BIOGEOCHEMICAL PROCESSES IN AMAZON SHELF SEDIMENTS

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Each year the Amazon River delivers approximately 1 billion metric tons of sediment to the equatorial Atlantic (Meade et al., 1985). A portion of this debris is highly weathered and contains abundant reactive Fe, Mn, Al, and Si oxides. The associated terrestrial organic matter is biologically refractory (Hedges et al., 1986). Upon entering the Amazon Shelf region, particles are exposed to seawater brine, mixed with labile planktonic organic matter, and incorporated into the seabed where a variety of geochemical reactions take place. Major research efforts within the AmasSeds Project include elucidating the types and rates of these diagenetic reactions (particularly those associated with the decomposition of organic matter), their governing factors, and their influence on the properties of overlying water and preserved deposits.

Modern views of early diagenetic processes in shallow marine sediments have largely derived from studies of temperate estuaries and shelves, particularly along the coasts of North America and Europe. In these environments, sedimentary organic matter is decomposed through the general sequential series of biogeochemical oxidants (O₂, NO₃⁻, Mn, Fe, SO₄²⁻, CO₂), although reactions are typically dominated over large vertical intervals by sulfate reduction or methanogenesis. Oxygen is often relegated to a largely indirect role as an oxidant of anaerobic metabolites. Other potential oxidants of organic matter such as NO₃⁻, Mn and Fe oxides, are of minor importance (Jorgensen, 1982; Chanton and Martens, 1987; Mackin and Swider, 1989). The chemical properties of sediments and patterns of authigenic mineral formation in deposits reflect these reaction paths, as well as diagenetic transport regimes strongly influenced by the bioturbation activities of benthic organisms (Berner, 1980). In contrast, early studies of Amazon Shelf sediments demonstrated extensive zones of suboxic reactions associated with Fe,Mn reduction and oxidation cycles, and little evidence of macrobenthic activity (Aller et al., 1986; Aller and Aller, 1986). Unlike other shelf deposits of similar organic content (0.6 ± 0.1% C), sulfate depletion is limited, dissolved sulfide is undetectable, and sulfide minerals are minor in the upper few meters (total reduced S is typically <0.1%). High concentrations of dissolved Fe⁺⁺ and Mn⁺⁺ in pore water (0.1–1 mM) are found throughout the upper 0.5–2 m of Amazon deposits compared with vertical intervals 0.01–0.05-m thick in many shelves. These suboxic zones extend across the Amazon Shelf for ~100 km, encompassing an area of at least 40,000 km² off Brazil alone. Because Fe and Mn oxides and their reduction products are carrier phases for numerous minor and trace elements, their redox cycling can critically influence oceanic chemistry. The input of highly weathered debris (ferricrete, laterite) and massive physical mobility/working of sediments are apparently critical factors determining these patterns.

Seabed geochemistry and associated benthic biological studies during the AmasSeds project (see introductory article by Nittrouer et al., this issue a) are designed to document large-scale diagenetic processes on the Amazon Shelf, to exploit unique biogeochemical and physical features for study of modern elemental cycles, and to provide insights into ancient sedimentary environments. Specifically, the objectives are to: (1) document seasonal variation in geochemical properties and reaction rates related to carbon and metal cycling in the upper 1–2 m of the seabed; (2) elucidate the major physical, chemical, and biological factors controlling diagenetic and benthic community patterns; (3) determine the source and reactivity of organic carbon driving seabed reactions; (4) examine the formation of authigenic minerals including Fe carbonates, phosphates, sulfides, and silicates; (5) estimate the impact of seabed dia-

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**Sampling Approach and Methods**

