Hydrothermal sediments record changes in deep water oxygen content in the SE Pacific

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[1] The distribution of redox-sensitive metals in sediments is potentially a proxy for past ocean ventilation and productivity, but deconvolving these two major controls has proved difficult to date. Here we present a 740 kyr long record of trace element concentrations from an archived sediment core collected at $\sim 15^{\circ}$ S on the western flank of the East Pacific Rise (EPR) on 1.1 Myr old crust and underlying the largest known hydrothermal plume in the world ocean. The downcore trace element distribution is controlled by a variable diagenetic overprint of the inferred primary hydrothermal plume input. Two main diagenetic processes are operating at this site: redox cycling of transition metals and ferrihydrite to goethite transition during aging. The depth of oxidation in these sediments is controlled by fluctuations in the relative balance of bottom water oxygen and electron donor input (organic matter and hydrothermal sulfides). These fluctuations induce apparent variations in the accumulation of redox-sensitive species with time. Subsurface U and P peaks in glacial age sediments, in this and other published data sets along the southern EPR, indicate that basin-wide changes in deep ocean ventilation, in particular at glacial-interglacial terminations II, III, IV, and V, alter the depth of the oxidation front in the sediments. These basin-wide changes in the deep Pacific have significant implications for carbon partitioning in the ocean-atmosphere system, and the distribution of redox-sensitive metals in ridge crest sediment can be used to reconstruct past ocean conditions at abyssal depths in the absence of alternative proxy records.

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1. Introduction

[2] Hydrothermal sediments derived from plume dispersion of vent derived particles accumulate around tectonically active margins such as mid-ocean ridges [Mills and Elderfield, 1995], but are rarely exploited for paleoceanographic reconstruction of past ocean conditions because of the complexity of inputs and postdepositional changes in these sediments. Iron-oxide hydrothermal particles coprecipitate certain trace elements (e.g., U, Mo, As, P, V) from seawater [Edmonds and German, 2004; Feely et al., 1996] and many trace metals (e.g., Fe, Cu, Zn, U, Mo, Cd, Pb) are enriched in sulfide particles which are transported along with the Fe oxides [Trocine and Trefry, 1988]. The sedimentary record of these elements depends on the nature and extent of postdepositional reactions at the seafloor [Poulton and Canfield, 2006]. The trace element record in metalliferous sediments can potentially reveal past changes in ocean chemistry if the seawater record is preserved [Edmonds and German, 2004; Feely et al., 1996], but problems in interpretation arise because the record can be perturbed during suboxic diagenesis [Schaller et al., 2000] and lowtemperature fluid flow [Mills and Dunk, 2010]. Furthermore,

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in sediments with low accumulation rates, bioturbation and sediment redistribution smooth the record of past changes.

[3] In the abyssal South Pacific the sediment records are significantly impacted by hydrothermal inputs, and generating a record of past change in this setting has remained a challenge. The southern East Pacific Rise (SEPR) between 13 and 18°S is a site of intense hydrothermal activity [Lupton and Craig, 1981], and particle-rich plumes are present along more than half of the ridge axis in this region [Feely et al., 1996]. The metalliferous sediment distribution in this region of the SE Pacific is controlled by westward flow at ~2500 m depth [Reid, 1982] which transports the hydrothermal plume over hundreds of km [Dymond, 1981]. Systematic downcore variation in oxyanion [Poulton and Canfield, 2006] and trace metal [Schaller et al., 2000] content has been observed in young crustal and ridge flank sediments from this sector. In particular, subsurface uranium concentration peaks are common along the EPR from 10°S [Schaller et al., 2000] to 20°S [Shimmield and Price, 1988] (Figure 1). The origin and implications of these subsurface uranium enrichments in EPR sediments have been debated over several decades but not satisfactorily explained. Subsurface enrichments have previously been attributed to diffusive uranium uptake from seawater and fixation in suboxic sediments under higher-productivity glacial conditions [Yang et al., 1995], scavenging of seawater uranium by variable inputs of reactive hydrothermal plume particles [Shimmield and Price, 1988] and within-sediment transport

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Figure 1. Map of southern East Pacific Rise showing coring location at 15°S (GS7202-35 [*Rydell et al.*, 1974]). Ridge crest cores with subsurface U enrichments are shown (Y-71-7-53P [*Dymond*, 1981; *Schaller et al.*, 2000], RIS65 and 69 [*Veeh and Boström*, 1971], EXCO [*Dunk and Mills*, 2006], V19-54 [*Bender et al.*, 1971], and 154-18 [*Shimmield and Price*, 1988]) along with the coring location for DSDP Leg 92, hole 598 [*Poulton and Canfield*, 2006].

[*Mills and Dunk*, 2010] and precipitation from uranium-rich fluids [*Rydell et al.*, 1974; *Schaller et al.*, 2000]. Deciphering the origin of these uranium peaks would provide new information for an ocean region with sparse Pleistocene proxy records.

[4] Volumetrically the deep Pacific Ocean makes up a significant proportion of the global ocean and thus changes in Pacific circulation, ventilation and deep water nutrient, and carbon content can have a major impact on biogeochemical cycles. Most late Pleistocene reconstructions of the Pacific have focused on productivity variations [e.g., *Martinez and Robinson*, 2010] and changes to ventilation of the upper water column [e.g., *Stott et al.*, 2009]. Recent combined proxy and modeling approaches suggest reorganization of the North Pacific deep water ventilation during the last glacial termination may have had a significant impact on global heat transfer [*Okazaki et al.*, 2010] but there are limited proxy records to test this hypothesis, particularly in the south Pacific.

[5] Paleoceanographic records from the SE Pacific are sparse because carbonate sedimentation is restricted to the hydrothermally dominated ridge region. The published sedimentary records along the EPR lack age models and often the depth sampling resolution is inadequate. Our knowledge and understanding of past change would be significantly enhanced if the hydrothermal plume input and postdepositional changes within the sediment record could be deciphered and interpreted to reconstruct paleoceanographic and paleoclimatic conditions for this ocean.

[6] Here we present new age controlled data for an archived sediment core originally studied by *Rydell et al.* [1974] that demonstrates that diagenetic overprinting of hydrothermal inputs from the SEPR records significant glacial-interglacial changes in ventilation of the deep ocean moderated by intense redox cycling at glacial terminations II, III, IV and V during the Pleistocene. Comparison with published records along the SEPR suggests that these changes were basin wide and may thus significantly impact the storage of respired carbon in the deep ocean and poleward oceanic heat transport.

[7] The EPR ridge crest from 10 to 20°S underlies lowproductivity oligotrophic subtropical waters and is bathed by North Pacific Deep Water (NPDW: T < 2°C, $O_2 < 150 \mu M$ [*Fiedler and Talley*, 2006]) which recirculates southward from the North Pacific as the major return flow of the overturning circulation in this basin. This water mass underlies the well-ventilated intermediate modal waters sourced from the Southern Ocean [Fiedler and Talley, 2006]. The nutrient maximum and oxygen minimum associated with the NPDW outflow is centered at ~2000 m depth in the modern ocean. This return flow is hypothesized to have deepened to 2500-3000 m depth during the Last Glacial Maximum [Sigman and Boyle, 2000; Matsumoto et al., 2002] sequestering nutrients and respired carbon to the abyss and thus lowering atmospheric pCO₂ [Boyle, 1988]. This hypothesis has been tested extensively by attempts to reconstruct glacial Pacific nutrient distributions but has been hampered by apparently contradictory proxy records (e.g., δ^{13} C [Matsumoto et al., 2002] and Cd/Ca [Boyle, 1992]). Redox-sensitive metal records have been used in various ocean basins to identify glacial decrease in abyssal water oxygen content [e.g., Galbraith et al., 2007]. These different proxy records have recently been reconciled by the identification of an enhanced glacial respired carbon pool in the deep NW Pacific with lower oxygen levels but unchanged phosphate content arising from shifts in preformed nutrient supply [Jaccard et al., 2009].

