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ACS Earth Space Chem., **Just Accepted Manuscript** • DOI: 10.1021/acsearthspacechem.8b00203 • Publication Date (Web): 19 Feb 2019

Downloaded from <http://pubs.acs.org> on February 22, 2019

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Redox processes impacting the flux of Fe(II) from shelf sediments to the OMZ along the Peruvian shelf

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For submission to ACS Earth & Space Chemistry special issue

Abstract:

Iron (Fe) is a limiting nutrient in many regions of the open ocean and can also play a key role in controlling primary productivity in Eastern Boundary Upwelling Systems (EBUS). In EBUS regions, where intense Oxygen Minimum Zones (OMZs) contact the continental shelf, significant iron inputs can result from the supply of Fe(II) from reducing sediments. How much of this iron makes it to the photic zone depends on physical processes mixing over different time scales (minutes to decades) and the kinetics of redox and complexation processes impacting the biogeochemical cycling of iron. In this work we examine the controls on Fe(II) release from shelf sediments across the Peruvian OMZ by measuring Fe(II) and H_2O_2 in the water column and benthic boundary layer (BBL) and applying a simple 1D mixing model, with either 1 or 2 layers, where the flux of Fe(II) to the water column is treated as analogous to radon, that the decay rate is constant within the mixing layer. Our modelling approach then allows us to compare our estimated decay rate against published oxidation rates for specific oxidants of Fe(II) in OMZ waters and check the validity of our approach. Our data indicate that throughout the OMZ, Fe(II) decay rates may be partially influenced by H_2O_2 , but that it is most likely that nitrate-dependent anaerobic Fe(II) oxidizing (NDFO) bacteria are the main oxidizers. In the secondary nitrite maxima (SNM), abiotic NO_2^- or biotic mediated processes may also be important. This work highlights the importance and uses of redox species in understanding biogeochemical cycles in the ocean.

Keywords: H_2O_2 , ROS, NDFO, Nitrite, Nitrate, BBL.

1. Introduction

Understanding and interpreting the distribution of elements in the ocean, and perhaps more importantly the biogeochemical and redox processes that drive the observed distribution, is a major goal of chemical oceanographers today. An important step towards this goal is the identification and quantification of processes occurring in the ocean over differing spatial and temporal scales as data on this is vital for correctly interpreting the impact of chemical and biological processes occurring. Redox processes involving iron are an important part of the biogeochemical cycling of iron ¹ and other elements in the ocean ² thus improving our ability to model iron chemistry in the oceans is critical for Earth system models ³. However deciphering iron redox cycling in the ocean are more complicated than in sediment systems as the observed signals are the results of abiotic and biotic processes superimposed on a mixing ocean varying both spatially and temporally. Over the last 30 years the use of chemical tracers for studying mixing processes has undergone considerable evolution since the first ground breaking studies using CFCs were published ^{4, 5}, measurements now include multiple tracers (e.g. CFCs, ³He, ¹²⁹I) with different lifetimes and sources, leading to the development of concepts such as the tracer age of a water mass ⁶. Thus, in the context of international research programs such as GEOTRACES (www.geotraces.org), SOLAS (solas-int.org) and IMBER (www.imber.info) information on redox and mixing processes is critical to improving our understanding of biogeochemical cycles in the ocean.

One group of chemical tracers that has so far been underutilized are the transient redox species that exist in the ocean. These are chemical species which while being thermodynamically unfavoured under seawater conditions may exist over a range of

different temporal scales due to kinetic limitations. In many cases the reduction step might be relatively quick but reoxidation (e.g. H_2O_2 , Fe(II) and I^-) or further reduction (e.g. H_2O_2) may take some time to occur ^{7, 8}. Recent advances in analytical chemistry have seen the development of new methods specific for the determination of redox chemical species at open ocean concentrations paving the way for their application to use as tracers ⁹. A further key requirement for a useful transient redox tracer is information on the processes that control the conversion rate (oxidation or reduction) of the specific tracer. A major controlling factor for the speciation of many of these redox species is the concentration of dissolved oxygen (O_2) and thus a natural laboratory for examining these tracers is in oxygen minimum zones (OMZ) ¹⁰.

