systems are dependent on physical-chemical, especially pH and redox changes (DO levels), and microbiological mechanisms, mediated by the species transformation [24,25], as is observed in the diagenetic processes. The physical-chemical mechanisms involved include metal speciation, adsorption, precipitation, coprecipitation and diffusion [22,26]. The capacity of sediments to adsorb and retain trace metals is



Fig. 2. A) Graphic representation of the oceanic relief (bathymetric profile) with exponential fitting curve (red line) and respective differential equations, behavior of sedimentation – deposition processes and physical-reworking patterns in the coastal zone and ACS; B) Distribution pattern of the SiO<sub>2</sub>/Al<sub>2</sub>O<sub>3</sub> ratio in the coastal zone and continental shelf with respective parabolic fitting curve (red line of tendency) and equation

	S	ediment		Pore water								
	Water depth	Porosity	Sand	Silt+Clay	00	Salinity	рН	<b>O</b> <sub>2</sub>	Fe <sup>2+</sup>	Fe <sup>3+</sup>		
	m	ml/cm <sup>3</sup>	%	%	%	(dS/m)		%	mmol/L	mmol/L		
100*	40.0	0.25	18.6	5.8	0.32	35.5	7.99	22.79	0.02	0.04		
AC3	98.5	0.40	94.2	81.4	1.78	38.4	8.21	57.60	0.03	0.05		
07	4.5	0.66	9.4	80.6	0.46	4.9	6.51	5.64	0.04	0.04		
UZ	8.3	0.70	19.4	90.6	2.70	6.0	7.84	18.63	0.06	0.07		
Innor	40.0	0.25	18.6	70.7	0.55	35.5	7.99	22.79	0.03	0.04		
Inner	42.2	0.40	29.3	81.4	0.88	37.7	8.21	38.54	0.03	0.05		
Outor	72.4	0.28	43.9	5.8	0.32	36.3	8.03	39.37	0.02	0.04		
Outer	98.5	0.34	94.2	56.1	1.78	38.4	8.18	57.60	0.02	0.05		

 Table 1. Some characteristics of the sediment and pore water determined at the ACS (minimum and maximum values by zone)

\*ACS= sites 1 to 3 plus 5; CZ= sites 4 plus from 6 to 10; Inner= 3 and 5; Outer= 1 and 2



Fig. 3. AB. medium concentration and trend of the trace metals in the CZ and ACS, blue Amazon – Brazil

also dependent of other variables as the availability of Fe and Mn, presence of carbonate and the clay minerals levels.

The accumulation of OM associated to fine fractions of clay increased the porosity of the sediments at the coastal zone  $(0.66 - 0.7 \text{ ml/cm}^3)$ , while in the outer shelf, owed mainly the uniformity of the grain size, the sandy sediments

presented the smallest porosities values  $(0.28 - 0.34 \text{ ml/cm}^3)$ , Table 1). The porosity is defined as the amount of water that a rock or sediment can store. For this reason, porosity also can interfere in the metallic ions levels and consequent mobilization and distribution in the pore water, allowing the accumulation (stock) of metals in the sediment compartment.

Table 2. Pearson's correlations: Bold correlations are significant at $p < 0.050$ and $n = 11$
(values with 3 decimals ±.xyz)

	Depth	οM	00	IC	Sand	Silt	Clay	Si/Al	DO	<b>O</b> <sub>2</sub> %	рΗ	Sal	Cr	Pb	Ni	Zn
Depth	1.0															
OM	703	1.0														
OC	476	.875	1.0													
IC	.424	718	860	1.0												
Sand	.915	715	556	.343	1.0											
Silt	737	.383	.205	202	912	1.0										
Clay	944	.776	.537	386	958	.760	1.0									
Si/Al <sup>1</sup>	.803	780	692	.085	.937	863	892	1.0								
DO	464	719	677	704	145	.082	.233	.105	1.0							
O <sub>2</sub> %	299	644	525	312	.011	125	.081	.249	.948	1.0						
pН	.686	713	754	.711	.565	429	599	.293	531	419	1.0					
Sal <sup>2</sup>	.882	830	583	.522	.849	466	709	.444	801	662	0.755	1.0				
Cr	565	.632	.601	226	715	.738	.729	840	408	307	.070	476	1.0			
Pb	258	.617	.540	507	425	.490	.437	650	592	444	.418	229	.909	1.0		
Ni	034	.783	.666	612	674	.529	.340	522	758	473	.573	474	.794	.926	1.0	
Zn	041	.732	.663	611	231	.634	.337	470	650	564	.586	409	.821	.945	.961	1.0
Hg	515	.516	.497	.013	691	.809	.624	714	324	251	.070	489	.789	.705	.627	.573
						1	SiO <sub>2</sub> /A	$l_2O_3$ ; <sup>2</sup> S	Salinity							