[8] Glacial terminations are the times when the strongest changes in deep water ventilation and thereby oxygen levels, sedimentation rates, and input of electron donors occurs [*Broecker and Denton*, 1989]. Glacial terminations with global evidence for reconfiguration of ocean circulation occur at the ends of Marine Isotope Stage (MIS) 12 (V), 10 (IV), 8 (III), 6 (II) and 2 (I). There is evidence from the Pacific for basin-wide maxima in export production during early deglaciation which in turn impacts on bottom water oxygen content [*Martinez and Robinson*, 2010]. The inferred deglacial shifts in deep Pacific ventilation will also have significant controls on abyssal water oxygen levels [*Okazaki et al.*, 2010].

[9] At glacial terminations where benthic redox processes are perturbed, nonsteady state diagenetic processes are initiated which lead to postdepositional diagenetic overprint within these sediment intervals [Thomson et al., 1996]. Sedimentary redox driven diagenesis occurs throughout the redox gradient surrounding the oxidation front [Thomson et al., 1996]. Here we use the term oxidation front in a general way to describe the region just above and just below the limit of oxygen penetration, where a range of elements undergo redox cycling at different levels within the sediment in response to rapidly changing redox conditions. Solid phase Mn(IV) accumulates above the oxidation front in oxic conditions, supplied by reduced Mn(II)(aq) from below. Conversely solid phase, authigenic U(IV) accumulates under postoxic conditions within the oxidation front, supplied by dissolved U(VI) from above. Several other redox-sensitive elements are mobile and reactive within the oxidation front and can form local enrichment peaks between those of Mn and U [Thomson et al., 1996].

2. Sampling and Methods

[10] The core GS7202-35 was recovered from 82 km west of the EPR ridge axis at 14°47.9'S, 113°30.1'W from a

water depth of 3044 m and comprised trigger and piston cores 1.7 m and 8.47 m in length, respectively [Rydell et al., 1974]. The cores are archived in the Miami Core Repository at 4°C and were resampled in contiguous 5 cm strips (~4 kyr increments) which were dried, ground and homogenized prior to geochemical analysis. Bulk carbonate δ^{18} O and δ^{13} C were measured on a Europa Scientific GEO 20– 20 mass spectrometer and all data are expressed relative to the VPDP standard. Carbonate and organic carbon content were determined by difference using coulometry and by Carlo Erba CHN analysis. Major (Fe, Mn, Ca, Al, Si, Ti, P, K, Na) and trace element analyses (U, Mo, Cd, V, Cu, Ba, Br, I, S, Cl) were carried out using a Philips PW1400 fully automatic wavelength dispersive XRF. A subset of 25 samples was subjected to a total digestion procedure [Dunk and Mills, 2006] for analysis by Inductively Coupled Plasma-Atomic Emission Spectrometry. Analysis was carried out on a Perkin Elmer Optima 4300DV Precision ICPAES. Accuracy of these ICPAES analyses was assessed by comparison with USGS standard MAG-1 and was better than 1%. XRF pellets were calibrated using matrix-matched standards comprising mixtures of USGS standards MAG-1, JLs-1, GXR-1, NIM-L, BE-N, SDo1, NOD-P-1 and NOD-A-1. Accuracy of the XRF analysis was also assessed by comparison with the ICPAES data and for all elements presented (Table 1) is better than 3% of the certified value.

[11] We avoid issues relating to variable carbonate dilution downcore by reporting elemental ratios; all ratios are expressed in mass units (g/g). A comparison of the bulk sedimentary trace element/Fe ratios with the SEPR plume [*Feely et al.*, 1996] and oxic metalliferous sediments from east of the ridge axis at the same latitude [*Dunk and Mills*, 2006] (Figures 4e, 6c, 6d, and 6e) allows assessment of the inputs and postdepositional changes occurring with time. All data are for bulk sediment analysis including dried pore water salt.

3. Results and Discussion

3.1. Age Model and Sediment Accumulation

[12] The trigger and piston cores were combined into one continuous record using a combination of the downcore bulk δ^{18} O and trace element profiles. The best Pearson correlation for all records was achieved when the piston and trigger core were overlapped by 27.5-32.5 cm; all piston core depths have therefore been adjusted by +135.25 cm relative to the published stratigraphy which assumed erroneously that the piston core sampled the sediment water interface [Rvdell et al., 1974]. An age model for the sediment accumulation was constructed by correlation of the bulk sediment δ^{18} O record with the astronomically tuned composite time series obtained from cores V19-30, ODP 677 and ODP840 [Shackleton et al., 1990] using the Analyseries software [Paillard et al., 1996]. We estimate the uncertainty of the correlation to be <10 kyr, with a clear identification of all glacial and interglacial periods in the records (Figure 2a).

[13] The 9.8 m combined core contains a 740 kyr record which spans Marine Isotope Stages (MIS) 1 to 20 [*Shackleton et al.*, 1990] with linear sedimentation rates