Analysis of time series data indicate that OMZs are expanding ¹¹ and this may result in an expansion of the distribution of redox sensitive species with resulting changes in the biogeochemical cycles of these elements ^{12, 13}. Recent work in OMZs and anaerobic zones suggest strong coupling between the Fe and N biogeochemical cycles; via chemodenitrification, where nitrite oxidises Fe(II) and is converted to N_2O , and by the microbially mediated nitrate-dependent anaerobic Fe(II) oxidation (NDFO) ^{14, 15}.

Anaerobic ammonium oxidation coupled to Fe(III) reduction (Feammox) ¹⁶ has been found in soils but not yet the ocean. Nitrate oxidation of Fe(II) has also been considered as possibly supporting life on Mars ¹⁷. The balance between microbial and abiotic oxidation rates for Fe(II) has implications for both our present day understanding of Fe cycling and that of the past ¹⁸. Thus, there are several compelling reasons to investigate the chemistry of transient redox tracers such as Fe(II) in OMZ waters. In the present work we report data on the vertical distribution of Fe(II) effluxed from reducing sediments into

the bottom waters in an OMZ region and examine potential processes controlling its distribution.

1.1 Mixing at Ocean boundaries.

Information on mixing processes in the surface ocean is critical to understanding the distribution and biogeochemical cycling of bio-important elements such as Fe. Vertical mixing in the ocean occurs over a range of spatial and temporal scales ¹⁹ and is an important constraint on the supply of nutrients and the exposure to light for phytoplankton ²⁰. Temporal variability in the processes affecting the surface ocean has seen the development of two terms to describe the surface mixed layer ²¹: (i) The *Active Mixing Layer* (AML) is defined as the depth zone which is actively mixed from the surface at a given time and generally corresponds to a zone where strong surface forcing induces turbulent mixing. (ii) The *Mixed Layer Depth* (MLD) is the maximum depth reached by the AML on time scales of 24 hours or more.

Early work on the mixing in the upper ocean were based on temperature or density profiles obtained using a conventional CTD and analysed in terms of Thorpe displacements ²². The recent development of microstructure profilers ²³ has led to improvements in the measurements of the turbulence in the upper ocean. Additional approaches have used tracer release experiments using SF₆ or high resolution velocity measurements made by a shipboard mounted acoustic doppler current profiler (ADCP) data to determine velocity shear this results in a number of approaches by which diffusion and mixing processes can be assessed in the upper ocean ²⁴⁻²⁹.

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3 The other boundary to the water column is the interface with the sediment. Here
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5 similar terms exist for the mixing zones adjacent to the ocean bottom. The benthic or
6
7 bottom boundary layer (BBL) is the region adjacent to the ocean bottom ³⁰, and in this
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9 region chemical and physical properties can be distinct from the overlying ocean interior
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11 ³¹. Turbulent boundary layer flow ³² in the BBL induces the formation of a bottom mixed
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13 layer (BML), which is typically 15-60 m in height ³³. The BML is in many ways
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15 analogous to the surface AML, and is the zone through which benthic-pelagic coupling of
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17 biogeochemical cycles can occur, as the surface AML is for air/sea interactions.
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21 Waters close to the sediment in the ocean have historically been assessed using
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23 light scattering (nephelometry) and are typically described by the following terms ³⁴:
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25 bottom nepheloid layer (BNL), which can extend up to 2000 m and the bottom mixed
26
27 nepheloid layer (BMNL) and which is equivalent to the BML. Above the BMNL there is
28
29 an approximately logarithmic fall-off in the intensity of light scattering up to the clearest
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31 water minimum that indicates the top of the BNL. Along continental margins,
32
33 intermediate nepheloid layers (INL) can be found and they result from interactions on the
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35 upper continental slope and at the depth of the shelf edge and may extend out across the
36
37 continental margin ³⁵. Determining mixing rates in the BBL is perhaps more complicated
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39 than in the AML as microstructure sensors are limited in their application here, instead
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41 recent work on mixing in the BBL have looked at different approaches to estimating
42
43 turbulent diffusion in this region ³¹ and employed specialized sampling devices to obtain
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45 samples in the BML ³⁶.
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51 The effect of turbulent mixing on the distribution of a chemical species can be
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53 simply described as quasi diffusive transport according to Fick's law of diffusion:
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55
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57