Table 3. Weight factor ( $F_w$ ) of the parameters analyzed in the distribution and mobility of trace elements

depth	OM	00	IC	sand	silt	clay	Si/Al	DO	<b>O</b> <sub>2</sub> %	рΗ	sal	Cr	Pb	Ni	Zn	Hg
-0.9	0.9	0.8	-0.6	-1.0	0.6	0.8	-1.0	-0.8	-0.6	0.1	-0.3	0.8	0.6	0.8	0.7	0.6
				*	Bold=	values	with sig	nificand	e at p<	0.050						

Table 4. Estimates of net production and net mineralization of organic matter from the incubation test (t= 96h) for sites 1 (outer shelf) and 4 (coastal zone)

Site	Depth	pth [CO <sub>2T</sub> ]			ΔI*	<b>F</b> i**	$\Delta NH_4^+$	Fe <sup>2+</sup> /Fe <sup>3+</sup>
	(m)	(mmol/	L)	(mmol/L)	(meq/L)	(log)	(µmol/L)	
		t <sub>o</sub>	t <sub>96</sub>					
1	0.0	3.30	4.10	1.44	1.81	-1.50	-9	0.35
1	0.5	3.50	4.80					0.54
4	0.0	3.60	6.60	4.94	3.26	-0.48	53	0.65
4	0.5	3.80	8.70					1.45

\*Activity coefficient variation for t=96h; \*\*Ionic force defined by Kielland [19]

The SiO<sub>2</sub>/Al<sub>2</sub>O<sub>3</sub> ratio followed the same trend presented for grain size, with a total variation in the study area from 3.7 (site 4) to 30.3 (site 1), and averages by zoning of  $13.3\pm10.8$  in ACS;  $4.3\pm0.5$  in the inner shelf;  $22.3\pm8.1$  in outer shelf; and  $6.7\pm1.3$  in the coastal zone. These results confirm the proportion of silicates as a function of sediment transported, with a strong distinction between the low SiO<sub>2</sub>/Al<sub>2</sub>O<sub>3</sub> ratio in the sediments deposited by the fluvial currents (Amazonas and Pará rivers) and the high ratio in the sandy sediments reworked by the marines currents (Fig. 2B). It is correct to say that the sediment characteristics found at each sampling site were influenced not only by the currents, but also by the depth of the bed. The shallower areas presented higher sedimentary deposition load, with accumulation of fine sediments. On the other hand, the deeper areas presented greater resuspension and swirling capacity, preventing the deposition of fine sediments.

#### 4.2 Distribution of Trace Elements

The trace metal contents varied between Cr 36.5 - 86.4 (71.0±13.2); Pb 54.2 - 108.0 ( $85.9\pm17.3$ ); Ni 15.8 - 34.0 ( $25.1\pm6.1$ ); and Zn 76.0 - 135.0 ( $108.0\pm21.7$ ) mg/kg and Hg 66.0 - 113.2



Fig. 4. Vertical profile of DO (μmol/L) and ΣCO<sub>2</sub> (mmol/L) according to the zoning process: A) and D) outer shelf; B) and E) inner shelf; C) and F) coastal zone



Fig. 5. Vertical profile of the Fe<sup>2+</sup> and Fe<sup>3+</sup> fractions in the pore water from sediments located in the vector 1 – 4 of the ACS, and indication of the possible transport routes of the associated trace elements

Legend: a) decomplexing; b) co-precipitation; c) fixation; d) mobilization

(77.8±11.4) ng/g (Fig. 3A). The following behaviors were observed: 1) tendency to greater fraction of fine sediments in the inner shelf by direct influence of the high sedimentary load of the Amazon River mouth; 2) absence of OM in the sediments of the outer shelf by the continuous flow of the ocean currents from southeast to northwest; 3) a strong seasonality in the volume of sediment transported due to rainfall. The concentration of trace metals in the sediments followed the same convergence observed for the sediment flow, establishing a pattern with higher levels of trace metals in the CZ and inner shelf, and lower levels in the outer shelf (Fig. 3B).