Sampling sites on the shelf were chosen to promote integration between AmasSeds research groups, thereby constraining interpretations and maximizing implications of measurements. Seasonal studies of diagenesis and sediment-water interactions are continuing at three sites along two shore-perpendicular transects: 1) off the river mouth (RMT), and 2) a more northern Open-Shelf Transect (OST) (Fig. 1). In addition, a transect (ST) south of RMT has been sampled for sediment-water solute exchange and benthic communities; and, a transect (NT) north of OST was sampled for benthos. These transects encompass a variety of major sedimentation and biological regimes. Sampling to date has taken place in August, February, and May during periods typical of descending, ascending, and peak flow periods of the Amazon River. These months also include periods of varying wind stress and influence of the North Brazilian Coastal Current (Nittrouer et al., this issue; Geyer et al., this issue). A future cruise during minimum river flow (October) will complete the temporal sampling plan.

Geochemical and benthic biological sampling along the two major transects have included collection of: one 0.25-m$^2$ box core, two 0.1-m$^2$ box cores, and one or two $\sim$ 3 m long kasten cores at each site. Plankton samples also were obtained to determine availability of colonizing benthic species. Subcores from the 0.25-m$^2$ box core were incubated aboard ship for O$_2$ nutrient, and dissolved Fe and Mn diffusive flux estimates. In situ incubations using benthic landers are precluded by physical conditions on the shelf. Box cores and kasten cores were sampled over $\sim$ 1–5 cm and 10 cm depth intervals, respectively, and pore waters were separated under anoxic conditions. Nutrients and CO$_2$ were analyzed aboard ship. Sediment incubation experiments were used to estimate selected reaction rates for Mn, Fe, SO$_4^-$ reduction, and for CO$_2$, NH$_4^+$, and HPO$_4^{2-}$ production as a function of depth in the cores. These measurements help constrain interpretations of pore-water distributions and transport processes. Experiments were set-up to determine the composition of authigenic mineral precipitates and are in progress. Pore-water samples were stored for C isotopic analyses. Core samples have been archived and frozen for solid-phase analyses, including organic geochemical and authigenic-mineral studies.

**Oxidation-Reduction Reaction Rates and Balances in the Seabed**

One common measure of decomposition rate and benthic metabolism is the flux of O$_2$ to the seafloor. If there is little storage of reduced products of decomposition in a deposit, O$_2$ flux can correspond closely to total C oxidation. The redox reactivity of Amazon bottom sediments, as measured by O$_2$ diffusive uptake, is relatively uniform and generally comparable to other, much colder temperate shelves. The yearly average of 18 samples is 11.8 ± 3.8 mmol m$^{-2}$ d$^{-1}$, with only a hint of seasonal variation (May: 10.5 ± 2.8; Aug: 13.2 ± 4.3) (Fig. 2). Oxygen measured in sediments by use of microelectrodes, is absent below 1–4 mm at all sites. This agrees with penetration depths estimated from a 1-dimensional diffusion model, the measured O$_2$ flux, and bottom-water oxygen concentrations. The average ratio of diffusive fluxes estimated from pore-water concentration gradients and of directly measured (incubation) O$_2$ fluxes is 1.07 ± 0.19 (N = 6), also in excellent agreement.

An additional, more general measure of total remineralization is the production of CO$_2$. Directly measured production rates of total CO$_2$ in the upper 1–1.5 m of sediment demonstrate that there is no dramatic spatial variation in decomposition rates within the sampled region (Fig. 3).
This is consistent with a lack of spatial pattern in O₂ fluxes measured during AmasSeds and in NH₄ production rates previously measured for Amazon surface sediments (Mackin et al., 1988). There is a remarkably small attenuation of reaction rates with depth in deposits. This is expected for sediments that are subject to rapid sedimentation and reworking. The depth-integrated production of CO₂ indicates that \( \sim 55 \pm 10 \) mmol C m\(^{-2}\) d\(^{-1}\) is actually decomposed in the upper \( \sim 1.5 \) m as compared to an O₂ diffusive flux of \( \sim 12 \pm 0.4 \) mmol m\(^{-2}\) d\(^{-1}\). Thus the diffusive flux of O₂ can account for a maximum of \( \sim 20\% \) of total C oxidation in these sediments and at least 80\% of C decomposition in the seabed is anaerobic.