$$J = -K_z \frac{\partial C}{\partial z} \quad (1)$$

Where J is the flux through the water column, K_z is the diffusion coefficient, C the concentration of the chemical species and z the depth. Modelling of the physical processes in the AML has evolved from the diel cycle studies of the Price-Weller-Pinkel (PWP) model³⁷ to the present day use of turbulent closure approaches³⁸ which form the basis of the 1D General Ocean Turbulence model (GOTM) model (www.gotm.net) which has been applied to iron redox processes in the surface ocean³⁹. Iron may be thought of as an unlikely tracer due to its particle reactivity and non-conservative behaviour in seawater, however total Fe has been used previously as a mixing tracer in an iron enrichment experiment⁴⁰ giving values similar to microstructure measurements²⁴. In the present work our aim was not to examine the mixing modelling approaches in detail but to focus on the chemical processes involved and evaluate the suitability of this approach to transient redox tracers for use in biogeochemical modelling studies.

1.2 Iron speciation and supply to the photic zone from sediments

In oxygenated seawater Fe(III) is the dominant redox species, as while Fe(II) is more soluble than Fe(III) at alkaline pH, Fe(II) is rapidly oxidized by reactive oxygen species (ROS), principally O_2 and H_2O_2 ^{41,42}, with the short lived radical species O_2^- and OH are also involved. However in seawater Fe(III) is poorly soluble and is rapidly hydrolysed resulting in the formation of various Fe(III) oxyhydroxide phases with differing chemical reactivities⁴³. Dissolved Fe(III) is strongly complexed by organic chelators in seawater^{44,45}, most likely produced by bacteria or phytoplankton, which overall increase the solubility of iron⁴⁶. The reduction of Fe(III) to Fe(II) by

photochemical or other processes is a possible mechanism by which colloidal iron is made more bioavailable to phytoplankton ^{47, 48}. In tropical waters, Fe(II) is viewed as a short lived intermediate in iron cycling ⁴⁹, existing at low concentrations (pM or less), however during spring blooms in colder coastal waters Fe(II) has been detected at elevated concentrations (~ 1 nM) ⁵⁰. More recently significant concentrations of Fe(II) have been detected at depth in the sub-oxic waters of oxygen minimum zones ⁵¹⁻⁵³.

Continental shelf sediments can be a significant source of iron, through simple resuspension and mixing, to the overlying water column ^{54, 55} and a primary source of iron for phytoplankton ⁵⁶. In EBUS regions, where an OMZ interacts with the continental shelf, iron fluxes can be considerably enhanced ⁵⁷⁻⁵⁹ and lead to what Bruland et al. termed the Blue and Brown waters of Peru ⁵⁹, where inshore high iron concentrations helped to supply large phytoplankton blooms fuelled by upwelling waters, while offshore waters were low in iron and optically very clear with little phytoplankton activity. In this context the cross shelf transport of iron is also important, and recent work has shown that in the Peru OMZ there is considerable mesoscale eddy activity ^{60, 61} and that these eddies reduce biological activity inshore ⁶² while increasing offshore activity and impacting N cycling there ^{63, 64}. Along the Peruvian continental shelf, significant concentrations of Fe(II) have been detected previously in the water column ^{53, 65, 66} and were associated with nitrite maxima and/or benthic sources. Similarly a well-defined INL was found previously along the Peruvian coast with a particle maximum at 200 m ⁶⁷ and was associated with the nitrite maxima within the core of the OMZ.

