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Benthic geochemistry of manganese in the Bay of Biscay, and sediment mass accumulation rate

Aurélia Mouret · Pierre Anschutz · Pascal Lecroart · Gwénaëlle Chaillou · Christelle Hyacinthe · Jonathan Deborde · Frans J. Jorissen · Bruno Deflandre · Sabine Schmidt · Jean-Marie Jouanneau

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Abstract Manganese is a major redox reactive element of benthic metabolism. We have built a database of existing knowledge on the benthic geochemistry of Mn in the Bay of Biscay, in order to comprehensively assess the behaviour of Mn in a variety of environments during early diagenesis. The database contains vertical profiles of particulate and dissolved Mn species of 59 cores collected during 17 cruises between 1997 and 2006 at nine stations positioned

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A. Mouret · P. Anschutz () · P. Lecroart · G. Chaillou · J. Deborde · B. Deflandre · S. Schmidt · J.-M. Jouanneau Université de Bordeaux, CNRS, UMR 5805-EPOC, 33405 Talence, France e-mail: p.anschutz@epoc.u-bordeaux1.fr

G. Chaillou

Département de Biologie, Chimie et Géographie, Université du Québec à Rimouski, G5L 3A1 Rimouski, QC, Canada

C. Hyacinthe

Department of Marine Sciences, University of Georgia, Athens, GA 30602, USA

F. J. Jorissen

Laboratoire d'Etude des Bio-indicateurs Actuels et Fossiles (BIAF) UPRES EA 2644, Université d'Angers, 49045 Angers Cedex, France

B. Deflandre

Université Paris 7, Laboratoire de Géochimie des Eaux, IPGP, 75251 Paris, France

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between 140 and 4,800 m water depths. At all studied stations, Mn species follow the conventional distribution, where Mn(III,IV) species are enriched in the oxic layer, and dissolved Mn is present in the anoxic sediments. A minor part of Mn-oxides originates from sedimenting particles. The major part is of diagenetic origin, and derives from the oxidation of upward-diffusing dissolved Mn(II). Mn-oxide inventories are higher at the deeper stations than at the shallower ones. This difference cannot be attributed to different sources of sedimenting particles, but it must depend on sedimentation rate and diagenetic processes. At depth, dissolved Mn(II) concentrations are constant. This probably reflects equilibrium with an authigenic Mn(II) phase, which is the ultimate phase into which Mn is fossilized. The Mn content of deeper anoxic sediments is similarly low in all the cores studied, associated with corresponding trends of Mn content in sedimenting particles of the Bay of Biscay. Bioturbation, rather than redox oscillations, can convey Mn(III,IV) species downwards into the anoxic sediments where they are reduced, associated with a peak of dissolved Mn. Because dissolved Mn(II) is re-oxidized when it diffuses towards the oxic layer, the inventory of the diagenetic Mn(III,IV) phase remains at steady state, especially at stations where the oxic layer is thick. It then becomes possible to calculate the residence time of diagenetic Mn(III,IV) particles within the oxic layer, using the upward-directed flux of pore water Mn(II). By applying this residence time to the accumulation of sediments within the oxic layer, we obtain the sediment mass accumulation rate. The values calculated for the sediments of the Bay of Biscay fit well with accumulation rates obtained from radionuclides or sediment traps. The method has also been validated with data collected in other marine sedimentary environments.



Introduction

Manganese is the most abundant transition metal in natural environments, after iron. Manganese oxides and oxyhydroxides represent significant oxidants for organic carbon in marine sediments (Aller 1990, 1994; Canfield et al. 1993; Thamdrup et al. 1994). In marine oxygenated environments, manganese is present in the form of Mn(IV) oxides and Mn(III) oxyhydroxides (Post 1999). These oxidized forms of Mn are very reactive and have a strong capacity for the adsorption of trace metals (Murray 1975; Stumm and Morgan 1996). Pore water Mn(II) is produced upon Mn-oxide reduction, which is coupled either directly to organic carbon oxidation (Froehlich et al. 1979; Myers and Nealson 1988a) or indirectly via the oxidation of reduced solutes like Fe²⁺, sulphide, ammonia or organic acids (Stone 1987; Myers and Nealson 1988b; Burdige et al. 1992; Luther et al. 1997; Hulth et al. 1999; Anschutz et al. 2000; Schippers and Jørgensen 2001; Hyacinthe et al. 2001; Luther and Popp 2002).

Marine sediments commonly contain an oxic surface layer enriched in insoluble Mn(III,IV) phases. Deeper, these oxides are reduced to soluble Mn(II) (Burdige 1993). Dissolved Mn(II) can diffuse upwards in pore waters across the oxic-anoxic boundary, and sometimes across the sediment-water interface, where it precipitates again in the form of Mn(III,IV) (Sundby and Silverberg 1985). Precipitation and adsorption on particles of upward-diffusing Mn(II) is a dominant phenomenon that enriches oxidized surface sediment in manganese (Sundby et al. 1981; van der Zee et al. 2001). Vertical profiles of dissolved manganese in pore waters often display a concentration maximum immediately below the oxic front, with a downward-directed gradient below the maximum. Precipitation of mixed Ca-Mn carbonates, driven by increased alkalinity production, traps Mn(II) in anoxic sediments (Middleburg et al. 1987; Mucci 1988, 2004; Jakobsen and Postma 1989).

Transient-state diagenesis is the rule for most continental margin sediments (Sundby 2006). For manganese species, non-steady-state diagenesis can be recognized on single vertical profiles when, for example, the depth where upward-diffusing Mn(II) is oxidized does not correspond to the depth of maximum enrichment of Mn(III,IV) phases. Bioturbation, or seasonal redox oscillations due to episodic organic matter sedimentation are the major causes of shortterm (days-months) transient profiles (Gehlen et al. 1997; Gobeil et al. 1997; Anschutz et al. 2000). Seasonality in the geochemical composition of suspended matter, especially in coastal environments, may also modify vertical Mn distribution (Dellwig et al. 2007). Changes in bottom water oxygen concentrations exert a strong control on the redox environment within surface sediments, and can induce the migration of redox boundaries and reaction fronts (Sundby 2006). Bioirrigation through burrows is also a significant process that can affect the flux of dissolved Mn(II) in coastal environments (Aller and Aller 1998).

Moreover, the transient- or steady-state behaviour of Mn in modern marine sediments depends on the timescale of observation. At a multi-annual timescale, steady-state diagenesis may be assumed for Mn species, particularly when bioturbation activity is moderate. At steady state, the flux of buried Mn corresponds to the mean flux of sedimenting Mn. Enrichment of Mn-oxide in the oxic layer is of diagenetic origin. Therefore, at steady state, the level of diagenetic Mn-oxide must be maintained constant, controlled by the flux of Mn to the sediment and by benthic metabolic processes. In other words, at steady state the residence time, or mean lifetime, of Mn-oxides within the enriched surface sediment layer must be constant. Consequently, the distribution of Mn species and estimation of residence time may be used as an independent method to estimate sediment mass accumulation rate (MAR). This concept needs to be examined by studying the distribution of the different phases of Mn in numerous sediment cores.

We have built a database that collates published information (Hyacinthe et al. 2001; Anschutz et al. 2002; Chaillou et al. 2002, 2003, 2006; Fontanier et al. 2002, 2003, 2005, 2006) and non-published data of vertical benthic profiles of dissolved and particulate manganese, and major diagenetic parameters in several fine-grained sedimentary environments of the Bay of Biscay. Also included are sporadic measurements of physicochemical properties of the water column and suspended particles. This unique database gives us the possibility to (1) compare the distributions of Mn species in different sedimentary environments, (2) examine the seasonal–annual variability of Mn-species distribution and (3) relate enrichment levels of Mn species to sediment MAR for this bay, as well as (4) strengthen universal relationships between reactive Mn species, an aspect otherwise not feasible if based on only a few cores.