An important aspect in the flow of metals is the high capacity of metallic adsorption by the clay minerals in the sedimentary transport regime (discussed in the item 4.1). The high cation exchange capacity observed in the clay and OM particles, associated to the slightly acidic pH conditions in the fluvial-marine svstem. contributed to the storage of the trace metals especially in the inner shelf, as bivalent forms. This explains the positive correlation pattern calculated for OM, clay and trace elements. The Pearson correlation indexes calculated between OM and trace elements ranged from 0.516 Hg to 0.783 Ni (Table 2). There was also observed a significant correlation between OM and fine clay particles (0.776). In relation to the adsorption and desorption processes of the particles, the increase of the silicate load in the sediments of the outer shelf reduced the OM and clay contents and, consequently, of trace elements available for mobilization. The bioavailability of a trace metal, and consequently its toxicity, both depend on the form in which the metal is found (degree of speciation). Factors such as pH, Eh, alkalinity, degree of oxidation, suspended solids (OM and clay minerals), oxygen and temperature interfere with the mobility/precipitation ratio of trace metals. The pH has an important controlling role in the precipitation of the metallic elements through its ability to attack the minerals of rocks, soils and sediments [22], inducing the leaching or solubilization of the metals. During the leaching process, oxygen can be partially consumed to oxidize OM and reducing it to simpler inorganic fractions. Thus, the contribution of O<sub>2</sub> transferred from the cooler ocean currents to the sedimentary layer, especially in the first half meter, interferes in the OM-metals and claymetals adsorption mechanism, a tendency observed in the negative correlation between the parameters. The depth of the sedimentary bed

also showed a significant correlation, in this case with OM, sedimentary fractions, pH and salinity (Table 2).

A normalization index was applied to the absolute values in order to establish the importance or 'weight factor' ( $F_w$ ) of each parameter under the distribution and mobility of trace elements in the studied area. The normalization to establish a 'weight factor' is a practice usual in the creation of indexes such as the Water Quality Index, first applied by Brown and McClelland [27] for the National Sanitation Foundation (US NSF). Equation 8 presents the applied normalization calculation. The numerical values in front of each parameter indicate their 'weight' of importance in the general tendency, and the signal (positive or negative) of each parameter indicates if the proportionality was direct or indirect in relation to the concentration of trace elements in the sediments. The results suggest that the mobility of Cr. Ni and Zn is strongly related to the depth and location of sampling sites; sand, clay, OM and OC levels, and the DO concentration (Table 3).

$$Fw = \frac{Avr \times (SD)^2}{Max^2 - Min^2} \qquad (eq.8)$$

Where:  $F_w$ = weight factor; Avr= average; SD= standard deviation; Max= maximum and Min=minimum.

#### 4.3 Diagenetic Processes

The series of physical-chemical transformations of OM, occurring in the water and sediment compartments, including pore water, is known as diagenesis. It is an essential process for the cycling of nutrients, allowing the renewal of marine life. In the diagenesis, several redox reactions occur due to the decomposition of OM by bacterial activity (reactions of Froelich et al. [28], whose main regulating factor is the concentration of DO in the sedimentary layer, especially in pore water. In aquatic sediments it is usual to continuously produce organic compounds through decomposition, by aerobic pathways, and anaerobic as well as mineralization of OM from the breathing and fermentation processes. Add this to autochthonous organic load the volume of organic compounds carried by the streams, especially bottom streams, and deposited over an area, incorporating organic compounds into the processes mentioned above.