Table 1. Downcore Age and Bulk Geochemical Data for Core GS7202-35^a

Depth (cm)	Age (ka)	Fe (%)	Mn (%)	Ca (%)	Al (ppm)	Si (%)	P (%)	U (ppm)	Mo (ppm)	Cd (ppm)	V (ppm)	Cu (ppm)	Ba (ppm)
5.25	4.5	5.92	1.68	28.5	700	1.34	0.286	5.0	8.6	11.8	178	245	1294
15.5	13.3	4.97	1.50	29.7	738	1.29	0.292	7.1	7.0	18.4	140	173	1101
20.5	18.3	5.84	1.65	28.1	670	1.33	0.314	6.7	8.5	16	169	237	1222
25.5	22.8	4.69	1./1	29.0	/33	1.14	0.244	4.8	10.5	10.4	134	211	1201
35.5	31.8	633	2.00	28.0	801	1.37	0.324	8.2	19.0	22.5	209	211 244	1319
40.5	36.3	5 30	1.96	28.6	691	1.47	0.407	8.5	15.8	18.4	164	200	1210
45.5	40.8	4 89	1.78	29.5	742	1.20	0.286	8.2	10.8	30.6	150	174	1189
50.5	45.3	4.96	1.78	29.9	746	1.23	0.287	8.3	11.7	22.8	146	178	1154
55.5	49.8	5.14	1.81	29.7	763	1.30	0.311	8.1	11.2	17.9	157	182	1284
60.5	54.4	5.69	1.97	28.6	792	1.47	0.372	8.3	14.2	16.4	182	204	1294
65.5	58.9	6.15	2.14	28.0	704	1.38	0.403	8.2	19.2	21.1	191	239	1287
70.5	63.4	6.04	2.26	28.2	717	1.35	0.375	7.4	21.7	20.9	203	237	1397
75.5	67.9	5.96	2.21	28.7	771	1.35	0.343	7.3	21.7	30.5	195	236	1478
80.5	72.4	5.83	2.07	28.8	830	1.31	0.307	4.7	19.7	14.4	195	225	1497
85.5	76.9	6.04	2.13	28.4	851	1.34	0.289	4.9	22.8	18.7	208	237	1626
90.5	81.4	5.99	2.16	28.7	818	1.30	0.260	5./	24.1	19	207	251	1764
95.5	83.9 00.4	5.88	2.45	27.0	803 720	1.57	0.262	5.7	29.7	15.4	230	265	1/04
115.5	103.9	5.88 6.01	2.27	28.6	746	1.25	0.201	5.0	28.1	20.3	108	264	1465
120.5	103.9	6.76	2.10	20.0	822	1 38	0.281	5.4	39.0	20.5	246	323	1837
125.5	113	6.17	2.34	28.2	860	1.30	0.258	41	40.5	21	220	294	1812
135.5	121.9	5.88	2.12	28.6	969	1.28	0.241	3.6	33.9	24.1	205	264	1667
138	123.9	5.59	1.99	29.0	910	1.19	0.222	4.0	32.5	20.3	183	252	1520
140.5	126	5.78	1.92	29.0	914	1.21	0.222	4.0	23.7	21.6	188	248	1442
143	128	5.69	1.93	29.1	885	1.20	0.224	6.0	19.6	6.0	184	250	1288
145.5	130	6.13	1.95	28.5	927	1.34	0.262	3.5	17.4	22.1	206	243	1403
148	132.1	6.72	2.10	28.3	897	1.27	0.279	8.3	21.7	9.2	218	303	1105
153	136.1	6.42	2.07	27.9	872	1.35	0.337	8.6	14.1	18.4	209	247	1165
155.5	138.2	5.53	1.88	29.1	801	1.27	0.325	8.3	/.1	13.2	172	196	1105
158	140.2	5.02	1.90	30.0 20.5	834	1.1/	0.304	86	18.5	27	1/0	224	1285
163	142.2	5.10	2.01	29.5	855	1.23	0.331	0.0 12.6	7.9	20.9	174	107	1231
165 5	146.3	5.95	2.01	28.5	784	1 39	0.360	11.3	7.3	20.9	190	208	1283
168	148.3	6.25	2.14	27.6	738	1.39	0.501	12.6	7.9	20.9	190	228	1231
178	156.5	5.91	2.21	28.0	717	1.34	0.463	11.9	5.7	22.1	192	213	1274
188	164.6	4.98	1.99	29.5	754	1.18	0.294	8.2	4.6	16.1	161	178	1334
193	168.7	5.05	1.96	30.4	733	1.06	0.241	6.9	3.8	22.8	148	204	1357
198	172.7	4.91	2.04	29.7	746	1.14	0.247	5.6	4.5	16.4	160	183	1385
203	176.8	4.89	2.06	29.7	742	1.12	0.231	5.7	3.6	15.2	154	186	1355
208	180.9	5.23	2.18	30.0	834	1.20	0.263	4.3	4.6	22.8	175	224	1498
213	184.9	5.08	2.06	29.4	/33	1.1/	0.278	4.8	2.7	18.1	105	199	1398
223	191	5.70	2.20	29.5	720	1.11	0.272	4.7	4.3	25.5	107	261	14/4
223	197.1	6 14	2.15	28.8	965	1.04	0.220	43	5.1	25.5	218	302	1773
238	200.2	6.47	2.72	28.3	897	1.37	0.299	4.3	6.5	33.7	236	334	1720
243	203.3	5.85	2.37	29.5	750	1.09	0.236	4.5	5.4	3.1	197	302	1375
248	206.3	6.32	2.53	28.8	944	1.26	0.273	5.8	4.3	23.9	221	326	1460
253	209.4	5.71	2.32	29.6	960	1.14	0.233	3.6	4.5	21.1	191	292	1287
258	212.4	5.62	2.31	29.4	1032	1.21	0.237	4.3	4.4	22.2	186	282	1388
263	215.5	5.63	2.34	29.8	805	1.12	0.243	4.3	4.5	13.8	185	280	1427
268	218.5	5.87	2.80	28.6	906	1.34	0.333	5.3	9.8	20.6	214	282	1520
273	221.6	5.79	2.52	29.1	/33	1.12	0.312	6.3	7.2	10.2	191	265	1343
2/8	224.0	0.12 5.20	2.02	29.1	091 754	1.11	0.310	5./ 2.5	8.1	18.5	219	280	13/0
283	227.7	5.39	2.48	29.8	734	1.15	0.240	5.5	/.4	52.9 10.2	214	249	1373
200	230.7	5.67	2.99	29.5	633	1.05	0.210	3.5	53	6.8	214	327	1331
298	236.8	7 38	2.57	28.0	574	1.13	0.203	6.0	4.6	16.7	265	370	1285
303	239.9	7.71	2.69	27.6	624	1.33	0.324	8.0	4.9	16.9	274	380	1318
308	242.9	4.43	1.55	31.7	591	0.92	0.211	8.1		12.9	122	176	944
313	246	4.64	1.75	30.7	775	1.19	0.287	11.3		28.2	136	187	1106
318	249	4.40	1.60	30.5	662	1.07	0.231	7.5		21.0	134	157	1055
333	258.2	4.27	1.63	30.5	696	1.13	0.296	9.9		26.3	124	154	1123
338	261.2	4.66	1.82	30.7	754	1.23	0.335	9.7		18.9	143	189	1120
343	264.3	5.13	1.97	29.9	771	1.32	0.414	11.9	1.0	20.5	162	211	1235
348 252	267.3	6.08	2.34	28.5	/92	1.44	0.484	14.8	1.8	24.1	198	259	1307
333 350	2/0.4	0.05	2.38	27.9	833 807	1.47	0.45/	12.2	1.4	20.4 22.9	220	293	1458
363	∠13.4 276.5	7 10	2.00	27.9 27.3	07/ 907	1.49	0.409	9.5 10.2	2.3 2.2	23.0 38.0	237	309	1554
505	410.0	1.17	2.01	41.5	104	1.01	U. T2J	10.4	<i></i>	50.7	∠- т 0	555	101/

Table 1. (continued)