Most of this anaerobic oxidation must be through metal reduction because there is little evidence of SO₄ depletion in the same zones. This imbalance also means that most reoxidation of anaerobic decomposition products occurs in the water column during resuspension/erosion events or during bioturbation not accounted for in core samples, or, that reduced products such as FeCO₃, MnCO₃, and FeS₂ are stored in the sediments and not reoxidized. These possibilities are not mutually exclusive and, in the latter case, require that authigenic minerals constitute only a small fraction of the sediment by weight. The integrated CO₂ production corresponds roughly to 25\% of the estimated average net primary production in overlying water (DeMaster et al., this issue), a percentage typical of many shallow marine environments.

**Source of Reactive Organic Matter**

Organic matter in Amazon deltaic sediments is composed of both terrestrial and marine-derived material. In the study area, \( \sim 70-90\% \) of the sediment carbon is probably terrestrial (Showers and Angle, 1986). Because of the refractory nature of Amazon River organic material (Hedges et al., 1986; Richey et al., 1990), marine organic matter was expected to be the reactive reductant driving many of the diagenetic reactions in the seabed. The sediment incubation experiments mentioned previously indicate that the N/C ratio of dissolved decomposition products released in the upper 10-20 cm is near the mean Redfield value for plankton, implying a marine source. In addition to analysis of nutrient-release stoichiometry, the carbon isotopic composition of pore-water dissolved inorganic carbon (DIC) has been examined directly for clues of the origin and fate of decomposing organic matter. For the river, \( \delta^{13}C \) values for organic C typically are \( \sim -27 \) to \(-28\%\), and for outer-shelf suspended matter are \( \sim -19.5\%\) (Hedges et al., 1986; Showers and Angle, 1986). Preliminary \( \delta^{13}C \) measurements suggest that the remineralized carbon in the sediments is isotopically marine (\( \delta^{13}C \sim -20\%\); Fig. 4). Any authigenic carbonates formed should have this isotopic composition despite the dominance of terrestrial organic carbon in the sediment. This provides a means of demonstrating a marine origin for any preserved deposits.

**Spatial and Temporal Variation in Pore Water Chemistry**

As found previously, SO₄ depletion in pore water usually begins below \( \sim 0.5-1 \) m and underlies a zone of elevated Fe\(^{2+}\), Mn\(^{2+}\) concentrations at all stations. Metabolite build-up (e.g., NH₄, CO₂) is generally higher in pore waters along the OST topset region compared to the RMT sites, despite the fact that reaction rates are not greatly different (Fig. 3). Relatively low DIC concentrations...
In some locations, dramatic seasonal variability in solute profiles is observed. Tensions (<5 mM) and somewhat variable δ¹³C values over the upper 1.5 m along RMT are consistent with frequent reworking of the seabed (see: Kineke et al., this issue; Nittrouer et al., this issue b). Preliminary modeling suggests that in some cases pore-water solute profiles over the upper ~1 m are only about 1–2 months old due to physical disturbance or extensive biological reworking. In some locations, dramatic seasonal variability in solute profiles also is observed, with lowest concentrations generally found during ascending river flow (Fig. 5). Thalassinid shrimp (Callianassa sp. and Upogebia sp.) are abundant at the innermost stations of RMT and may be at least partly responsible for the apparent, massive sediment-water exchange. Deep burrowing sipunculids, brittle stars (ophiuroids), and gobid fish are also present at these stations. Solute exchange is apparently less extensive along OST, but rapid transport takes place in at least the upper 20–60 cm, due in some instances to physically mobile fluid-mud layers.

DIC reaches >20 mM at OST stations and experimental incubations indicate that authigenic carbonate precipitation occurs in this region.