The relationship between the production and consumption of organic material defines the metabolism of an ecosystem, in this case benthic or sedimentary. In the euphotic zone, the processes of production exceed the mineralization in the diurnal phase  $[\Delta(CO_{2T})_b =$ n], reversing the direction of the reaction at night  $[\Delta(CO_{2T})_{b}$  = +n]. In addition to O<sub>2</sub>, physical and physical-chemical factors such as stream flow, sediment porosity, sedimentation rate, watersediment interface temperature, pH, alkalinity, cation exchange capacity (CEC), trace element concentration, OM, respiration rate and intensity of benthic biological activity determine the way diagenesis occurs, especially in the time of nutrient regeneration rate. By analyzing the sampling sites by zoning, it was possible to identify areas of low O<sub>2</sub> content, with an increase in CO<sub>2T</sub> levels. These results suggest moments of reduction of  $Fe^{3+}$  and  $Mn^{4+}$  (equations 9 and 10) [28], especially in areas with high OM content and protected from currents such as site 4 below 0.2 meters (Fig. 1). Degradation of the organic matter by the reduction of Mn4+ (eq.9) is thermodynamically more favorable than the reduction of  $Fe^{3+}$  (eq.10). However, the iron reduction pathway is considered to be more important for the mobility and/or co-precipitation of trace elements in the coastal zone and inner shelf, because the total Fe concentration in the Amazonian continental waters is about 100 times higher than the concentration of manganese. The Fe Mn concentration mean total and determined in the Amazon River waters is 1.3 -3.4 mgFe/L and 0.009 - 0.08 mgMn/L (Darwich and Aprile, unpublished data). Sequence of oxidant species observed in the sedimentary layer [28]:

 $\begin{array}{rrrr} (CH_2O)_{106}(NH_3)_{16}(H_3PO_4) & + & 236MnO_2 & + \\ 472H^+ \rightarrow 236Mn^{2^+} + & 106CO_2 & + & 8N_2 & + & H_3PO_4 \\ + & 366H_2O & (9) \end{array}$ 

$$(CH_2O)_{106}(NH_3)_{16}(H_3PO_4) + 212Fe(OH)_3 + 636H^+ \rightarrow 212Fe^{2+} + 106CO_2 + 16NH_3 + H_3PO_4 + 636H_2O$$
 (10)

Although in the coastal zone and inner shelf areas occurs large deposition of OM, factors such as slightly alkaline pH and low interstitial  $O_2$  levels may be contributing to the reduction of mobility of trace elements. Considering that part of the DO content is being consumed by the OM oxidation, the process of deposition and reduction of the metallic mobility intensifies, ensuring that this sedimentary layer acts preferentially as a storage compartment (stock)

of trace elements. Vertical profiles of the O<sub>2</sub> and  $\Sigma CO_2$  contents in the pore water of the sedimentary layer of the sampled sites were elaborated (Fig. 4). In general, the sandy sedimentary layer had a greater flow of oxygenation than the mud layers of the coastal zone and inner shelf. DO contents in the 0.0 -0.5 m ranged from 95.6 - 178.1 µmol/L on the outer shelf; of 66.5 - 112.5 µmol/L on the inner shelf; and 3.1 - 54.4 µmol/L in the coastal zone (Fig. 4). The bottom currents in the outer shelf, cooler and oxygenated, they interfered in the sedimentary O<sub>2</sub> flux, facilitated by the porosity of the sandy sediments. It can be pointed out that the diffuse oxygen flux in the sediments was directed by the consumption in the oxidation process. The oxidation of organic matter in fluvial-marines sediments is considered the most important form of respiration in fluvial-marine sediments [29]. Besides, another factor may be contributing to diffuse oxygen flow, in this case a biological action involving benthic respiration, especially in coastal sediments. Studies on oxygen flow and oxireduction of nitrogenous and sulfated forms (NO<sub>3</sub> and SO<sub>4</sub>) performed in the same region demonstrated the importance of the benthic respiration in diagenetic processes [29-31]. In general, aerobic respiration takes place in the oxic surface layer and is followed by nitrate and sulfate reductions. However, in coastal sediments the oxic zone often is only a few millimeters thick, as suggest the studies of Aller et al. [12] and Froelich et al. [28], and exactly as observed in the sediments of the coastal zone and inner shelf of the ACS, where the anaerobic respiration becomes dominant. Besides, much of the oxygen uptake is used to reoxidate the products of anaerobic respiration as  $H_2S$ ,  $NH_4^+$  and  $CH_4$  at the oxic/anoxic interface of the sediment [32].