Depth (cm)	Age (ka)	Fe (%)	Mn (%)	Ca (%)	Al (ppm)	Si (%)	P (%)	U (ppm)	Mo (ppm)	Cd (ppm)	V (ppm)	Cu (ppm)	Ba (ppm)
373	282.6	6.46	2.69	28.2	725	1.39	0.374	8.4	2.4	27.7	230	309	1456
378	285.6	5.78	2.29	29.6	582	1.09	0.316	8.0	2.2	18.7	184	272	1182
383	288.7	5.74	2.30	29.4	662	1.24	0.349	6.6	2.3	26.6	196	262	1230
388	291.7	5.8/	2.51	29.1	691	1.24	0.317	0.0	2.2	20.3	206	275	1318
393	294.8	5.88 5.79	2.62	29.0	763	1.25	0.292	/.0	3.5 5.0	9.5	210	281	13//
398	297.8	5.78	2.39	29.2	703	1.23	0.280	0.8	3.0	20.4	210	270	1365
403	303.9	5.60	2.34	20.0 28.4	675	1.24	0.293	83	3.4	20.4	201	251	1331
408	303.9	5.08	2.41	20.4	679	1.23	0.339	0.5 7.8	3.4	23.0	201	231	1296
413	310	6.42	2.30	28.6	889	1.31	0.336	6.2	2.1	16.5	221	207	1421
423	313 1	6.65	2.57	28.3	729	1.30	0.377	8.6	2.1	25.7	233	307	1381
428	316.1	6.08	2.26	29.3	775	1.31	0.314	6.2	2.0	31.0	213	271	1400
433	319.2	5.52	2.06	29.9	628	1.06	0.220	4.0	2.5	19.9	180	257	1197
438	322.2	5.69	2.39	29.3	683	1.23	0.250	3.8	5.3	21.0	201	268	1268
443	325.3	5.85	2.26	29.7	553	1.09	0.246	5.8	4.9	37.5	195	279	1160
448	328.3	6.76	2.06	29.1	507	1.16	0.319	6.3	1.9	21.6	217	314	922
453	331.4	7.67	2.45	26.1	599	1.54	0.431	7.0		25.4	287	329	1124
458	334.4	6.35	2.07	28.0	607	1.30	0.358	6.4	1.5	20.5	225	262	874
463	337.5	5.81	2.05	28.6	662	1.26		3.8		16.1	209	221	916
468	340.5	5.55	1.95	29.0	649	1.20	0.309	5.5		24.8	194	203	946
473	343.6	5.30	2.06	29.9	687	1.16	0.331	6.6	1.6	21.4	177	211	1002
478	346.7	5.17	2.02	29.9	708	1.19	0.338	7.6	1.2	31.4	174	207	1039
483	349.7	5.48	2.21	28.7	645	1.21	0.388	10.4		9.0	180	203	998
488	352.8	5.85	2.17	28.1	679	1.29	0.476	14.3		18.2	189	217	1020
498	358.9	5.31	1.93	30.4	696	1.07	0.328	12.9		14.1	158	226	989
503	362	5.83	2.08	29.7	712	1.15	0.365	14.8		8.3	176	253	1119
508	365.1	6.53	2.45	28.6	729	1.23	0.391	14.7	1.7	36.7	203	297	1250
513	368.1	6.67	2.69	26.5	7/1	1.38	0.402	12.1	1.4	22.6	221	282	13/4
518	3/1.2	0.4/ 6.66	2.85	27.0	/05	1.57	0.3//	10.7	2.3	32.0	224	284	1401
525	374.3	6.37	2.70	27.9	834	1.41	0.343	9.7	2.0	24.7	231	303	1420
533	380.4	6.47	2.00	20.1	780	1.37	0.328	7.0	1.9	10.5	217	203	1453
538	383.5	6.92	2.00	27.0	826	1.30	0.302	11.4	2.5	23.4	228	347	1455
543	386.5	6.62	2.85	27.7	775	1.41	0.251	73	1.8	22.4	240	323	1533
548	389.6	8 81	3.89	24.2	780	1.65	0.279	7.6	4 4	26.6	343	477	1816
553	392.7	8.50	3.62	24.9	830	1.61	0.273	8.2	2.5	24.6	325	447	1792
558	395.7	7.93	3.53	24.7	708	1.49	0.251	8.5	3.0	30.6	304	412	1622
568	401.9	5.42	2.39	29.8	725	1.18	0.214	6.9	2.5	10.3	191	263	1288
573	404.9	5.18	2.10	30.0	540	0.96	0.196	5.7		21.4	174	220	1092
578	408	6.03	2.17	29.8	612	1.07	0.231	8.5		23.7	197	271	1070
603	423.8	5.56	2.16	28.6	637	1.25	0.425	13.3		10.6	180	198	1011
613	430.1	4.67	1.86	30.0	641	1.10	0.286	9.4		21.1	145	167	906
623	436.5	5.74	2.32	28.7	662	1.29	0.397	16.8		10.5	191	215	1018
628	439.6	6.25	2.61	27.6	637	1.42	0.538	27.9		10.7	210	238	1120
633	442.8	6.42	2.55	28.0	700	1.48	0.505	30.3		20.2	223	262	1142
638	445.9	5.64	2.29	28.2	637	1.32	0.453	26.2		18.6	179	211	990
648	452.3	5.32	2.07	29.4	557	1.05	0.335	14.1		22.1	166	212	942
653	455.4	5.20	2.12	29.4	645	1.1/	0.3/6	13.4	1 1	20.5	164	202	957
663	458.0	5.41	2.24	29.0	075 725	1.18	0.364	12.5	1.1	20.9	1/5	214	1070
668	401.7	5.95	2.41	28.0	723	1.20	0.404	17.0	1.8	19.3	199	244	1233
673	404.9	7.60	2.90	27.1	/00	1.57	0.345	14.0	1.0	21.4	233	354	1233
683	408.1	8.55	3.01	23.5	826	1.55	0.340	18.5	7.5	20.3	327	408	1505
688	477.6	7.84	3.06	26.0	809	1.58	0.296	9.9	2.2	13.0	299	372	1274
693	482.5	7.48	2.88	26.6	750	1.50	0.290	11.4	18	26.0	288	359	1179
698	487.7	7.92	3.05	25.9	712	1.54	0.319	13.7	110	16.6	302	384	1197
703	493	8.40	3.28	25.3	721	1.62	0.361	14.7	2.7	24.3	332	416	1309
708	498.3	7.59	3.10	26.2	717	1.52	0.325	12.4	1.0	12.9	302	364	1203
713	503.6	7.31	2.82	26.7	754	1.49	0.297	10.7		24.5	287	331	1076
718	508.8	7.58	2.93	26.4	729	1.47	0.305	12.3	1.4	17.3	297	341	1113
723	514.1	8.06	3.11	25.9	733	1.55	0.343	12.9		17.3	320	368	1190
728	519.4	8.85	3.41	24.6	733	1.74	0.446	17.3	1.8	21.4	352	420	1309
733	524.7	7.64	2.96	26.1	687	1.53	0.425	16.1	1.5	9.2	293	346	1169
738	529.9	6.78	2.64	27.1	704	1.45	0.402	15.2	1.0	14.0	247	289	1087
743	535.2	6.45	2.59	27.6	704	1.39	0.386	14.6		19.3	236	266	1100
748	540.5	6.49	2.64	27.5	696	1.37	0.402	13.3	1.1	27.5	230	269	1101
753	545.8	6.92	2.91	26.7	746	1.47	0.434	14.0		25.3	259	287	1288
758	551	6.45	2.74	27.2	759	1.41	0.412	13.6		24.7	238	270	1220
763	5563	6.26	2.57	277	813	1.37	0.356	10.5		24 5	230	264	1226

 Table 1. (continued)

Depth (cm)	Age (ka)	Fe (%)	Mn (%)	Ca (%)	Al (ppm)	Si (%)	P (%)	U (ppm)	Mo (ppm)	Cd (ppm)	V (ppm)	Cu (ppm)	Ba (ppm)
768	561.6	6.72	2.76	27.0	792	1.42	0.395	13.1	1.4	17.7	243	296	1308
773	566.9	7.45	3.12	26.0	767	1.51	0.417	16.0	1.1	27.0	279	348	1448
778	571.7	7.89	3.59	25.6	784	1.54	0.360	14.0	3.8	23.0	319	403	1517
783	575.6	7.38	3.37	26.3	830	1.47	0.305	11.9	3.6	29.7	288	376	1516
793	583.3	7.32	3.44	26.0	729	1.50	0.401	16.6	5.7	18.0	286	370	1463
798	587.2	7.10	3.09	26.3	708	1.45	0.449	14.0	4.0	15.8	267	321	1290
803	591.1	6.59	2.87	27.3	725	1.36	0.368	11.8	2.4	17.6	243	294	1379
808	595	6.69	3.00	27.0	763	1.43	0.362	15.5	4.1	23.4	246	308	1329
813	598.9	7.04	2.89	26.7	830	1.47	0.348	14.3	2.1	25.1	259	321	1488
818	602.8	7.26	2.89	26.7	759	1.48	0.318	11.9	1.6	24.8	274	336	1465
823	606.6	7.40	2.96	26.6	754	1.44	0.273	10.8	1.1	26.0	296	355	1491
828	610.5	7.42	2.93	26.6	792	1.49	0.276	10.4	1.4	36.5	285	354	1435
833	614.4	8.96	3.45	24.6	830	1.74	0.352	14.3	2.3	21.0	352	453	1465
838	618.3	10.44	4.03	22.5	733	1.96	0.540	18.8	3.9	28.5	420	538	1435
843	622.2	8.78	4.65	23.8	687	1.68	0.463	15.6	13.1	27.8	360	415	1461
848	626.1	6.90	4.57	26.0	679	1.43	0.396	11.2	16.8	22.6	269	304	1303
853	630	6.35	2.39	27.8	662	1.33	0.411	10.1		12.5	223	210	947
858	634.2	6.49	2.51	27.6	670	1.37	0.450	11.4	1.0	29.2	231	220	985
863	638.5	6.19	2.35	28.0	754	1.36	0.435	9.0		28.7	214	205	1019
868	642.7	6.41	2.50	27.7	704	1.34	0.437	11.4	2.3	17.8	221	210	1067
873	647	7.75	6.24	23.9	742	1.58	0.493	14.8	30.8	29.4	327	363	1596
878	651.2	8.49	3.06	24.9	717	1.67	0.569	12.9	3.3	22.4	313	310	1157
883	655.5	7.82	3.14	25.7	754	1.61	0.488	11.1	3.0	22.0	293	298	1260
888	659.7	7.08	2.70	26.9	704	1.39	0.436	9.2	2.2	27.1	253	269	1181
893	664	6.84	2.46	27.3	767	1.39	0.372	8.6	1.5	19.8	245	261	1215
898	668.2	7.18	2.58	27.0	788	1.42	0.362	7.9	1.3	26.2	270	286	1340
903	672.5	9.08	3.34	24.6	805	1.70	0.451	10.5	3.4	24.7	368	408	1481
908	676.7	8.33	3.24	25.4	881	1.62	0.342	9.5	2.6	24.3	344	392	1575
913	681	9.11	3.53	24.5	864	1.63	0.321	10.5	3.2	26.2	374	457	1640
918	685.2	10.38	3.95	22.7	822	1.92	0.477	12.7	4.1	25.8	441	548	1569
923	689.5	8.22	3.53	25.3	822	1.62	0.338	11.0	5.2	10.0	341	419	1367
928	693.7	8.18	3.12	25.6	805	1.62	0.336	8.9	1.8	23.0	340	392	1285
933	698	7.42	3.02	26.4	847	1.52	0.319	10.7	2.7	25.8	298	351	1266
938	702.2	7.43	3.21	26.2	872	1.56	0.356	11.1	2.5	21.3	285	344	1241
943	706.5	7.57	3.19	25.6	1078	1.73	0.471	17.5	2.7	35.3	280	328	1202
948	710.7	6.66	2.77	26.6	1162	1.63	0.432	11.6	2.0	25.0	228	270	1075
953	715	7.19	3.06	26.0	1007	1.61	0.480	10.0	4.5	7.2	263	294	1073
958	719.2	7.00	3.33	26.2	923	1.53	0.447	8.6	6.3	15.9	266	298	1081
963	723.5	7.52	2.72	26.2	1091	1.70	0.500	9.9	1.9	16.8	284	263	965
968	727.7	6.69	2.84	27.0	1082	1.58	0.393	8.4	2.7	22.8	255	240	1076
973	732	6.58	4.07	26.5	927	1.50	0.337	10.3	13.8	23.3	275	296	1354
978	736.3	7.47	3.42	25.8	893	1.69	0.403	9.7	5.7	24.0	311	301	1215
983	740.5	7.92	3.94	25.0	868	1.76	0.457	8.8	9.7	21	341	353	1378