Materials and methods

Sampling

The Bay of Biscay is located on the eastern side of the Northern Atlantic Ocean, influenced by oceanic waters from the North Atlantic Drift. Several rivers in France and Spain (the Loire, Charente, Gironde, Adour, Bidassoa and Nervion) feed the sedimentary basin. The Gironde and Loire rivers are presently the main sources of fine sediments to the continental margin (Jouanneau et al. 1998b).

The Bay of Biscay database consists of 104 studied cores, for which vertical profiles of redox-sensitive properties have



been measured. The distribution of manganese was studied in 59 undisturbed cores collected at nine stations between 140 and 4,825 m water depths in the south-eastern part of the bay, on the Aquitaine continental slope (stations A, B, C, D and F), and in the north-eastern part, close to Cap Ferret canyon (stations H, I, 2 and 11; Fig. 1, Table 1), during 17 cruises from 1997 to 2006. Seasonal variations in benthic parameters were estimated at stations A, B and D, which were sampled more than ten times in different seasons. Other stations were sampled less frequently. The deepest stations are influenced by North Atlantic deep water (NADW) with a temperature of 4°C, those shallower than 1,200 m by warmer Mediterranean outflow water and Northern Atlantic central water (about 10-12°C). At all stations, the sediments consist of mud composed of silico-clastic clay and silt, and less than 30% carbonates (Hyacinthe et al. 2001). Water column samples were taken in February 2002 and March 2003 at stations A. B and D.

Material

Cores were collected with a Barnett multi-corer, which enabled us to sample the upper few decimetres of the sediments, the overlying bottom water, and the undisturbed sediment—water interface. The cores were sliced in thin horizontal sections (generally, 0.5 cm thick for the top 2 cm, 1 cm thick down to 8 cm, and 2 cm thick below this). For each depth interval, a sub-sample was immediately sealed in a pre-weighed vial, and frozen for subsequent analyses of

Fig. 1 Map of the south-eastern part of the Bay of Biscay, showing the locations of the stations (*squares*)

ATLANTIC OCEAN Bay of Biscay SPAIN FRANCE Bay of Biscay SPAIN FRANCE C FRANCE SPAIN FRANCE SPAIN FRANCE FRANCE	W1°		W2°	W3°	W4°
A B D FRANCE	N45°	\(\)	***************************************		Bay of Biscay
B D FRANCE	N44		A		4.
SPAIN		1 = [1000		
	W1°	/	~~~~	SPAIN	S. S.

Table 1 Study site characteristics

Station	Geographical p	Geographical position					
A	44°10.24′N	2°20.06′W	1,000				
В	43°50.31′N	2°03.47′W	550				
C	43°40.08′N	1°38.87′W	250				
D	43°42.00′N	1°33.45′W	140				
F	44°17.10′N	2°44.95′W	1,250				
11	44°30′N	2°40′W	1,600				
Н	44°32.52′N	3°37.23′W	2,000				
I	44°49.46′N	2°33.78′W	2,800				
2	45°30′N	6°30′W	4,825				

porosity and solid fraction. Another sub-sample was centrifuged under inert N_2 atmosphere at 5,000 rpm for 15 min in order to collect pore waters. A part of the supernatant was filtered (0.2- μ m cellulose acetate syringe filter) and acidified with ultrapure HNO₃ for dissolved Mn analyses.

Occasionally, surface sediments from a sister core were collected for excess ^{210}Pb ($^{210}\text{Pb}_{xs}$) and excess ^{234}Th ($^{234}\text{Th}_{xs}$) analyses.

Water column samples were taken using a multi-sampler/carousel with 12 bottles (12 l each), at depths close to the bottom (5 m above the seafloor), and at intermediate depths. Two or three bottles were sampled at each depth in order to collect enough suspended particles. The material was filtered on board through pre-weighed 0.4-µm cellulose acetate membranes for selective leaching to extract reactive Mn-oxides.



Analyses

Porosity was calculated from water content determined by comparison of the weights of wet and freeze-dried sediments. The specific density of whole particles was estimated at 2.65 (Berner 1980), which is the mean density of alumino-silicate and calcium carbonate minerals. Each freeze-dried solid fraction was homogenized, and the water content used to correct for the presence of sea salt.

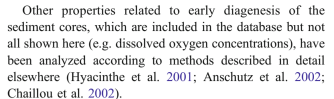
Solid-phase samples were subjected to two different extraction techniques for the determination of reactive solid-phase Mn. The most reducible fraction was extracted with an ascorbate solution (50 g NaHCO₃, 50 g Na-citrate, 20 g ascorbic acid for 1 l solution, buffered at pH 8; Kostka and Luther 1994; Anschutz et al. 2005). Another extraction on a separate aliquot was carried out with 1N HCl to determine acid-soluble Mn (Mn-HCl). We had tested for any influence of sediment pre-treatment by performing extractions on freeze-dried, on frozen but not dried, and on fresh sediment. The selective leaching solutions eventually extracted almost the same quantity of particulate Mn in the three cases. Consequently, we decided to work on dried samples, because the control on particle weight was much better than for wet sediments.

For the ascorbate and 1N HCl procedures, about 100 mg of dried sample was leached with 10 ml solution during 24 h, shaking continuously at ambient temperature. The supernatant was diluted with 0.2 M HCl for the ascorbate extraction, and with water for the HCl extraction. Manganese was measured by flame atomic absorption spectrometry (Perkin Elmer AA 300), using an external aqueous standard for calibration. The reproducibility of analyses was better than 5%. Mn-HCl represents the whole fraction of Mn-oxides and Mn associated with carbonates. Mn extracted with ascorbate (Mn-Asc) is only the most reducible part of Mn(III,IV) oxides and oxyhydroxides.

Mn-Asc and Mn-HCl inventories were calculated from depth \boldsymbol{a} to depth \boldsymbol{b} according to

$$I = \sum_{a}^{b} C_{z} \times (1 - \phi_{z}) \times \Delta_{z} \times \rho \tag{1}$$

where I is the inventory of Mn-Asc or Mn-HCl, Δ_z the depth interval between depths a and b, C_z the Mn-Asc or Mn-HCl concentration for Δ_z , ϕ_z the porosity for Δ_z , and ρ the particle density. All inventories were integrated for the upper 20-cm sediment column. This depth was chosen because it corresponds to the minimum length of the cores. The results show that for a 1-cm² horizontal core section, a 20-cm depth column represents a rather constant sediment inventory. The values vary between about 12 g at stations D and C, where mean porosity is highest, and 16 g at stations A, F and 11.



Maximum sediment mass accumulation rates and biodiffusion coefficients were estimated from vertical profiles of ²¹⁰Pb_{xs} (half-life=22.3 years) and ²³⁴Th_{xs} (half-life= 24.1 days). Activities of ²¹⁰Pb and of its parent ²²⁶Ra were determined in about 5 g of freeze-dried material sealed in a counting vial, by high-resolution and low-background gamma spectrometry with a semiplanar and a well-type detector for 4-24 h (Jouanneau et al. 1998a; Schmidt et al. 2007). The γ detectors were intercalibrated with IAEA reference materials (RGU-1 and RGTh-1). Supported ²¹⁰Pb was determined by three independent methods, which gave the same result within the counting errors: (1) average ²¹⁰Pb activities were estimated in deeper sections of the cores (except at stations C and D), (2) ²²⁶Ra-supported ²¹⁰Pb activities were estimated from direct measurements of ²²⁶Ra activities by gamma counting and (3) ²²⁶Ra activities were obtained from ²¹⁴Bi and ²¹⁴Pb. By subtracting the activity of ²²⁶Ra from the ²¹⁰Pb total specific activity, we obtained the unsupported ²¹⁰Pb component and used this to determine accumulation rates.