The mineralization of the anaerobic organic matter involves several oxidation processes, consuming nitrates, Fe and Mn oxides, sulfates and carbonic acid to form CO<sub>2</sub> and fermentation, in this case with formation of CH4 (methanogenesis). Due to its greater stability in relation to H<sub>2</sub>S and SO<sub>4</sub><sup>2-</sup>, the CO<sub>2</sub> becomes an important indicator of diagenetic processes. In general, is considerate for analysis the sum of all dissolved carbonate forms, which are defined as total inorganic carbon or  $CO_{2T}$ . The concentration oscillation of both free  $CO_2$  and  $CO_{2T}$  depends directly on the production and consumption in the processes already mentioned, pH, alkalinity of the carbonates and ionic composition of the water. The carbonate content, which interferes

with pH and alkalinity, has as main source in the marine environments the contribution carried by the waterways and biogenic production, whose main raw materials are plankton and mollusks. In relation to carbonate content, in the Amazonian plain the main source of carbonate is The predominantly HCO<sub>3</sub> [33]. biogenic production of carbonate in shallow waters and its dissolution in deep waters causes a dynamic equilibrium system, affected by ocean currents, and that controls, among other factors, the concentration of total inorganic carbon in the system. The CO<sub>2T</sub> contents presented some variability as a function of zonation, ranging from 3.3 - 4.3 mmol/L in the outer shelf; of 3.3 - 4.7mmol/L on the inner shelf; and 3.2 - 7.8 mmol/L in coastal zone sediments (Fig. 4). The higher CO<sub>2</sub> levels in the sediments are associated with decomposition processes and complete oxidation of organic matter and mineralization partially anaerobic of the sedimentary particles. All the sampling sites showed a considerable capacity for a rapid change in pore water constituents associated with the remineralization suggesting the presence processes, of abundant organic compounds, especially in the inner shelf that, as mentioned above, act primarily as a storage compartment of thin sediments, depending on their morphology and direction. At some sampling sites. flow decay of  $O_2$  occurred the exponential simultaneously to CO<sub>2</sub> production, especially at depths of 0.0 - 0.2 meters in the sedimentary layer. This behavior reflects the nature of particle deposition and reworking of particles. It should be remembered that the samplings and measurements were carried out during the months of the lowest flow in the Amazon and Pará rivers and at the ACS. Thus, it is plausible to believe that seasonality can alter the flow pattern of oxygen as well as CO2 during the hydrological year, especially during the periods of higher flow of the Amazon River, when the sediment load transported can reach double volume, as has been suggested by Meade et al. [10] and Gibbs [13].

# 4.4 Sedimentary Incubation

Incubation tests can be classified into open and closed systems. In the open system the loss of  $CO_2$  by diffusion is allowed, which does not return to the system. However, the closed system is often questioned by 'imprisoning' the gases, preventing the changes at the water-sediment interface. The fact is that both tests

have positive and negative points, and it must be assumed that there is an imprecision in the results. The great trump of incubation tests, however, is the ability to quantitatively estimate the processes involved. It is based on this perspective that a closed test was applied to estimate the net productivity and mineralization content of OM in the coastal zone and ACS.