^aData gaps indicate measurement below detection limit.

(LSR) of 0.95-1.64 cm kyr⁻¹ (Figure 2c). There is no systematic glacial-interglacial change in inferred LSR but there is a notable decrease in LSR at the end of MIS 7, when the core location was ~68 km from the ridge axis. The lower sedimentation rate in the upper core ($\sim 1 \text{ cm kyr}^{-1}$) means that glacial-interglacial differences are not as clearly resolved as those deeper in the core. The observed sedimentation rates are consistent with estimates from 230 Th_{xs} decay in the upper 3 mbsf [Rydell et al., 1974] and estimated mean accumulation from seismic profiling [Hauschild et al., 2003]. At 15°S the sediment appears to drape the seafloor uniformly with little indication of lateral sediment transport [Hauschild et al., 2003]. There is little basement topographic variation and therefore minimal basal fluid advection through the sediment on this ridge flank compared with the eastern flank at this latitude [Mills and Dunk, 2010]. Based on estimated spreading rates [Grevemever et al., 2002], core GS7202-35 was located ~25 km west of the

EPR axis at 740 ka and has moved a further 57 km off axis since that time.

[14] Pleistocene sedimentary records are characterized by the mid-Pleistocene transition (MPT) which is manifest in the Pacific by significant changes in the bulk carbon isotope composition (Figure 2b) [Shackleton and Hall, 1995] and marked shifts in the inferred deep water ventilation by the South Pacific upper circumpolar deep water (UCDW) at 870 ka and 450 ka [Venuti et al., 2007]. The early deposition at site GS7202-35 (740-400 kyr) occurred during the MPT. The major shift in bulk carbon isotope composition recorded in Equatorial Pacific sediment has been interpreted as evolutionary succession in the nanofossil assemblage [Shackleton and Hall, 1995]. The bulk δ^{13} C record for this core exhibits the main features seen in the Equatorial Pacific Pleistocene record, providing additional stratigraphic constraints for the proposed age model (Figure 2b). There is evidence for a number of shorter-lived shifts in carbon



Figure 2. Plots of (a) the δ^{18} O for the benthic foraminifera composite (gray line) [*Shackleton et al.*, 1990] with the bulk δ^{18} O data for this study (black dots and solid line), (b) the bulk δ^{13} C composition for ODP leg 138 eastern equatorial Pacific (gray line) [*Shackleton and Hall*, 1995] with the bulk δ^{13} C data for this study (black dots and solid line), and (c) the Linear Sedimentation Rate (LSR) inferred from the age model presented in Figure 2a. Distance from ridge axis is estimated from sediment age and spreading rates [*Grevemeyer et al.*, 2002]. Grey shading indicates glacial stages; numbers indicate marine isotope stages (MIS) [*Imbrie et al.*, 1984].

isotope partitioning at this site at the ends of MIS 12, 10 and 8 (Terminations V, IV and III; Figure 2b).

3.2. Sediment Composition

[15] The sediment of core GS7202-35 comprises a mixture of hydrothermal material derived from plume fallout (Fe sulfides and Fe and Mn oxyhydroxides) and biogenic carbonate (CaCO₃) with minor biogenic silica and aluminosilicates (Table 1 and Figure 3). The maximum lithogenic (basaltic) input at this site estimated from the bulk sediment Al content and the Al content of basalt at this latitude (Al = 8.88% [Schramm et al., 2005]) is small (0.52–1.57% of bulk sediment) and this component can be ignored in our discussion. Note that the highest basaltic inputs are observed toward the base of the core when the site was closer to the ridge axis. The biogenic carbonate content varies from 58 to 72% (Figure 4a) with lower values in the deeper parts of the core that are diluted by the more intensive hydrothermal input (up to 40%) when the core location was closer to the ridge axis. The main control on CaCO₃ burial away from the equatorial productivity maximum is inferred to be carbonate preservation rather than changes in productivity [Murray et al., 2000; Anderson et al., 2008]. The characteristic changes in CaCO₃ preservation are damped at the shallow ridge crest and the dramatic glacial-interglacial fluctuations seen else-



Figure 3. Plot of Fe (%), Mn (%), Si (%), and Al (%) against CaCO₃ content for core GS7202-35.



Figure 4. Plot of (a) CaCO₃, (b) Fe, (c) Mn, (d) Mn/Fe ratio, and (e) Mo/Mn ratio against age. Dashed horizontal lines indicate the end-member hydrothermal Mn/Fe ratio for Nazca Plate ridge crest sediments [*Dymond*, 1981] and oxic sediment Mo/Mn ratio [*Shimmield and Price*, 1986; *Dunk and Mills*, 2006]. Arrows indicate glacial terminations and the active and relict oxidation fronts (see text).

where in CaCO₃ records in Pacific cores retrieved from greater water depths (>4200 m [*Murray et al.*, 2000]) are not observed in this shallower ridge flank 15°S record. However, the significant dissolution event associated with MIS 11 [*Murray et al.*, 2000] is associated with a minima in CaCO₃ content in this record (Figure 4a). The hydrothermal material preserved in the sediment largely consists of X-ray amorphous Fe and Mn oxides with a minor clay component (<1%). Consideration of the sea salt elements S, Na, Cl, Br indicate the presence of 100–2500 ppm excess S content relative to Cl throughout the core, and Fe sulfides have been observed by scanning electron microscopy of impregnated sections of sediment from the ridge crest at this latitude [*Dunk and Mills*, 2006]. So while sulfide oxidation will

have occurred during 25 years of core storage, the presence of primary sulfides at this site is inferred from the excess S measured by XRF. The organic carbon content of the sediment is low throughout (<0.2%) with no measurable variation with depth; these low values are consistent with published data from this oligotrophic region [*Schaller et al.*, 2000].