The uppermost sediment layers were measured for 234 Th by counting the 63.3 KeV gamma emission. These measurements had to be completed within 1 month after sampling, due to the rapid decay of 234 Th. Sediment layers were investigated downcore until a rather constant 234 Th activity was reached, which was considered as the supported activity used to calculate the 234 Th $_{xs}$ data. 234 Th activities were corrected for radioactive decay occurring between sample collection and counting.

Results

At all stations, the concentration of bottom water oxygen is always above 190 μ mol 1⁻¹. Oxygen penetration depth varies from 5 mm at the shallowest station D to 10 cm at the deepest station 2. Profiles of pore water oxygen are not shown here. For all studied cores, the depth where oxygen disappears corresponds to the depth interval where dissolved Mn appears (cf. Fig. 2).

Mn-Asc contents in suspended particles are always less than 2 μ mol g⁻¹ for water samples collected on the continental slope below 250 m water depth. Actually, the Mn-Asc contents of suspended particles measured in February 2002 and March 2003 are 2 μ mol g⁻¹ at station B and 1 μ mol g⁻¹ at station A. Each Mn-Asc profile in the sediment cores shows a subsurface maximum. Maximum



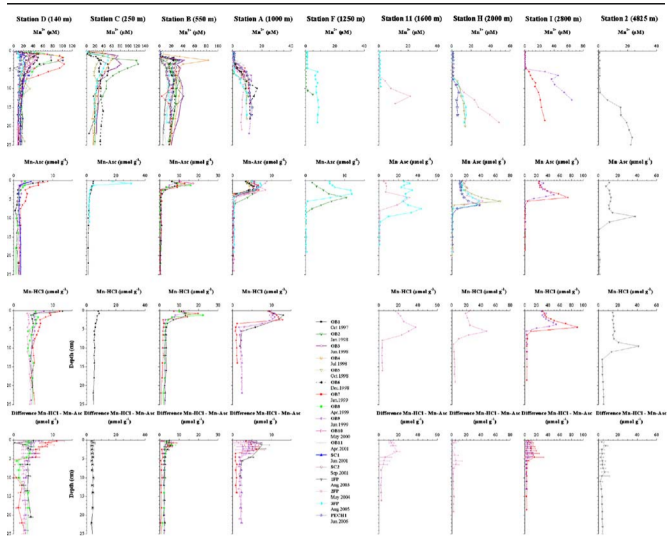


Fig. 2 Vertical depth (cm) profiles of pore water Mn (Mn²⁺, μM), Mn extracted by ascorbate (Mn-Asc, μmol g⁻¹), Mn extracted by HCl (Mn-HCl, μmol g⁻¹), and the difference between Mn-HCl and Mn-Asc at stations D, C, B, A, F, 11, H, I and 2. Distinct symbols represent the different cruises during which cores were collected: legend: *OB* Oxybent, *SC* Sedican, *FP* Foramprox cruises. The oxygen penetration depth of each core is contained in the sample where

dissolved Mn(II) concentration begins to increase. The error on Mn-HCl and Mn-Asc values is about 10% but not indicated here, for better readability. The sum of the error bars for the difference between Mn-HCl and Mn-Asc represents the sum of errors on Mn-HCl and Mn-Asc. The full dataset is available online in the electronic supplementary material for this article

values are above 30 μ mol g⁻¹ dry sediments for the deeper stations in contact with NADW, whereas the contents are lower than 20 μ mol g⁻¹ for the shallower stations (Fig. 2; the full dataset for this figure is available online in the electronic supplementary material for this article). Mn-Asc content is always higher in surface sediments than in suspended particles. For almost all stations, Mn-Asc contents decrease abruptly below the oxic front, and reach values close to zero. At the shallowest stations D and C, reactive Mn-oxides occur below the top of the anoxic sediments.

Mn-HCl profiles show similar trends but Mn-HCl contents are 2 to 5 μ mol g⁻¹ higher than Mn-Asc contents. The difference between the Mn-HCl and Mn-Asc values is close to Mn-HCl at the bottom of the studied cores (Fig. 2).

Mn-Asc and Mn-HCl inventories are reported in Tables 2 and 3, and the difference between Mn-HCl and Mn-Asc (cf. mainly Mn associated with carbonates) in Table 4. Sediments collected above 1,250 m water depth have Mn-Asc inventories of between 8 and 30 μmol per cm² (μmol cm⁻²) for a 20-cm-long core, the values being between 110 and 166 μmol cm⁻² for the deeper stations. Sediments collected above 1,250 m water depth have Mn-HCl inventories of between 30 and 80 μmol cm⁻², whereas the deeper stations have values between 130 and 190 μmol cm⁻². The inventories of Mn associated with the Mn-HCl minus Mn-Asc fractions are approx. constant (40±20 μmol cm⁻²).

Dissolved manganese concentrations are less than 2 μ mol Γ ⁻¹ in the oxic sediments. At the deeper stations, Mn^{2+}



Table 2 Mn-HCl inventories (μmol cm⁻²; SD standard deviation)

Cruise	Station										
	D (140 m)	C (250 m)	B (550 m)	A (1,000 m)	11 (1,600 m)	H (2,000 m)	I (2,800 m)	2 (4,825 m)			
OB1	69.0	71.5	58.4	60.9	=	=	=	=			
OB7	71.0	_	36.8	35.5	_	_	192.6	_			
OB8	66.6	_	50.3	_	_	_	_	_			
OB9	63.3	_	47.8	54.9	_	_	145.6	_			
1FP	_	_	_	_	_	_	_	181.2			
2FP	63.2	_	_	50.6	132.5	130.1	_	_			
Average	66.6	71.5	48.3	50.5	132.5	130.1	169.1	181.2			
SD	5%	_	16%	19%	-	-	14%	-			

concentration increases below the oxic layer and then reaches an almost constant value, between 10 and 20 μ mol l^{-1} . At the shallower stations, where Mn-Asc was detected in the anoxic sediments, Mn²⁺ profiles show a maximum below the oxic layer, with values reaching 100 μ mol l^{-1} . In the deeper anoxic zone, concentrations are close to 20 μ mol l^{-1} .

We observe a very similar overall shape for Mn-species profiles from cores collected in different months—years at a given station. However, the profiles are not identical. The difference becomes significant at the deeper stations, such as in replicate cores at stations F, 11 and H (Fig. 2). Here, two cores each collected during a different cruise to a given station can be situated several hundreds of meters apart because of ship drifting during multi-corer deployment. Thus, the seven cores studied for stations F, 11 and H can

be considered as representing seven distinct sites. For clarity, the profiles have been grouped by station on Fig. 2.

At the shallower stations, like stations D and C, vertical $^{210}\text{Pb}_{xs}$ profiles show almost no gradient (Fig. 3). At the deeper stations, by contrast, surface $^{210}\text{Pb}_{xs}$ activities are close to 1,000 Bq kg $^{-1}$, and the values decrease exponentially downcore to reach less than 100 Bq kg $^{-1}$ at depths varying between 2 cm at station 2 and 10 cm at station B. A distinct deviation in the gradient occurs at about 1 cm depth in the profiles of stations A and B.