The variation of the  $CO_{2T}$  and  $O_2$  in the waters  $(\Delta CO_{2T}(t) \text{ and } \Delta O_2(t))$ , including the pore waters, is controlled not only by the biological processes but also by the diffusion rate of the gases in the interface zone  $(\Delta(CO_{2T})_a \text{ and } \Delta(O_2)_a)$ , and the facilitation or not of the gas flow as a function of the degree of sedimentary porosity. Thus, the study of the diagenetic processes involving the production and mineralization from the CO<sub>2</sub> and O<sub>2</sub> variations in the water should be understood as an estimate, since there are several factors controlling the concentration and flow of these gases in the sediments, including hydrodynamics and seasonality. The closed incubation technique attempts to minimize the problem of gas exchange. Despite this, incubation reduces the flow of biogenic elements. The system's buffering capacity is of great relevance to the results, since any change in pH and alkalinity, especially alkalinity of carbonate (Alc<sub>c</sub>), interferes with the concentrations of  $H_2CO_3$ ,  $HCO_3^-$  and  $CO_3^{2-}$ , and consequently with CO<sub>2T</sub> levels. It is tried to minimize this variation, considering that the chemical reactions and the diffusions present kinetics of the same order. Thus, when CO<sub>2</sub> adsorption occurs, a part of the CO<sub>2</sub> that enters the system is transformed into carbonates, and when the CO<sub>2</sub> desorption occurs, a fraction of carbonates from the system is converted into CO<sub>2</sub>. Incubation tests confirmed that the range of 0.0 - 0.2 m is essentially the most active from the point of view of sediment reworking by the flow and inflow of the dissolved gases. As expected, CO2T rates increased rapidly during the incubation period. Increasing the concentration of inorganic carbon forms may mean an increase in aerobic decomposition rates (complete decomposition), due to the increase in the supply of reactive organic compounds. Generally, the saturation of mineral carbonates and other forms of inorganic carbon are guite high in the sediments from ACS, as were observed by Aller et al. [2], Aller et al. [12].

In the comparison of the results between the incubation of sediments from site 1, sandy and

under strong influence by marine currents, to site 4, mud and located in protected area with marked sedimentation, the amplitude of variation for the incubation period (t\_0  $\rightarrow$  t\_{96}) was much higher at site 4. The  $\Delta CO_{2T}$  determined were respectively; site 1= 1.44 mmol/L and site 4= 4.94 mmol/L (Table 4), suggesting that the net mineralization exceeded the net production of organic matter to  $[\Delta(CO_{2T})_b > 0 \text{ for } \Delta(O_2)_b < 0].$ The coefficient of activity (I) also showed a greater amplitude of variation in site 4 ( $\Delta I$ = 3.26 meg/L) compared to site 1 ( $\Delta I$ = 1.81 meg/L). The activity coefficient is a function not only of the chemical species present, but also of the ionic interactions in pore water. In silica sediments, the ionic interactions are weak and easily disrupted by the current underflow, which eventually penetrates the sediments due to their high porosity. Already in silt-clayey and clay-silty sediments, the low porosity (<0.7 ml/cm<sup>3</sup>) reduces the mobility of the mineral elements, accentuating the ionic interactions, which present strong connections, especially cationic, with the negative external surface of the particles of clay and organic matter. The ionic forces (F<sub>i</sub>), determined as a function of their logarithms, followed the same trends, being greater in the mud sediments. These behaviors are confirmed in the CEC analysis. Another important aspect is the variation rate in the productivity of  $NH_4^+$  ions (Table 4), which showed influx in sandy sediments (site 1 = -9umol/L), and high efflux for mud sediments (site  $4 = 53 \,\mu mol/L$ ).

Extrapolating the results of the incubation test to the study area, it is possible to establish some hypotheses: 1) the flow of marine currents at site 1 may be allowing the aerobic oxidation in these sedimentary layers, with the nitrification route more pronounced than the ammonification rote; 2) in the region of the coastal zone and inner shelf the routes of oxidation and reduction may be alternating according to the physical, physicalchemical and seasonal factors; 3) in the coastal zone and inner shelf the net mineralization rate has exceeded the net production rate of organic matter, since the calculations show  $\Delta(CO_{2T}) > 0$ for  $\Delta(O_2) < 0$  (Table 4) for the vector from site 1 to site 4 (Fig. 1). It should be considered, however, that the rate of sedimentation or continuous contribution of sediments from the Amazon River to the inner shelf is immense, as already discussed.

Daily fluctuations in the rate of  $NH_4^+$  production at the same site, with changes in the axis of