[16] Plume particles collected from the neutrally buoyant plume overlying the superfast spreading EPR consist of a mixture of Fe and Mn oxides and sulfide/elemental sulfur phases [*Feely et al.*, 1996]. Plume particles collected from the neutrally buoyant plume ~25 km west of the ridge axis at 15°S contain significant particulate sulfur (~30 nmol/L; particulate molar S/Fe ~0.04) and particulate Mn (1–2 nmol/

L; particulate molar Mn/Fe up to 0.1 [Feely et al., 1996]). Plume particles collected from further south on the EPR contain a particulate component characterized by high levels of reduced S, particulate Mn and methane [Feely et al., 1996]. Plume transported hydrothermal sulfides are inferred to contribute significant transition metals to sediments from 10°S and elsewhere on the EPR [Schaller et al., 2000; Poulton and Canfield, 2006]. Thus the two sources of electron donor to the sediment at this site are organic carbon export from overlying oligotrophic waters and hydrothermally derived reduced sulfur species. The electron donor input to the sediment is inferred to be cyclical, either through glacial-interglacial variation in overlying productivity and carbon export or by variations in hydrothermal inputs from the EPR which in turn are moderated by bottom water oxygenation which is determined by abyssal ventilation rates.

3.3. Fe and Mn Diagenesis

[17] The core is highly enriched in Fe (4.2-12%) and Mn (1.4–6.9%) throughout (Table 1 and Figures 3, 4b, and 4c). The Fe content covaries inversely with the CaCO₃ content (Figure 3) so that the dominant control on Fe content is the variable dilution with CaCO3 downcore. Mn/Fe ratios are variable and high (0.24-0.47; Figure 4d). These values are higher than oxic sediments east of the axis at this latitude (EXCO: Mn/Fe = 0.21-0.33 [Dunk and Mills, 2006]) and elsewhere on the EPR (10°S: 0.17-0.33 [Schaller et al., 2000], 20°S: 0.29–0.35 [Shimmield and Price, 1988]). These Mn/Fe ratios are also higher than the inferred hydrothermal sediment end-member calculated for Nazca Plate metalliferous ridge crest and flank sediments (Mn/Fe = 0.323 [Dymond, 1981]) throughout the core below 25 cmbsf, although Mn/Fe values approach this modern sedimentary end-member value at glacial terminations V, IV, III and II (Figure 4d). The source of additional Mn is inferred to be the hydrothermal plume inputs associated with superfast spreading hydrothermalism as characterized by Mn- and S-rich particles collected on the SEPR [Feely et al., 1996].

[18] Dissolved Mn(II) is the product of dissimilatory Mn reduction during early diagenesis and Mn(II) can also be derived from in situ reaction of Mn oxides with plumederived sulfides [Aller and Rude, 1988; Schippers and Jørgensen, 2001]. The resultant Mn(II) is retained within the sediment both by sorption on carbonate surfaces and reprecipitation as diagenetic Mn(IV) phases in the oxic surface layer [Shimmield and Price, 1986; Thomson et al., 1986; Yang et al., 1995]. The relatively high Mn/Fe ratios compared with ridge crest and flank sediments from 10°S [Heath and Dymond, 1981; Schaller et al., 2000] indicate that the oxic surface layer at this site must always have been thick enough to trap a significant proportion of the upwardly diffusing Mn within the sediment for most of the last 740 ka. This is in contrast to sediment records from 10°S, 34 km from the ridge axis, where significant Mn loss is observed below 4 cmbsf (Mn/Fe = 0.17-0.33 [Schaller et al., 2000]). There is no Mn redox cycling apparent in oxic sediments from the same latitude east of the ridge axis; sedimentary Mn/Fe ratios are relatively constant and no measurable Mn(II)(aq) is observed in pore waters [Dunk and Mills,

2006] although inferred carbon export and bottom water oxygen [*Fiedler and Talley*, 2006] is similar east and west of the ridge axis at 15°S. Thus the main control on redox status at the EPR sites is inferred to be the enhanced hydrothermal supply of reduced sulfide species on axis and the western flank that are subsequently oxidized in situ during early diagenesis.

[19] Hydrothermally derived Mo is associated with hydrothermal sulfides which are transported to the ridge crest and flank sediments. In situ oxidation of sulfides releases Mo to pore waters and the Mo content of postoxic sediments is subsequently controlled by the redox cycling of Mn within the oxidation front [Schaller et al., 2000]. The molybdate oxyanion (MoO_4^{2-}) has a high affinity for MnO_2 and Mo is remobilized when the MnO₂ substrate is reduced during early diagenesis [Shimmield and Price, 1986]. The Mo/Mn ratio of oxic deep sea sediments and ferromanganese deposits is relatively constant (0.002 [Shimmield and Price, 1986]) and the average ratio for oxic sediments (with lower hydrothermal Mn content) from the eastern EPR flank at 15°S is 0.0016 [Dunk and Mills, 2006]. The Mo/Mn ratio at 15°S does not reach oxic values at any depth within the core (Figure 4e). The hydrothermal supply of particulate Mn at this site [Feely et al., 1996] vastly exceeds the inferred Mo supply which is not adequate to achieve normal pelagic oxic Mo/Mn ratios even within the upper oxic layers: Mn is significantly enriched throughout the core. The low downcore Mo/Mn ratio indicates that significant Mn cycling (and preferential Mo loss) occurred while the core was relatively close to the ridge axis under postoxic conditions and Mo remobilization and accumulation in the upper core is apparent (Figure 4e). The elevated Mo/Mn ratios in the upper core are consistent with a shift to more oxic sedimentary conditions with time which is inferred to relate to the distal accumulation of more oxidized hydrothermal material and an overall increase in Pacific ventilation and/ or decrease in carbon export since the mid-Pleistocene transition [Venuti et al., 2007].

[20] The active oxidation front in this core is inferred to be at ~140 cmbsf (120 kyr; Figure 4e) based on the Mo distribution and the maximum inflection in the Mo/Mn profile. The shape of the Mo/Mn profile is consistent with progressive deepening of the oxidation front to 140 cmbsf and Mo preferentially accumulates above the oxidation front over time as continued Mo and Mn cycling occurs. Metastable remnants of relict Mo/Mn peaks are observed in MIS 7 and 9 (Figure 4e). Deep (>1 m) oxidation fronts have previously been identified in Atlantic cores where nonsteady state diagenesis is controlled by turbidite emplacement with additional organic carbon supply to the seafloor [Colley and Thomson, 1985] and deep oxygen penetration (several meters) is observed in hydrothermal sediments collected from the same latitude on the eastern flank of the EPR [Dunk and Mills, 2006]. Cores underlying the oligotrophic South Pacific gyre show oxygen penetration to basement (~8 m depth [Fischer et al., 2009]). The slow sediment accumulation in the upper core and the inferred low electron donor input relative to deeper in the core mean that while similar processes may well occur transiently at the end of MIS 2 (termination I), evidence for nonsteady state redox cycling



Figure 5. Plot of (a) U against age, (b) Cd against age, (c) P (open circles) and K (filled circles) against age, (d) Cu against age, and (e) V against age. All shading as for Figure 2.

is not preserved in the upper sediment record and the active oxidation front has now penetrated to termination II (140 cmbsf).

[21] Plume-derived hydrous Fe oxides which include ferrihydrite phases will undergo transformation during oxic and suboxic diagenesis to a range of more crystalline Fe rich phases [*Dunk and Mills*, 2006]. Phase transformation reactions exclude large cations and oxyanions from the more crystalline Fe oxide structures and diffusive transport and reaction of these elements can overprint the primary sedimentary record. The in situ oxidation of Fe sulfides by a range of electron donors including O_2 , NO_3^- and MnO_2 [*Aller and Rude*, 1988] leads to ferrihydrite formation during early diagenesis and this ferrihydrite transforms to crystalline phases over time, further complicating the diagenetic history of this core.