Sediment porewaters are typically lower in pH and oxygen than the overlying seawater resulting in an environment that favours Fe(II) this leads to the iron flux from

the sediments being predominantly Fe(II) ^{68, 69} and fluxes increasing with decreasing oxygen concentrations ⁷⁰. There is some data suggesting sediments are also a source of iron binding ligands ^{71, 72} to the water column, oxidation of the Fe(II) and subsequent complexation of Fe(III) has been proposed as a mechanism by which the flux of iron from the sediment could be mixed further through the water column ⁷³. How far the iron is transported than is a balance between the kinetics of oxidation, complexation and scavenging on to particles that are then sedimented out, processes that occur in the AML ⁷⁴ as well as the BBL.

1.3 Oxidation kinetics of Fe(II) in seawater

There have been a number of laboratory studies undertaking precise measurements of Fe(II) oxidation rates in seawater at nM levels ^{41, 42, 75-79}. However the measurement of Fe(II) in the open ocean is complicated by low concentrations (< 100 pM) and the potential for artefacts induced by the analytical method employed ⁸⁰. The recent development of flow injection chemiluminescence techniques ⁸¹ now permits studies into the speciation and cycling of Fe(II) in natural waters with minimal sample perturbation. There are only a few measurements of Fe(II) oxidation rates at low O₂ concentrations ⁸²⁻⁸⁴ and deviations from the rate laws found at higher O₂ concentrations are thought to be due to H₂O₂ being the dominant oxidant and back reactions between Fe(III) and O₂⁻. H₂O₂ has previously been shown to be a key oxidant for Fe(II) in Antarctic waters ⁸. Table 1 summarizes data on Fe(II) oxidation rates in the literature that is of relevance to the present study.

In the absence of ROS species, attention more recently has focused on Fe(II) oxidation by NO_x species in abiotic and biotically mediated pathways⁸⁵. The abiotic oxidation of Fe(II) by NO_3^- in the absence of O_2 has been found in earlier studies to have a maximum rate around pH 8^{86, 87} and while it is slower than oxidation by NO_2^- at neutral pH, it is efficiently catalysed by Cu⁸⁸. Contrastingly Cu was not found to catalyse the reaction between NO_2^- and Fe(II) at pH 8 but did alter the ratio of the products, N_2O and N_2 ⁸⁹. Heterogeneous reactions involving Fe(II) containing minerals (e.g. nontronite or green rust) have been shown to be very important as they can catalyse the reaction with NO_2^- ⁹⁰. Recent works have focused on identifying and quantify the processes involved during Fe(II) oxidation by NDFO bacteria in order to assign biotic and abiotic rates as at the high concentrations of Fe(II) typically employed the heterogeneous reactions may mask any microbial oxidation^{91, 92}.

1.4 Application of Fe(II) as a tracer in the BBL of OMZ waters

In regions where the OMZ overlies the continental shelf or slope, Fe(II) diffusing from the sediments may persist for some time due to the low O_2 concentrations present there. Under the conditions of constant O_2 concentrations in the OMZ and in the absence of significant advective transport the vertical distribution of Fe(II) may be considered analogous to the release of radon from the sediments if it is assumed that there is a constant decay rate. Radon has been used in a number of studies to determine mixing rates in the BBL⁹³⁻⁹⁶. A brief outline of the approach is given below:

The OMZ waters in the BBL above the sediment are assumed to have constant temperature, pH, salinity and that the decay rate for Fe(II), λ , is constant throughout the BBL and can be considered analogous to the half-life of a radiotracer such as ^{222}Rn .