Activities of 234 Th_{xs}, measured in the surface layers of the cores (Table 5), decrease strongly with depth, to below the detection limit at 1 cm depth for most of the cores. The values recorded in the upper 0.5 cm vary from cruise to cruise at a given station.

Table 3 Mn-Asc inventories (μmol cm⁻²; SD standard deviation)

Cruise	Station										
	D (140 m)	C (250 m)	B (550 m)	A (1,000 m)	F (1,250 m)	11 (1,600 m)	H (2,000 m)	I (2,800 m)	2 (4,825 m)		
OB1	14.5	19.3	19.0	12.7	_	_	_	_	_		
OB2	_	_	_	15.7	23.0	_	_	_	_		
OB3	_	_	_	12.5	_	_	_	_	_		
OB4	_	_	_	8.1	_	_	_	_	_		
OB5	_	_	_	11.7	_	_	118.5	_	_		
OB6	_	_	_	8.7	_	_	_	_	_		
OB7	27.8	_	18.0	13.9	_	_	_	123.9	_		
OB8	15.2	_	14.6	_	_	_	_	_	_		
OB9	20.7	_	14.3	15.1	_	_	_	100.1	_		
OB10	_	_	19.2	13.9	_	_	_	_	_		
OB11	_	_	_	11.3	_	_	_	_	_		
SC1	20.3	_	_	_	_	_	_	_	_		
1FP	_	_	_	_	_	_	_	_	124.5		
2FP	20.5	_	_	18.9	_	111.2	96.3	_	_		
3FP	_	28.9	_	16.4	27.6	166.9	109.1	_	_		
PECH1	_	_	_	_	_	_	91.0	_	_		
Average	19.8	24.1	17.0	13.2	25.3	139.1	103.7	112.0	124.5		
SD	22%	20%	13%	22%	9%	20%	10%	11%	_		



Table 4 Mn-HCl minus Mn-Asc inventories (μmol cm⁻²)

Cruise	Station										
	D (140 m)	C (250 m)	B (550 m)	A (1,000 m)	11 (1,600 m)	H (2,000 m)	I (2,800 m)	2 (4,825 m)			
OB1	54	52	39	48	_	_	_	_			
OB7	43	_	19	22	_	_	69	_			
OB8	51	_	36	_	_	_	_	_			
OB9	42	_	33	40	_	_	45	_			
1FP	_	_	_	_	_	_	_	57			
2FP	43	_	_	32	21	34	_	_			
Average	47	52	32	36	21	34	57	57			

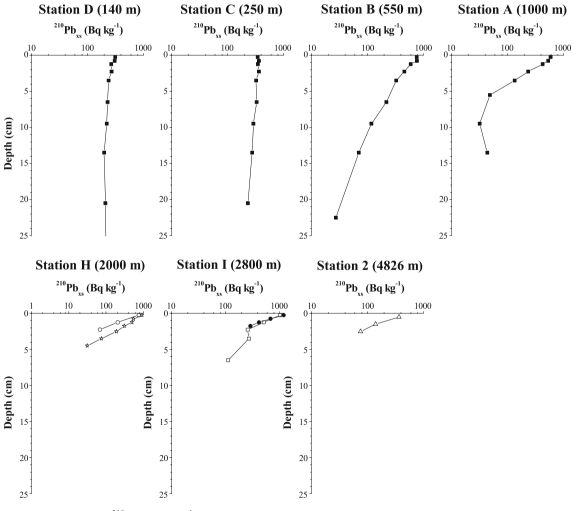


Fig. 3 Vertical profiles of log $^{210}\text{Pb}_{xs}$ (Bq kg $^{-1}$). The profiles at stations D, C, B and A (*closed squares*) were obtained from cores collected during the Oxybent 1 cruise. The *open dots* represent log $^{210}\text{Pb}_{xs}$ of surface sediments at station H collected during Oxybent 5,

and *open stars* during Pech 1. At station I, *open squares* represent data from Oxybent 7, *and closed dots* from Oxybent 9. Station 2 was sampled during the Foramprox 1 cruise (*open triangles*)



Table 5 $^{234}\text{Th}_{xs}$ data used for the calculation of biodiffusion coefficients (see text; nd not detected)

Depth	$^{234}\mathrm{Th}_{\mathrm{xs}}$
(cm)	$(Bq kg^{-1})$
0-0.50	110
0.50 - 1.00	33
0-0.25	2,704
0.25-0.50	325
0.50-0.75	538
0.75-1.00	118
0-0.50	801
0.50 - 1.00	66
0-0.50	270
0.50-1.00	32
0-0.50	375
0.50 - 1.00	237
1–2	80
0-0.50	102
0.50 - 1.00	nd
0-0.50	407
0.50 - 1.00	107
0-0.50	312
0.50-1.00	nd
0-0.50	263
0.50-1.00	nd
0-0.50	451
0.50-1.00	341
	0-0.50 0.50-1.00 0-0.25 0.25-0.50 0.50-0.75 0.75-1.00 0-0.50 0.50-1.00 0-0.50 0.50-1.00 1-2 0-0.50 0.50-1.00 0-0.50 0.50-1.00 0-0.50 0.50-1.00 0-0.50 0.50-1.00

Discussion and conclusions

Transport process characterization

Bioturbation is classically described as an eddy-diffusive transport process occurring within a surface mixed layer

(Goldberg and Koide 1962). Biodiffusion is a local transport associated with spatially random and small-scale sediment displacements between discrete mixing events (Boudreau 1986a; Meysman et al. 2003). Nonlocal mixing, which implies the exchange of material between non-adjacent points, can exist in natural environments (Boudreau 1986b) but, due to the absence of peaks along the radiogenic profiles reported for the present study (Fig. 3), this process is not considered further in this case.

Surface sediments can be characterized by a mixed layer of thickness L with both a constant biodiffusion coefficient (Db) and a constant burial rate (w). The mixed-layer depth is generally defined as the depth of strong slope inflection along the 210 Pb profile. Biodiffusion coefficients are computed from 234 Th profiles (Table 6). Recently, Lecroart et al. (2007) reported that short-lived radioisotopes are highly suitable to constrain bioturbation in coastal and shelf sedimentary environments. At steady state, and assuming constant porosity, the classical advection-diffusion model equation adapted for short-lived radionuclides is used for the mixed layer:

$$0 = Db(t)\frac{\partial^2 A}{\partial z^2} - w\frac{\partial A}{\partial z} - \lambda A \tag{2}$$

where A is the activity of the radionuclide within the mixed layer, z the depth in the mixed layer relative to the sediment–water interface, and λ the decay rate of 234 Th (λ =10.51 year⁻¹). Solution of Eq. (2) enables computation of Db:

$$Db = \frac{\lambda z^2}{\left[\ln(A(z)/A_0)\right]^2} + \frac{wz}{\ln(A(z)/A_0)}$$
(3)

where A_0 is the activity of ²³⁴Th at the sediment–water interface. Db can be characterized in terms of depth profiles of ²³⁴Th if an independent tracer is used to compute the

Table 6 Burial rates and biodiffusion coefficients computed from ²¹⁰Pb and ²³⁴Th

	Water depth (m)	Cruise #	Burial rate (cm year ⁻¹)	Mixed-layer depth (cm)	Biodiffusion coefficient (cm ² year ⁻¹)
Station D	140	OB1	_	>30	=
Station C	250	OB1	_	>20	_
Station B	550	OB1	0.120	1	1.80
		OB2	_	_	0.80
		OB8	_		0.40
		OB10	_	_	14.80
Station A	1,000	OB1	0.062	1	0.56
		OB2	_	_	6.70
		OB3	_	_	0.12
		OB10	_		1.45
Station I	2,800	OB7	0.045	<1	< 0.08
		OB9	0.033	<1	< 0.08
		OB10	_	_	0.55



burial rate. Below the mixed layer, the ²¹⁰Pb profile is affected only by burial and decay. At steady state, the conservation equation enables us to compute the burial rate

$$w = -\frac{\lambda}{p} \tag{4}$$

where p is defined in a semi-logarithmic diagram by the slope of the ²¹⁰Pb profile beneath the bioturbated layer.