3.4. Trace Metal Diagenesis

[22] The U content in core GS7202-35 is extremely high compared with normal hydrothermal sediments (Figure 5a) with maximum values found consistently in sediments laid down during glacial periods, and a peak of 30 ppm at 6.33 mbsf (443 kyr, MIS 12). Previously published $^{234}U/^{238}U$ ratios [*Rydell et al.*, 1974] are within error of the age corrected seawater activity ratio (1.14 [*Chen et al.*, 1986]) in the upper 4.5 m of the core, indicating a seawater origin for the sedimentary U. Uranium has been shown to be elevated on hydrothermal sulfide surfaces [*Mills et al.*, 1994] and in sulfidic plume particles on the northern EPR [*German et al.*, 2002]. Thus fine-grained plume-derived sulfides are a major source of U input to metalliferous sediments.

[23] The Cd content is also significantly elevated throughout the core (10–40 ppm; Figure 5b), reflecting the



Figure 6. Elemental ratio plots for core GS7202-35 plotted against age. (a) U/Fe ratio (g/g), (b) P/Fe ratio (g/g), (c) Ba/Fe ratio, (d) V/Fe ratio (g/g), and (e) Cu/Fe ratio (g/g). Arrows indicate regions of U burn down and the active and relict oxidation fronts (see text). Dashed lines indicate the U/Fe ratio for oxic metalliferous sediments [*Mills et al.*, 1994], Ba/Fe for the plume-derived component of EXCO sediments [*Dunk and Mills*, 2006], V/Fe for oxic EPR sediments [*Dunk and Mills*, 2006], and Cu/Fe for the oxic SEPR end-member [*Dymond*, 1981] and oxic EPR sediments [*Dunk and Mills*, 2006]. Shaded area in Figure 6b indicates the range in observed P/Fe for SEPR sediments (see text). All other shading as for Figure 2.

plume sulfide input at this site. Trans-Atlantic Geotraverse (TAG) plume sulfides have Cd contents of 10–500 ppm [*Metz and Trefry*, 1993] and in situ oxidation of sulfides will lead to Cd association with oxyhydroxide phases and retention in the sediment. Comparison with a ridge crest core at 10°S [*Schaller et al.*, 2000] implies that the Mo, U and Cd are all primarily sourced from sulfidic plume input and are variably mobile during suboxic diagenesis.

[24] The U/Fe ratio for oxic ridge flank metalliferous sediments is relatively constant and low (U/Fe = 0.3×10^{-4} [*Mills et al.*, 1994]) and the U/Fe ratio is enhanced relative to this value throughout most of this core (Figure 6a). At the active oxidation front (120 kyr), U is inferred to have been relocated into the underlying glacial section by oxidation [*Colley and Thomson*, 1985]. Oxygen penetration and oxi-

dation remobilizes a fraction of plume-derived U that is reprecipitated as U(IV) below the oxidation front in MIS 6 (Figure 6a). The U/Fe peaks in the upper oxic core are inferred to be nonsteady state, metastable features that result from more recent hydrothermal sedimentation during MIS1– 5. The decreased plume-derived electron donor supply as the core moves off axis and the slower sediment accumulation rate (Figure 2c) allow oxygen burn down to MIS 6. The U/Fe ratio approaches the low oxic sedimentary value toward the end of MIS 6, 8, 10 and 12 (Figure 6a) indicating that oxic burn down and redistribution of U occurred after glacial terminations V, IV, III and II as the electron donor supply to the seafloor waned and/or the bottom water oxygen content increased during the following interglacial. Maximum U/Fe ratios occur during glacial periods and in



Figure 7. Schematic representation of pore water and solid phase response to an interglacial-glacial-interglacial transition through time from left to right. Interglacial oxygen penetration is relatively deep, and Mn^{2+} , and other redox-sensitive metals and those associated with Fe-Mn oxyhydroxide phases, are recycled below this depth. The glacial shift to a shallower oxidation front leads to significant Mn and Mo remobilization from the solid phase and relocation of these elements above the new oxidation front. Differential mobility of Mn and Mo, and the significant hydrothermal Mn input at this site, leads to variable Mo/Mn ratios within the shallow oxic zone. Lower bottom water oxygen levels and a shallower oxidation front lead to enhanced plume U supply to the seafloor during glacial periods. Glacial termination is inferred to be associated with a pulse of carbon supply to the seafloor that shallows the oxidation front further [*Martinez and Robinson*, 2010]. Subsequent deepening of the oxidation front during interglacial periods leads to oxygen burn down and relocation of the plume-derived U below the oxidation front.

particular the largest peak occurs at 443 ka in MIS 12 (Figure 6a). This largest U/Fe peak coincides with the MIS 12 ventilation minimum in the South Pacific [*Venuti et al.*, 2007] which is followed by the long, warm MIS 11 interglacial period. These mid-Pleistocene climatic extremes lead to maximal relocation of the deposited U.

[25] Diffusive uptake and fixation as U(IV) under suboxic conditions can conceivably lead to an influx of authigenic U in slowly accumulating sediments, but bioturbation and oxidation leads to net efflux of U from the sediments [McManus et al., 2005]. Significant authigenic U accumulation by this mechanism only occurs in sediments where the redox interface is within a few cm of the surface [Morford et al., 2005]. The surface oxic layer thickness at 15°S is inferred to have exceeded 5 cm to retain the Mn(II) within the sediment throughout much of the last 740 kyr and significant authigenic U uptake from seawater could only occur when the oxidation front is close to the sediment surface under glacial conditions. The main source of the U enrichment at 15°S is likely to be derived both from allochthonous inputs of seawater U associated with sulfidic plume particles [German et al., 2002] and enhanced by authigenic U uptake

during glacial periods. The subsurface U peaks record periods of nonsteady state diagenesis and burn down associated with post glacial decrease in electron donor supply to the sediment which is associated with increasing bottom water oxygen content. This interpretation challenges the assertion by *Rydell et al.* [1974] and *Schaller et al.* [2000] that U enrichment along this sector of the EPR arises from subsurface fluid flow. The inferred variation in redox status and impact on the Mo/Mn and U/Fe ratios over glacialinterglacial cycles is summarized schematically in Figure 7.

[26] Phosphorus and K content covary downcore (Figure 5c) and peaks occur in glacial stage sediments. Copper and V content also covary with depth and are inferred to be sourced from seawater and hydrothermal inputs and their distribution modified during early diagenesis (Figures 5d and 5e). These trace metal profiles will be discussed in turn to develop an understanding of the diagenetic overprinting on the inferred primary hydrothermal input.

[27] Phosphate is highly mobile during early diagenesis and the signature preserved is inferred to be a combination of the primary hydrothermal input overprinted by diagenetic processes. High P/Fe ratios coincide with glacial periods and



Figure 8. Comparison of U/Fe and P/Fe ratios downcore from (a) 10°S [*Schaller et al.*, 2000] and (b) the present study. High-precision organic carbon data for the 10°S site are plotted (dashed line). Shading indicates glacial periods for the 15°S core determined from the age model (Figure 2a).

lower P/Fe with interglacials (Figure 6b). The low sedimentary P/Fe ratio at this site and at 10°N (Figure 8) compared with SEPR plume particles (P/Fe = 0.1 [*Feely et al.*, 1996]) reflects the significant sulfidic plume input to this site of particles that have little affinity for seawater PO_4^{3-} [*Feely et al.*, 1996]. These low ratios are similar to P/Fe ratios at other sites along the SEPR (0.055 [*Marchig et al.*, 1986], 0.049 [*Schaller et al.*, 2000], and 0.044 [*Poulton and Canfield*, 2006]) which have also been attributed to dilution of plume-derived material with plume sulfides and P loss during oxic diagenesis that predominates in the upper sediment during interglacial periods. The higher glacial P/Fe ratios are consistent with exposure to a nutrient-rich, oxygen-depleted water mass during these periods.

[28] The close correlation between P and K at this site is striking and the most likely phase hosting K is hydrogenous Fe-hydroxyphosphates and minor phillipsite or other zeolite generated in situ during early diagenesis. However, there is no mineralogical evidence for the presence of any such crystalline phase in any abundance.