$$\lambda = k_{O_2}[O_2] + k_{H_2O_2}[H_2O_2] + \dots \quad (2)$$

Where k_{O_2} and $k_{H_2O_2}$ are the calculated rates constants for the oxidation of Fe(II) by O_2 ⁴² and H_2O_2 ⁴¹ respectively, other terms for NO_2^- or NO_3^- mediated oxidation could also be include (see Table 1 for details). Under the conditions of constant vertical diffusivity (K_z) in the BBL the vertical distribution of Fe(II) (denoted below as C) can be formulated as follows^{94, 97}:

$$K_z \frac{\partial^2 C}{\partial z^2} - \lambda C = 0 \quad (3)$$

1.4.1 Single layer model

This simple differential equation can be solved using a simple set of boundary conditions;

(i) Sediment/Water interface: $C=C_0$ at $z=0$, (ii) Top of the water column: $C \rightarrow 0$ as $z \rightarrow \infty$.

Gives the following solution

$$C = C_0 e^{-z \sqrt{\frac{\lambda}{K_z}}} \quad (4)$$

The flux (J) of Fe(II) at the sediment water interface can be related to C_0 as follows

$$J = -K_z \frac{\partial C}{\partial z} \bigg|_{z=0} \quad (5)$$

which then reduces to

$$J = C_0 \sqrt{K_z \lambda} \quad (6)$$

The steady state inventory of Fe(II) in the water column, (I_{ss}), from the benthic source can also be described:

$$I_{ss} = \int_{z=0}^{z=\infty} C \partial z \quad (7)$$

which leads to the following relationship

$$I_{ss} = \frac{J}{\lambda} = C_0 \sqrt{\frac{K_Z}{\lambda}} \quad (8)$$

Thus the steady state inventory of Fe(II) is related to the flux from the sediment and the half-life for Fe(II) in the overlying waters. Knowledge of one of these variables (e.g. the flux from sediment porewater profiles or in situ measurements of the decay rate) will allow the calculation of the other. Similarly, rearranging equation 8 allows a solution for K_Z

$$K_Z = \frac{J^2}{\lambda C_0^2} \quad (9)$$

Alternatively, a value of K_Z can be estimated by relationships to the inverse of the buoyancy frequency (N)^{27, 96} or via other tracers.

1.4.2 Two layer model

The two layer model originates from the Air/Sea interface radon model of Peng *et al.*⁹⁸ which was applied to benthic fluxes of radon by Sarmiento *et al.*⁹⁶. In this case we designate the layer closest to the sediment as layer 2, with thickness D , and that in the interior as layer 1. As above λ denotes the decay rate of Fe(II), with K_i and λ_i the vertical diffusivity and Fe(II) decay rate in each layer respectively.

Thus, the equation for the upper layer (layer 1) is

$$C_1 = C_D e^{\left(\frac{-(z-D)}{\sqrt{\frac{K_1}{\lambda_1}}} \right)} \quad (10)$$

Where C_D denotes the concentration at the interface between layer 1 and 2. The equation for the lower layer is

$$C_2 = \frac{C_D}{2} \left(1 + \frac{\sqrt{\frac{K_1}{\lambda_1}}}{\sqrt{\frac{K_2}{\lambda_2}}} \right) e \left(\frac{-(z-D)}{\sqrt{\frac{K_2}{\lambda_2}}} \right) + \frac{C_D}{2} \left(1 - \frac{\sqrt{\frac{K_1}{\lambda_1}}}{\sqrt{\frac{K_2}{\lambda_2}}} \right) e \left(\frac{(z-D)}{\sqrt{\frac{K_2}{\lambda_2}}} \right) \quad (11)$$

We simplify here by using the term $a_i = \sqrt{(K_i/\lambda_i)}$ for each layer.

$$C_2 = \frac{C_D}{2} \left(1 + \frac{a_1}{a_2} \right) e \left(\frac{-(z-D)}{a_2} \right) + \frac{C_D}{2} \left(1 - \frac{a_1}{a_2} \right) e \left(\frac{(z-D)}{a_2} \right) \quad (12)$$