At stations C and D, the thickness of the bioturbated layer exceeds the length of the cores (Fig. 3, Table 6), meaning that bioturbation dominates over sedimentation, making it impossible to determine maximum MAR in this case. At the deepest stations (A, B and I), the bioturbated layer is thinner ($L \le 1$ cm) than that commonly found in sedimentary environments (Boudreau 1998). This may be due to the high proportion of mostly refractory organic matter of terrestrial origin present at these stations (Etcheber et al. 1999; Grémare et al. 2005), which probably does not favour deep endomacrobenthic activity. The corresponding biodiffusion coefficients range from 0.08 to 14.8 cm² year⁻¹. Bioturbation could have slightly affected the manganese redox front at station B but less so at the other deep stations. Nevertheless, because the sediment could have been subjected to disturbance by burrowing organisms at all stations, ²¹⁰Pb_{xs} profiles may represent maximum accumulation

To conclude, accumulation rates deduced from ²¹⁰Pb_{xs} and porosity profiles have maximum values of 80, 36, 14, 17 and 20 mg cm⁻² year⁻¹ at stations B, A, H, I and 2 respectively. Calculations of accumulation rates are less robust at stations H, I and 2, because of the shortness of the ²¹⁰Pb_{xs} profiles. Grain size and calcium carbonate content are constant along cores collected at stations A, B, D and I (Hyacinthe et al. 2001), suggesting that mass accumulation rates can be assumed to be constant along the first decimetres of the sediment column of the Bay of Biscay slope.

Diagenetic recycling of Mn at the oxic-anoxic front

The ascorbate method enables us to extract bio-available Mn-oxides selectively, although it is not possible to differentiate between Mn(III) oxyhydroxides and Mn(IV) oxides (Hyacinthe and Van Cappellen 2004; Anschutz et al. 2005). The 1N HCl leaching method yields the ascorbate-extractable fraction and Mn associated with acid-soluble phases, such as carbonates and hydrous aluminium silicates. The general shape of the Mn-species profiles in sediments of the Bay of Biscay reflects the common vertical redox sequence observed in modern marine sediments. Manganese(III,IV) oxides extracted with ascorbate are present in the oxic layer, whereas dissolved Mn(II) becomes detectable in the anoxic sediments (Fig. 4). The

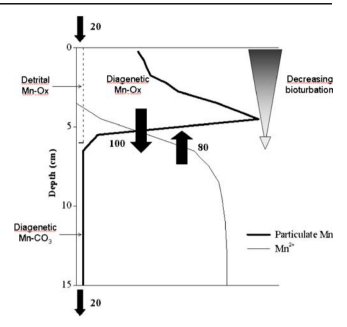


Fig. 4 Cycle of manganese in sediments of the Bay of Biscay. At steady state, the flux of buried Mn-HCl is equal to the flux of settling detrital manganese oxides from the water column. The major part of the Mn-oxide pool is of diagenetic origin. This pool is at steady state. The downward flux of diagenetic Mn-oxides at the redox front is equal to the upward flux of Mn(II). Flux values in the figure are indicative, with an arbitrary unit

disappearance of Mn(III,IV) species below the oxic layer is generally explained by their dissimilatory reduction by heterotrophic bacteria (Froehlich et al. 1979). The oxidation of Mn(II) with O₂ explains the decrease in Mn(II) concentrations from the anoxic sediments to the oxic boundary. At steady state, when the redox front does not migrate relative to the sediment surface, the shape of Mnoxide profiles should show only one peak at the depth of the manganese redox front (Burdige and Gieskes 1983). Actually, our database reveals high Mn-Asc contents in the whole oxic zone for all profiles, and these cannot be explained by corresponding enrichment levels in settling particles (cf. the latter are substantially less enriched; see above). This points to essentially four alternative explanations, not necessarily mutually exclusive.

- The Mn redox boundary and oxygen penetration depth may migrate within the Mn-oxide-enriched layer as a consequence of episodic fresh organic matter sedimentation. These redox fluctuations can redistribute Mnoxides in the oxic zone (Katsev et al. 2006).
- Episodic bioirrigation through burrows may enable anoxic water advection to the oxic layer, where Mn (II) can be oxidized.
- Mn-oxides may be mixed throughout the oxic layer by biological activity. Because of their stability under oxic conditions, they would accumulate in this depth interval. A relatively low level of bioturbation may



- suffice to redistribute Mn-oxide particles while maintaining a sharp gradient of Mn-Asc at the redox boundary, and a decreasing $^{210}{\rm Pb}_{\rm xs}$ slope.
- 4. Mn-oxide may be locally reduced to Mn(II) with ammonia produced by organic nitrogen mineralization within the oxic layer (Luther et al. 1997; Anschutz et al. 2000). It also can be reduced to dissolved Mn(III) (Trouwborst et al. 2006). Thus, local concentration of dissolved Mn could generate a short phase of diffusive transport before re-oxidation with oxygen to Mn-oxide. Repeated occurrence of this process could disperse the Mn-oxide pool uniformly in the oxic layer up to the sediment—water interface, without producing detectable peaks in dissolved Mn concentrations.

In cases 1 and 2, we would expect that at least a few of the studied cores would present evidence of significant dissolved Mn enrichment extending up to the Mn-oxide-enriched layer. Our database, comprising more than 30 cores collected at different places and times, and in which Mn-oxides and dissolved Mn were measured concurrently, reveals no indication of high dissolved Mn concentrations above the Mn-oxide maximum. Therefore, the bottom of the Mn-oxide-enriched layer is probably the only depth where Mn(II) is oxidized. Explanation 4 suggests that some dissolved Mn crosses the sediment—water interface and escapes from the sediment. Below, we argue against sediment loss of Mn. Therefore, a low level of bioturbation may account for the uniformly high Mn-Asc levels recorded in the oxic layer of our cores.

Mn-HCl at the bottom of the studied cores is probably representative of the reactive buried Mn fraction. At steady state, the content of buried Mn must be equal to that of settling detrital Mn-oxides (Fig. 4), especially at stations where the concentration of dissolved Mn(II) is close to zero in the oxic layer. At stations B and A, the Mn-oxide contents of settling particles are very similar to the difference between the Mn-HCl and Mn-Asc pools of the anoxic sediment layer (Fig. 2). Therefore, Mn is probably at steady state at a multi-annual timescale.