[29] Barium in hydrothermal sediments is dominantly associated with the Fe oxyhydroxide fraction with additional contributions from carbonate bound barium and biogenic barite from the upper water column [*Dymond*, 1981]. The Ba/Fe ratio of the hydrothermal input at 14°S has been determined by sequential leach analysis of metalliferous sediments (Ba/Fe = 0.013 [*Dunk and Mills*, 2006]) and the Ba/Fe ratio of core GS7202-35 is consistently higher than this ratio (Figure 6c) confirming the multiple sources of this element at this site. Low Ba/Fe ratios are associated with the minima in Mn/Fe (Figure 4d) and regions of U relocation (Figure 6a) which are relict oxidation fronts suggesting that Ba has been significantly redistributed during Mn and Fe recycling at this site and therefore cannot be used as a productivity proxy in this setting.

[30] Other redox-sensitive metals such as Cu and V are dominantly controlled by the recycling of MnO₂ (Figures 6d and 6e). Elevated Cu/Fe and V/Fe ratios correlate with maximum Mn/Fe ratios downcore and the relict oxidation fronts indicated by minimum Mn/Fe ratios correlate with minima in the Cu/Fe and V/Fe profiles. Redox cycling of MnO₂ releases Cu and V to pore fluids and they accumulate preferentially in the oxic interglacial regions overlying the active and relict oxidation fronts. The correspondence between low Mn/Fe and V/Fe ratios has previously been seen at 10°S where low V/Fe ratios are associated with Mn loss from the suboxic ridge crest sediments [*Schaller et al.*, 2000].

3.5. Mechanisms for Changing Depth of the Oxidation Front

[31] There are three main controls on the depth of oxidation front: bottom water oxygen content, electron donor supply, and sedimentation rate. We have demonstrated that sediment accumulation at this site does not control the observed nonsteady state diagenesis at terminations V, IV, III and II (section 3.1 and Figure 2c). Comparison of the suboxic sediment record at 15°S with elemental ratios for fully oxic sediments collected at the same latitude east of the ridge axis [*Dunk and Mills*, 2006] demonstrate the impact of plume-derived electron donors to the seafloor at this latitude. Hydrothermally derived electron donors have a major impact on the redox state of sediments in this region.

[32] Studies of productivity proxies at 5°S in the SE Pacific across glacial cycles including MIS 11 and 12 suggest no systematic change in export production through time but instead identify significant shifts in lysocline depth and carbonate preservation [*Murray et al.*, 2000; *Anderson et al.*, 2008] that are due to differences in bottom water carbonate ion activity related to ocean circulation changes at ~4200 m water depth. The ridge flank setting of the core studied here is well above the modern lysocline depth of ~3500 m and significant carbonate dissolution is unlikely. Records from the equatorial Pacific imply that pulses of enhanced

export occurred during glacial terminations [*Martinez and Robinson*, 2010]. Thus the primary control on oxidation front depth on the ridge flank at 15° S is likely to be glacial-interglacial changes in bottom water ventilation and oxygen content which in turn controls the supply of hydrothermal sulfides to the sediment moderated by short-term pulses in carbon export to the seafloor during glacial terminations. The low CaCO₃ levels in MIS 11 sediments suggest that carbonate dissolution may also occur in the surficial sediments at this time. It is not possible to quantify this dissolution effect, but it would be expected to have an effect similar to a decrease in accumulation rate in that it would lead to an increase in the oxidation depth in the sediments enhancing the apparent U accumulation.

[33] Lower glacial oxygen levels lead to a shoaling of the redoxcline in the sediment and increased accumulation of plume-derived sulfides and associated metals at the seafloor. Pulses of enhanced carbon export during glacial terminations lead to a shallow redoxcline. Mn cycling below the oxidation front leads to remobilization of oxyhydroxide bound metals (Mo, Cu, V) and reaccumulation above the oxidation front. Subsequent systematic deepening of the oxidation front occurs after glacial terminations V, IV, III and II, relocating the plume-derived U into the deeper suboxic sediment. Oxidation of the remnant plume-derived sulfides in situ provides a reactive ferrihydrite phase that effectively sorbs elements released during Fe oxide transformation reactions.

[34] One test of this hypothesis is comparison with data from other EPR locations with documented subsurface U enrichments (Figure 1). Few EPR cores have a Pleistocene age model and only two records have appropriate downcore sample resolution [Schaller et al., 2000; Shimmield and *Price*, 1988]. The former includes multielement data and a comparable sediment accumulation rate (~1.4 cm kyr⁻¹ [Schaller et al., 2000]) which can be compared with our data set (Figure 8). Downcore depth profiles of U/Fe and P/Fe for the upper ~ 2 m provide a record back to MIS 6 where the glacial age U and P peaks can be correlated between the two sites (Figure 8) supporting the case for a wider scale impact. The 10°S core is significantly more reducing (inferred active oxidation front at \sim 4 cmbsf) than the 15°S core because it is closer to the ridge axis (higher sulfide electron donor input) and underlies more productive waters at the periphery of the gyre. Thus the U/Fe ratios are higher (more U-rich sulfides supplied to sediment, less oxygen burn down and remobilization of U) whereas the P/Fe ratios are similar to 15°S. The upper core record at 10°S is dominated by redox cycling and metal flux (Mo, Mn, V) to the overlying water column [Schaller et al., 2000]. The high-precision organic carbon record for the 10°S site (Figure 8) is consistent with glacial shallowing of the oxidation front and enhanced preservation of the organic carbon flux to the seafloor during these periods.

[35] Comparisons with other cores are more problematic for the reasons stated above. For example, U peaks observed at 20°S previously attributed to enhanced hydrothermal scavenging [*Shimmield and Price*, 1988] are hypothesized to be associated with termination I deglaciation but in the absence of any Fe data and a poorly constrained age model this is impossible to test without resampling the archived cores. Comparisons with NW Pacific climate records from a similar depth (3244 m) suggest that glacial deepening of the oceanic respired carbon pool is a widespread phenomenon in the Pacific [*Jaccard et al.*, 2009].

4. Conclusions

[36] New paleoceanographic records from the SE Pacific show that MIS 12, 10, 8 and 6 were associated with significant reduction in ocean ventilation at ridge crest depth (\sim 3000 m), followed by a recharge of the bottom water with more oxygenated water after glacial terminations V, IV, III and II. This variation in ocean ventilation leads to changes in electron donor (hydrothermal sulfide) supply to the sediment that is moderated by enhanced carbon export during glacial terminations.

[37] The large hydrothermal plume at 15° S on the EPR is inferred to have delivered variable and decreasing amounts of particulate sulfide to the core site as it moved off axis over the last 740 ka. The main control on sedimentary accumulation of hydrothermal sulfide particles is distance from the hydrothermal source and bottom water oxygen content which in turn also controls the depth of the transition metal oxidation fronts in the sediment and the exposure time to oxidation at the seafloor. As a result only trace sulfide is now found in the sediment, and the transitory presence of sulfides in the sediments has been inferred from the very high contents of the chalcophile elements Cu, Cd, and U. In situ oxidation of sulfides and authigenic U uptake leads to further overprinting the inferred primary hydrothermal signature.

[38] After glacial terminations V, IV, III and II the ridge crest depth Pacific is inferred to be ventilated with higher oxygen content seawater and the active oxidation front progressively burns down during interglacial periods into the underlying glacial deposits modifying the Mn, Ba, Cu, V and U distributions. Suboxic ridge crest metalliferous sediments thus provide a Pleistocene history of past bottom water oxygen in a region that has sparse paleoceanographic records.

[39] Published sedimentary records from the southern EPR from 10°S through to 20°S exhibit subsurface U (and in some cases documented P) peaks that are largely undated (Figure 1). We demonstrate that these subsurface peaks may also be relicts of the basin-wide changes in glacial SE Pacific ventilation, but this hypothesis remains to be tested through determination of good sediment chronologies for each site and multielement analysis at an appropriate spatial resolution.

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