The concentration at the sediment water interface is then

$$C_0 = \frac{C_D}{2} \left(1 + \frac{a_1}{a_2} \right) e \left(\frac{D}{a_2} \right) + \frac{C_D}{2} \left(1 - \frac{a_1}{a_2} \right) e \left(\frac{-D}{a_2} \right) \quad (13)$$

The flux throughout the water column is then:

$$J = K_2 \frac{\partial C}{\partial z} \Big|_{z=0} \quad (14)$$

$$J = -K_2 \frac{C_D}{2a_2} \left(\frac{a_1}{a_2} + 1 \right) e \left(\frac{(D-z)}{a_2} \right) - K_2 \frac{C_D}{2a_2} \left(\frac{a_1}{a_2} - 1 \right) e \left(\frac{-(D-z)}{a_2} \right) \quad (15)$$

Thus at the boundary where $z = 0$:

$$J = -K_2 \frac{C_D}{2a_2} \left(\left(\frac{a_1}{a_2} + 1 \right) e \left(\frac{D}{a_2} \right) + \left(\frac{a_1}{a_2} - 1 \right) e \left(\frac{-D}{a_2} \right) \right) \quad (16)$$

See the S.I. for this manuscript for more details on the derivation of the steady state concentration. A key observation here is that it is not possible as in the 1D case to separate the oxidation rate from the diffusive mixing rate using the steady state concentration as the number of unknowns has increased to 5 (or 6 if you solve also for D). However as for the 1D case this could be reduced to 3 unknowns if estimates of K_i are made using other approaches, though in practice with a well-mixed BBL, fitting of K_2 only provides a lower bound for the mixing rate.

2. Experimental Section

2.1 Study sites

2.1.1 *Benthic-Pelagic coupling on the Peruvian shelf*

The study area considered in this work was the OMZ off the coast of Peru. Results are presented for the RV Meteor M77-1, Talcahuano, Chile to Callao, Peru (Oct 23 – Nov 21, 2008) an expedition which focused on benthic-pelagic coupling on the Peruvian shelf between 10° and 12° S^{99, 100}.

2.2 *Analytical Measurements*

2.2.1 *Dual measurement of Fe(II) and H₂O₂*

Seawater samples were obtained either using Niskin bottles on a standard Seabird CTD and rosette. A key element of this study was sampling close to the sediment using the CTD and this was accomplished successfully to within 1-2 m above the bottom by virtue of a well calibrated altimeter and stable weather conditions with minimal swell. Niskin bottles were checked for gas leaks prior and after each deployment. In this study samples were drawn first (before any other samples) from the Niskin bottles into brown high density polyethylene (HDPE) bottles (Nalgene) using a silicone tube in the same manner as for low level dissolved O₂¹⁰¹ with no headspace or bubbles in the samples. HDPE bottles are significantly less oxygen permeable than LDPE or Teflon. Use of clear glass bottles is avoided due to the potential for photoreduction of Fe and/or photo-formation of H₂O₂. Samples were analyzed in triplicate, within 1-2 hours of collection and were unfiltered. All sampling and analysis was performed in a trace metal clean laminar flow cabinet (AirClean Systems).

Fe(II) was determined using a modified version of an existing chemiluminescence method⁸¹. In brief the system was adapted to run Fe(II) and H₂O₂ simultaneously on the