Constant values of dissolved Mn recorded at depth in most of the cores can be explained by an equilibrium between the aqueous Mn(II) and an authigenic Mn phase. An increase in alkalinity due to sulphate reduction observed in the shallowest cores (Chaillou et al. 2003), and observations reported in the literature suggest that authigenic Mn associated with carbonate can represent the main phase under which reactive Mn is ultimately buried in anoxic sediments (Holdren et al. 1975; Grill 1978; Suess 1979; Aller 1980; Pedersen and Price 1982; Thomson et al. 1986; Middelburg et al. 1987; Mucci 1988, 2004; Jakobsen and Postma 1989). Under oxic conditions, authigenic Mn-carbonate precipitation is not favoured, because of low Mn enrichment and acidity

produced by aerobic respiration (e.g. Jahnke et al. 1997). Although errors on the difference between the Mn-HCl and Mn-Asc pools become high in the oxic layer, there is an enrichment of this fraction in the oxic zone at some stations (A, B and 11). Some refractory Mn-oxides that are not leachable with ascorbate but extracted by HCl may explain this excess. Adsorption of Mn(II) onto particles such as iron oxides also may enrich the oxic layer in extractible Mn (e.g. van der Zee et al. 2001).

To conclude, authigenic Mn-oxides precipitate close to the oxic-anoxic boundary and are advected via particle mixing. At depth, the reactive Mn content of the anoxic sediments corresponds to the Mn content of settling particles. Diagenetic Mn recycling at the oxic-anoxic front modifies Mn solid speciation but not Mn inventories.

Shape of pore water Mn(II) profiles

Dissolved Mn profiles exhibit different shapes depending on the presence or absence of Mn-Asc in the anoxic sediments. A dissolved Mn peak is present in the upper part of the anoxic layer when Mn-Asc occurs below the oxic layer. Such peaks can be built when the reductive dissolution of Mn-oxides exceeds the upward and downward Mn(II,III) diffusive fluxes triggered by Mn(II,III) re-oxidation and authigenic carbonate formation respectively.

Bioturbated (shallower) stations

With strong bioturbation, as observed at the shallowest stations, significant amounts of Mn-oxides can be conveyed to the anoxic zone where they are reduced, resulting in a dissolved Mn(II) peak. A fraction of dissolved Mn may consist of transient Mn(III) (Trouwborst et al. 2006). When bioturbation is permanent, as at stations C and D, this Mn (II) peak can persist (Fig. 2). Otherwise, such peaks would be temporary or less pronounced, as detected in some cores of station B.

The presence of Mn-oxide in the anoxic sediment layer could also be explained by an abrupt upward shift of the redox boundary (oxygen penetration depth, Mn redox boundary), which would "bury" the formerly active Mn spike into anoxic conditions (Katsev et al. 2006). Because the Mn peak was detected in all cores collected at stations D and C, the bioturbation scenario is more likely in the present case.

Less bioturbated (deeper) stations

At the deepest stations (A, F, H, 11, I and 2), the finding that several profiles show a gradual increase of dissolved Mn below the oxic layer (without a maximum concentration peak) suggests that manganese oxides are transported to the anoxic zone only by slow sedimentation. Oxides are



reduced at the bottom of, or immediately below the oxic layer via heterotrophic bacterial respiration or by means of reduced compounds that diffuse from below, such as dissolved sulphide, iron or ammonia (Anschutz et al. 2002).

To conclude, an Mn(II) peak below the oxic zone can be an indication that bioturbation affects the anoxic part of the sediments. The fact remains that a part of reducible Mn must reach the anoxic sediments far below (several cm) the oxic—anoxic front in order to sustain the upward flux of dissolved Mn(II), and the formation of Mn associated with carbonates.

Spatial heterogeneity of particulate Mn distribution

Mn-Asc profiles can show temporal variations for a given station (Fig. 5a). We initially suspected that this was due largely to heterogeneous sub-sampling of cores. In a simple model, we calculated and drew the profiles of manganese oxides obtained for different resolutions of core slicing along a porosity gradient (Fig. 5b; details are given in the caption to this figure). The resulting profiles are very dissimilar from one slicing step to another, whereas the overall inventory of Mn-oxide is always the same. Thus, the shape of profiles is not appropriate to meaningfully compare site- and time-specific vertical distributions of a given species. Better would be to integrate the Mn inventory for each core, since the vertical slicing resolution varied during our sampling campaigns.

The data show that, at a given station, inventories of Mn-Asc and Mn-HCl are of the same order of magnitude for the

different sampling time periods. The standard deviation is always lower than 22% (Tables 2 and 3), and can largely be explained by a likely 20% error in manganese and porosity measurements. We can also hypothesize that such fluctuations could at least partly be due to transient-state processes. However, quantitative examination argues against this. For example, at station A, the Mn-Asc inventory of the core collected during the Oxybent 4 cruise is $7.6\pm4.7 \,\mu\text{mol cm}^{-2}$ higher than that for the core collected during Oxybent 2 (Table 2). Considering the maximum diffusive flux of dissolved Mn(II) at station A (Table 7), a diffusion time of 20±12 years would be necessary to explain this enrichment in Mn-Asc, but the time span between the two cruises is only 7 months. On the other hand, spatial heterogeneities may well be important in this respect, since what we call "a station" in fact encompasses a relatively large area because of ship drifting during coring operations, particularly at the deepest stations. Indeed, in such deep-sea sampling campaigns, it is difficult to discriminate between patchiness and any temporal evolution trends (e.g. Fontanier et al. 2003).

Sediments collected above 1,250 m water depth have Mn-Asc inventories lower than for the deeper stations (Fig. 6). It is noteworthy that the deeper stations are located in the northern sector of the study area, so that this contrast with the southern stations could be due to differences in terrigeneous material sources. This hypothesis remains to be verified, because the assessment of terrigeneous material origin in the Bay of Biscay has to date focused only on the

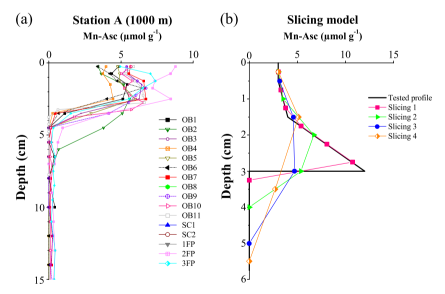


Fig. 5 a Mn-oxides (Mn-Asc) concentration profiles (μmol g⁻¹) vs. depth at station A for the period 1997 to 2006. **b** Model of different ways of slicing a core. *Thick black line* Mn-Asc content profile tested in the model, *slicing 1* slicing step of 0.5 cm, *slicing 2* initial slicing of 0.5 cm and then a step of 1 cm, *slicing 3* slicing step of 1 cm for the upper 2 cm of the core and of 2 cm for the remainder, *slicing 4* initial slicing of 0.5 cm and then a step of 2 cm. The resulting profiles are

different for the slicing steps, whereas the overall inventory of Mnoxides is always the same. The Mn-oxide enrichment peak is smoothed because of slicing, and the disappearance depth of Mnoxides is not accurately assessed. Mn-oxides are not present below 3 cm depth; depending on the slicing step used, Mn-oxides disappear between 3.25 and 5.50 cm depth. In the slicing 4 profile, one data point shows Mn-oxides at a depth where they should not be present



Table 7 Dissolved Mn²⁺ fluxes (µmol cm⁻² year⁻¹)