**Benthic Communities and Bioturbation Potential**

The benthic community distributions mirror the physical regime and provide insight into diageneric transport processes and timescales of physical disturbance within deposits. There are marked increases from inshore to offshore and from north to south in macro- and meiofaunal abundances, species richness, and functional group diversities (Fig. 6). Animals between 0.5 to 1 mm in size, dominate the bottom community. As in most sedimentary deposits, highest abundances occur near the sediment-water interface and decrease with depth in the deposit. Polychaete worms dominate the fauna and include surface deposit feeders such as early colonizing spionids, as well as deep deposit-feeding species. Other bur-
rowing organisms on the shelf include mud shrimps (thalassinids), brittle stars, lumbrinerid and cossurid polychaetes, sipunculids, and gobiid fish; which are found to depths of 25 cm below the surface. Large numbers of crab and shrimp larvae, as well as meiofaunal-size crustaceans (from 0.5 to 0.062 mm) including copepods and cumaceans, are also found to >25 cm, presumably having been passively transported downward by sediment reworking.

Macrofauna reach largest size and greatest abundances during ascending river flow at all shelf stations (max = <4000 m² at ST2, depth = 47 m). Inshore of 15 m, for example, abundances are 20 to 50 times greater than during descending flow. This is the time when biogeochemical properties of the sediment show evidence of recent major exchange, as mentioned previously. Model calculations indicate that burrow irrigation by the numbers and sizes of organisms present could be a significant exchange process in some locations. Also during this time, larval crustaceans of many groups are present in tens of thousands per m². Some crustaceans (both adults and larvae) remain during peak flow periods, but have a patchy distribution. Presumably, there is a relatively constant supply of larvae and adult organisms from the south, in the near-bottom waters associated with the North Brazilian Current. These organisms may settle out of the water column when overall current speeds decrease. The lack of any evidence of sexual maturity among both adult macro- and meiofaunal organisms (except for Upogebia sp.) suggests that very few of the organisms reproduce in these physically disturbed sediments.

The Present and Past Shelf

Seasonal variation in sediment transport/re-mobilization is apparently the major cause of short-term change in diagenetic and biological properties of the upper 1–2 m of the present shelf deposits. The relative importance of organic-matter decomposition pathways and the nature of benthic communities and sedimentation regime may have changed over longer timescales on the Amazon shelf. For example, methane concentrations (<40 μM) in the upper 2–3 m of sediment indicate that methanogenesis is not an important mode of organic matter diagenesis in modern Amazon shelf sediments and appears to be restricted to microenvironments or narrow zones. This is in contrast to greater depths within the seabed where seismic profiling and piston-core analysis demonstrate gas-charged sediment (Figueiredo et al., 1990). This gas is biogenic (δ¹³C = −84‰) and could result from the slow degradation of low-reactivity organic matter over a 1000-year timeframe. Alternatively, these methane deposits could have formed during a period when the flux of marine carbon to the seabed was much greater than today and perhaps sediment reworking was less intense. Sampling of deeper deposits should resolve this issue.

Future Research

This article represents a preliminary outline of some of our work on seabed diagenesis and benthic communities as part of the AmasSeds Project and of the basic research questions to be...
addressed. There remain a variety of major unknowns regarding biogeochemical processes within Amazon Shelf deposits, including the timing and relative importance of physical and biological transport events in determining diagenetic properties at different locations. There are major classes of reactions occurring which deplete pore water K⁺, Mg⁺⁺, Ca⁺⁺, Fe⁺⁺, Mn⁺⁺ and Si(OH)₄, the characterization of which has only begun. Also, it is not clear how and when benthos are recruited or immigrate to this dynamic region, as no evidence of sexually mature fauna has been found as yet. Numerous new questions have arisen in addition to those originally posed. Some will be resolved by additional sampling of low river flow conditions (and lower wind stress), and by deeper sampling of sediment deposits using piston cores. In addition, completion of sample analyses and experiments from earlier cruises, modeling, and interdisciplinary information exchange should further constrain interpretations.

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