orientation of influx - efflux (nitrification/ ammonification) are not uncommon, having been observed even in incubation tests [29,30]. In specific case of the sediments located in the axis of orientation of the vector shown in Fig. 1, it is believed that the great difference in porosity and OM contents of the sediments, associated to the depth of the layer, were the predominant factors in the change of the orientation axis of inflowefflux. This was confirmed by determination of iron fractions. Most of the time  $Fe^{2^+}/Fe^{3^+}$  ratio was <1.0 throughout the sedimentary profile (Fig. 5), suggesting a mild to moderate oxidation pattern in the first 0.4 m, especially for outer shelf sites (sampling sites 1 and 2). A sensitive reduction in the degree of oxidation of the ferric ions was observed in sites 2 and 3 from 0.4 m. However, site 4 (Silt + Clay> 80%) presented higher iron ratio, with  $Fe^{2+}/Fe^{3+} > 1.0$ , indicating a predominance of the maintenance of the reduced form of iron (Fe<sup>2+</sup>), which by electric affinity remains in the sedimentary layer adsorbed the particles of clay minerals and organic compounds not totally mineralized. Metal profiles sensitive to oxidation variation, such as Fe and Mn, may indicate redox potential changes within the sedimentary layers, being influenced by the concentration of DO in the pore water. The pore water column can define alternate zones of oxidation and reduction of chemical species as a function of depth, according to sedimentation rates. oxvaen demand. nutrients and microbiological processes [34]. The processes of dissolution and precipitation of minerals can affect the nutrient concentration (C, N and P) and trace elements in the sediments [8], because these chemical species may be adsorbed by the surface of minerals as iron oxides and hydroxides. The behavior of the trace elements in the bottom sediments is strongly associated with organic matter rates; pH (increase of the pH implies increase of the metallic adsorption); O<sub>2</sub> content; redox potential; seasonality (flow of alternating currents); and diagenetic processes, which occur in the fluvial-marine system. In this case, three routes of adsorption of trace elements to the iron in the sediments and pore water can be established, taking into account the depth and oxygen content: 1) adsorption in Fe<sup>3+</sup> oxides and hydroxides, especially at the watersediment interface and with increased metal mobility; 2) dissolution of Fe<sup>2+</sup> oxides and hydroxides in the layers of less interference of O<sub>2</sub>, in this case the trace elements may follow the oxidative route towards the interface, or to stay associated with reduced forms (S2-) and follow the co-precipitation way; and 3) fixation on the mineral surface or precipitation in the autigenic mineral phase, in or near the anoxic environment (Fig. 5). Reactive trace elements such as Cu, Ni and Zn may have their concentrations controlled by co-precipitation with Fe sulfides, which would result in the reduction of the metallic mobility of these elements. Comparing the trace element contents with the vertical distribution profile of the iron fractions, it was possible to evidence an increase in trace element concentrations in Fe reduction zones, indicating processes of desorption of iron oxides and hydroxides, and mineralization of organic matter.

# 5. CONCLUSION

There is a strong zonation in the sedimentary transport and deposition processes, influenced by the granulometry and texture of the sediments, and the fluvial-marine currents, which alternate in direction seasonality. Thus, it was possible to identify that the sediments of the coastal zone and inner shelf are predominantly silt-clayey and clay-silty as opposed to sandy and sand-silt sediments of the outer shelf. The trace elements presented a distribution and concentration pattern equivalent to that of the sedimentary distribution pattern, with higher metallic contents in the deposits of clav and organic matter of the coastal zone and inner shelf. The determination of a weight factor  $(F_w)$ calculated to estimate the degree of importance each physical and physical-chemical of parameter in the distribution and mobility of trace elements showed that these elements were strongly related to the depth and location of sampling sites, sand, clay, OM and OC levels, and DO concentration, especially to Cr, Ni and Zn metals. The vertical distribution of  $O_2$  and  $CO_2$ and the  $Fe^{2+}/Fe^{3+}$  ratio in the pore water suggest a predominance of organic matter oxidation in the sedimentary layer between 0.0 and 0.2 m, with partially anaerobic mineralization of the mud sediments below 0.4 m, especially in the coastal zone and inner shelf. However, the results indicate that there may be verv significant differences between aerobic and anaerobic degradation of organic matter. Increases in trace element concentrations were observed in Fe reduction zones. indicating processes of desorption of Fe oxides and hydroxides and net mineralization of OM. The results of the incubation test indicated that  $\Delta(CO_{2T}) > 0$  for  $\Delta(O_2) < 0$ , suggesting that especially for the coastal zone and inner shelf the net mineralization rate has

exceeded the net production rate of organic matter.

#### **COMPETING INTERESTS**

Authors have declared that no competing interests exist.

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