Cruise	Date	Station									
		D (140 m)	C (250 m)	B (550 m)	A (1,000 m)	F (1,250 m)	11 (1,600 m)	H (2,000 m)	I (2,800 m)	2 (4,825 m)	
OB1	Oct. 1997	7.67	1.10	1.54	0.24	_	_	_	_	_	
OB2	Jan. 1998	4.75	5.48	0.49	0.25	0.18	_	_	_	_	
OB3	Jun. 1998	3.65	6.94	0.48	0.14	_	_	_	_	_	
OB4	Jul. 1998	10.59	1.83	0.57	0.11	_	_	_	_	_	
OB5	Oct. 1998	8.40	5.48	0.35	0.22	_	_	0.26	_	_	
OB6	Dec. 1998	6.21	1.83	0.50	0.12	_	_	_	_	_	
OB7	Jan. 1999	10.59	_	0.56	0.26	_	_	_	0.19	_	
OB8	Apr. 1999	8.03	_	0.55	_	_	_	_	_	_	
OB9	Jun. 1999	5.11	_	0.38	0.22	_	_	_	1.06	_	
OB10	May 2000	3.65	_	0.32	0.38	_	_	_	_	_	
OB11	Apr. 2001	2.92	_	_	_	_	_	_	_	_	
SC1	Jun. 2001	3.65	_	_	_	_	_	_	_	_	
SC2	Sep. 2001	6.57	_	_	_	_	_	_	_	_	
1FP	Aug. 2003	_	_	_	_	_	_	_	_	0.22	
2FP	May 2004	2.92	_	0.22	0.15	_	0.19	0.26	_	_	
3FP	Aug. 2005	1.10	6.94	_	0.13	0.44	_	0.24	_	_	
PECH1	Jun. 2006	4.73	_	1.31	0.18	_	_	0.15	_	_	
Average	_	5.66	4.22	0.60	0.20	0.31	0.19	0.23	0.63	0.22	

continental shelf (Jouanneau et al. 1998b), and not the deeper zones. Nevertheless, the data discussed below argue against this explanation for these north/south gradients.

Because the inventories of Mn associated with the Mn-HCl minus Mn-Asc fractions are approx. constant, Mn(III, IV) oxides extracted by ascorbate largely control the Mn-HCl levels. Manganese carbonate levels in these anoxic sediments (representing the major part of the Mn-HCl minus Mn-Asc fractions) are very similar at the northern and southern stations (Fig. 6), with values very close to that of 2 µmol Mn-Asc g⁻¹ recorded in suspended particles in the Bay of Biscay waters. Because manganese is a conservative element (i.e. buried Mn-HCl content would be equal to the content of settling detrital manganese oxides), it follows that input at the northern stations is not enriched in Mn, compared to that at the southern stations.

To conclude, the shape of vertical profiles of particulate Mn changes with time. However, changes in the distribution of particulate Mn are low, and they can be attributed to patchiness and to the procedure of core sub-sampling. At a given station, inventories of Mn-oxide present in the upper sediment layer do not change significantly with time. This suggests that the quantity of diagenetic Mn-oxide that crosses the Mn(III,IV)–Mn(II) redox front due to burial is balanced by the upward flux of dissolved Mn(II). Integration of particulate Mn inventories enables us to compare the stations studied. The differences in Mn-oxide inventories would not be caused by variable sources of settling

material. One possible explanation would be distinct combinations of sedimentation rate and benthic metabolism, which would maintain an Mn-rich layer at the northern stations in contact with North Atlantic deep water, and a less enriched layer at shallower stations.

Diagenetic Mn(III,IV) oxide formation

Fluxes of dissolved Mn(II)

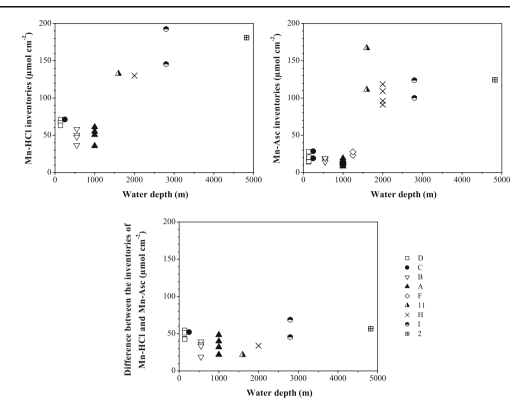
Diagenetic Mn-oxide authigenesis results from the oxidative precipitation of dissolved Mn(II) that diffuses upwards from the anoxic sediments (Fig. 4). The flux of dissolved manganese in pore waters can be calculated (assuming transport by molecular diffusion) from the concentration gradients according to Fick's first law:

$$J = \phi D_s dC/dX \tag{5}$$

where J is the flux, ϕ the porosity, $\mathrm{d}C/\mathrm{d}X$ the concentration gradient, and D_{s} the bulk sediment diffusion coefficient corrected for tortuosity, i.e. $D_{\mathrm{s}} = D_0/\theta^2$ where θ is the tortuosity, and D_0 the diffusion coefficient in water (Berner 1980). D_0 values were obtained from Li and Gregory (1974), and θ^2 is assumed to be equal to $1 - \ln(\phi^2)$ (Boudreau 1996). Fluxes were calculated from the linear gradient of Mn^{2+} in the upper part of the anoxic zone. These gradients were smoothed, i.e. the narrow peaks of dissolved Mn presumably caused by sporadic bioturbation were averaged out (e.g. at station B).



Fig. 6 Inventories of Mn extracted by HCl (Mn-HCl), Mn extracted by ascorbate (Mn-Asc), and the difference between the inventories of Mn-HCl and Mn-Asc vs. water depth at stations D, C, B, A, F, 11, H, I and 2



Bioturbated (shallower) stations Upward diffusive fluxes of dissolved Mn are highest (>2 μmol cm⁻² year⁻¹) at the shallowest stations D and C (Table 7). Flux values are not constant with time at a given station, indicating that non-steady-state conditions influence Mn cycling at the redox front. Nevertheless, the values remain in the same order of magnitude.

The high flux values calculated at stations D and C can be explained by the presence of an Mn(II) peak observed below the oxic layer. In turn, the magnitude of this peak depends on the amount of Mn-oxides transported, and thus reduced, in the upper part of the anoxic layer. Dissolved Mn fluxes vary with time, probably because bioturbation is not a steady-state process at these stations at the timescale corresponding to the frequency of the cruises.

Less bioturbated (deeper) stations At the deepest stations, diffusive fluxes are less than 1 μmol cm⁻² year⁻¹ (Table 7). It should be noted that the Mn(II) concentration gradient is deduced from only two or three data points for each station. This is not enough to obtain sufficiently high precision for these flux calculations.

Calculations of mass accumulation rates

Because the content of settling detrital Mn-oxides ($<2 \mu mol g^{-1}$) is low in terms of the total Mn-oxide pool, the enrichment recorded in the oxic layer is mostly of diagenetic origin,

and not inherited from sedimenting particles (Fig. 4). Therefore, we can assume that manganese oxide inventories result largely from the oxidation of dissolved Mn(II) that diffuses from below. The progressive development of an Mn-enriched layer in the oxic zone, i.e. the gradual increase with time of the Mn-Asc inventory, is a transient process that has not been observed at the studied stations. When the Mn-enriched layer reaches steady state, the burial flux of diagenetic Mn-oxides towards the anoxic zone must be essentially equal to the upward flux of Mn (II) (Fig. 4). Therefore, we can use this average upward flux of Mn(II) (J) and the inventory of Mn(III,IV) (I) to calculate the residence time T of Mn-oxides in the oxic layer, according to

$$T = T/J \tag{6}$$

Calculations of *T* were performed for the deeper stations, for which Mn-oxide inventories can be considered to be essentially constant. *T* is 28, 66, 82, 747, 460, 179 and 559 years for stations B, A, F, 11, H, I and 2 respectively (Table 8). This corresponds to the time necessary for Mn-oxides to move out of the oxic zone, due to burial. Mn is a proxy for the time needed to bury, below the oxic front, a pool of particles included in the oxic layer. Thus, *T* is equivalent to the time required for the accumulation of particles contained in the oxic layer. Therefore, the distribution of Mn species can be used to determine mass



Table 8 Results and parameters required to calculate residence time T (see text)

Station	Average Mn ²⁺ flux (µmol cm ⁻² year ⁻¹)	I: average Mn-Asc inventory (μmol cm ⁻²)	T: residence time of Mn-Asc (years)	M: mass of the oxic layer (mg cm ⁻²)	Accumulation rate deduced from Mn (mg cm ⁻² year ⁻¹)	Accumulation rate deduced from $^{210}\text{Pb}_{xs}$ (mg cm $^{-2}$ year $^{-1}$)
B (550 m)	0.60	17.0	28	1,517	54	80
A (1,000 m)	0.20	13.2	66	2,270	34	36
F (1,250 m)	0.31	25.3	82	3,565	44	_
11 (1,500 m)	0.19	139.1	747	5,073	7	_
H (2,000 m)	0.23	103.7	460	4,309	9	14 (7°)
I (2,800 m)	0.63	112.0	179	3,287	18	17 16 ^a 18 ^b
2 (4,825 m)	0.22	124.5	559	8,418	15	20
20	2.25	27.7	12	1,352	110	113 ^d
25	3.49	68.6	20	2,279	116	175 ^d
PB (3,890 m)	1.53	276.8	181	1,100	6	7 ^e
BT C (980 m)	0.49	208.0	421	6,323	15	40

Accumulation rates are maximum values deduced either from ²¹⁰Pb_{xs} values or from the distribution of Mn compounds (see text for details)

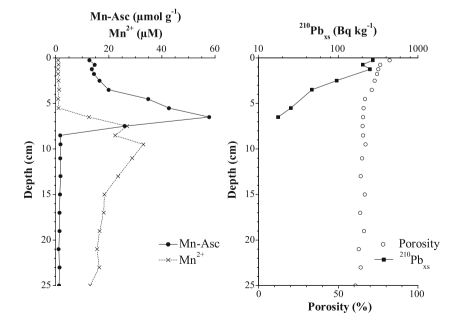
accumulation rate. The method consists of calculating T, and applying this time according to

$$MAR = M/T \tag{7}$$

where MAR is the mass accumulation rate, and M the mass of particles present in the oxic layer. Here, M was calculated for a projected area of 1 cm² by integration of the dry mass of particles deduced from particle volumes

(i.e. $1-\phi$), and a mean particle specific density of 2.65 (Berner 1980). We used this Mn method to estimate the MAR of those stations for which the Mn-oxide inventory was included largely within the oxic layer, i.e. the deeper stations (Table 8). Moderate bioturbation, which would redistribute Mn-oxide in the oxic layer, does not affect the concept of our model. The calculated values are between 34 and 54 mg cm⁻² year⁻¹ at the stations with low Mn-Asc

Fig. 7 Vertical distributions of particulate Mn extracted by ascorbate (Mn-Asc, μ mol g⁻¹), dissolved Mn(II) (Mn²⁺, μ M), 210 Pb excess activity (210 Pb_{xs}, Bq kg⁻¹) and porosity (%) at station BT C in the Mediterranean Sea (980 m, 42°43′18N, 4°46′58E)





^a Data from sediment trap at MS2 site, 3,000 m water depth, close to station I (ECOFER program, Heussner et al. 1999)

^b Data estimated with ²¹⁰Pb budget at MS2 site, 3,000 m water depth, close to station I (ECOFER program, Radakovitch and Heussner 1999)

^c Data from sediment traps (Schmidt, unpublished data)

^d Data from sediment traps, St. Lawrence estuary (Silverberg et al. 1986)

^e Data from sediment traps, Panama Basin (Honjo 1982)

inventories (B, A and F), and between 18 and 7 mg cm⁻² year⁻¹ at the deeper stations.

Validation of the method

We have attempted to validate the Mn method with other independent data from sediment traps and ²¹⁰Pb_{xs} profiles. Sediment trap data are available from the ECOFER program, for the Cap Ferret canyon at the site MS2 located close to our station I (2,800 m), and from the FORCLIM program, at our station H. Site MS2 at 3,000 m water depth provided a mean MAR of 16 mg cm⁻² year⁻¹ (Heussner et al. 1999). Radakovitch and Heussner (1999) obtained a value of 18 mg cm⁻² year⁻¹ at the same site, using a budget of ²¹⁰Pb. From sediment trap data, Schmidt (unpublished data) obtained a mass accumulation rate of 7 mg cm⁻² year⁻¹ at station H. These values are in excellent agreement with those obtained with the Mn method at stations I and H—18 and 9 mg cm⁻² year⁻¹ respectively (Table 8).

Mass accumulation rates estimated with ²¹⁰Pb_{xs} in sediments collected at stations B, A, H, I and 2 are remarkably close to values deduced from the distribution of Mn phases (Table 8), although these represent independent calculations. At stations A and I, Mn values are within 10% of ²¹⁰Pb_{xs} values. At stations H and 2, they are within 35%. The limited number of dissolved Mn profiles at these stations could explain this greater difference. Because pore water Mn²⁺ levels can be unstable close to the redox front, any Mn²⁺ profile in a transient state would cause deviations from the overall flux representative of processes at steady state. The difference between the Mn method and the $^{210}\text{Pb}_{xs}$ method is even greater at station B (Table 8), where the ²¹⁰Pb_{xs}-based MAR is higher, probably because bioturbation significantly affects the slope of the ²¹⁰Pb_{xs} profile, but not the Mn-oxide inventories.

Application to other environments

In order to better validate the Mn method for MAR calculation, we applied it to sedimentary environments for which published profiles of Mn species exist, together with MAR measured alternatively. Porosity data are necessary to calculate the inventories of Mn-oxide. This parameter is often lacking in the articles we examined. A complete set of data has been found for sediments from the outer St. Lawrence estuary (Sundby et al. 1981; Silverberg et al. 1986; Silverberg and Sundby 1990), and from the Panama Basin (Honjo 1982; Aller 1990; Aller et al. 1998). Here again, the mass accumulation rates obtained with the Mn method are remarkably close to those based on sediment trap data for both locations (Table 8).

Finally, we tested the method using data from sediment cores we collected in September 2006 in the western

Mediterranean Sea, at 980 m water depth (Fig. 7). The upward diffusive flux of dissolved Mn obtained from this single profile is 0.49 µmol cm⁻² year⁻¹. The time *T* to form the inventory of Mn-Asc in the oxic layer with this flux would be 421 years. The particulate material *M* contained in the Mn-oxide-enriched layer is 6,323 mg cm⁻², corresponding to a deduced MAR of 15 mg cm⁻² year⁻¹. The ²¹⁰Pb_{xs} profile gives a value of 40 mg cm⁻² year⁻¹. The presence of a bioturbation-induced mixed layer at the top of the core (Fig. 7), like at station B in the Bay of Biscay, or non-steady-state conditions probably explains this difference between the two MAR estimates.

To conclude, steady-state particulate Mn inventories and fluxes of dissolved Mn make it possible to estimate the residence time for diagenetic Mn-oxides in the oxic layer. By extension, the distribution of Mn species enables us to calculate mass accumulation rates. At the stations studied in the Bay of Biscay, mass accumulation rates obtained from the Mn method fit well with values obtained from the conventional ²¹⁰Pb_{xs} method and sediment trap data. This indicates that the diagenesis of Mn in these environments is at steady state at the multi-annual timescale. Compared to radionuclide techniques of determining mass accumulation rates, the analytical expense of the Mn method is high, including pore water as well as ascorbate extractions and analyses. It is, however, a suitable alternative approach, and could prove useful where geochemical investigations are anyhow being, or have been undertaken.

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