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3 same machine by continuously running sample to the detector and using an injection
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5 valve to separate the two independent reagent flows which had been prepared as
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7 described previously. This differs from the usual setup employed for Fe(II) as previously
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9 it was run in sample injection mode and here it is reagent injection mode. Samples were
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11 maintained at close to the ambient seawater temperature to maintain oxidation of Fe(II) at
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13 *in situ* rates and N₂ gas was used to fill the headspace of the bottles as the sample was
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15 removed so as to prevent atmospheric O₂ from entering the sample. The detection limit
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17 for this technique during this work (all analysis) ranged from 4 to 20 pM and depended
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19 mostly on the background chemiluminescence from the luminol reagent. Peak width and
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21 shape was used as criteria for the determination of Fe(II) at <100 pM levels as the
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23 presence of a small injection peak can bias peak height based analysis to anomalously
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25 higher Fe(II) levels.
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31 Standards for Fe(II) were run in oxygen deficient water from the core of the
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33 OMZ, a key point often overlooked in other studies is that the Fe(II) analytical signal is
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35 dependent on O₂ in the sample and in the reagent stream ¹⁰². In the case of samples from a
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37 strongly oxygen deficient OMZ, the only O₂ supplied is in the reagent stream and thus
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39 the size of the sample loop and flow rate are critical as too big a loop, or too slow a flow
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41 rate, will lead to a double peak as the reaction is minimized when the ratio of sample to
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43 reagent is highest in the mixing zone of the flow cell. Another aspect of this is that values
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45 of Fe(II) in oxygenated waters are likely over estimated due to the increased O₂ available
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47 for reaction. Standard curves were fitted using a quadratic function due to the non-
48
49 linearity of the Fe(II) response over the wide range of Fe(II) concentrations encountered
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52 20 pM to 200 nM.
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Previously we have applied a further criteria for the detection of Fe(II) ²⁸ that discrete samples should be measurable for at least one half-life ($t_{1/2}$), in the present work the slow oxidation rates in the OMZ waters meant it was not practical to run the samples over a complete half-life for these samples. An earlier study on Fe(II) in seawater suggested that V(IV) may be an interference to the Fe(II) method ⁵¹. In the present work we suggest that there is little efflux of V(IV) from the sediments as while vanadate can be reduced to vanadyl by H₂S or organic compounds, the vanadyl ion readily adsorbs to particle surfaces ^{103, 104} and would not readily diffuse out, though it may be present in resuspended particles. Additionally recent work has indicated that in the oxygen-depleted bottom waters on the Peru margin the removal of dissolved V was not observed ¹⁰⁵ suggesting no V(IV) formation.

H₂O₂ was measured using a flow injection chemiluminescence (FIA-CL) reagent injection method ¹⁰⁶ as described previously ¹⁰⁷. H₂O₂ standards were made by serial dilution from a primary stock solution (30% Fluka - Trace Select). The concentration of the primary standard was determined by direct spectrophotometry of the solution ($\epsilon = 40.9 \text{ mol L}^{-1} \text{ cm}^{-1}$, ¹⁰⁸). Secondary standards were analysed with a spectrophotometric method using Cu(II) and 2,9-dimethyl-1,10-phenanthroline ¹⁰⁹. Seawater samples were measured directly by FIA-CL. Sample concentrations were corrected daily for the reagent blank ¹⁰⁶, which was typically less than 0.2 nM. We note that another recent work on H₂O₂ in seawater ¹¹⁰ using this method included catalase as part of their blank measurement, however as noted in the original method ¹⁰⁶ this is prone to a range of artefacts due to catalase's low efficiency at low H₂O₂ concentrations and its interference

with the luminol assay ¹¹¹. Samples precision was 2-3% through the concentration range 1-100 nM, the detection limit (3σ) was typically 0.6 nmol L⁻¹.

As mentioned above O₂ concentrations in the sample influence the sensitivity of the Fe(II) measurements ¹⁰², but they don't appear to interfere with H₂O₂ analysis ¹⁰⁶. In the original method for the H₂O₂ analysis it was recommended to allow samples for H₂O₂ with Fe(II) in them to set for an hour before analysis to allow the Fe(II) to oxidize. However in OMZ waters we chose to sample both the Fe(II) and H₂O₂ simultaneously, as if there is O₂ and Fe(II) in the sample you will generate H₂O₂ as the Fe(II) oxidises, alternatively if the sample is anoxic then the oxidation of the Fe(II) will be very slow (e.g. hours to days) if the bottle is sufficiently gas tight, thus there is the potential to oxidize the initial H₂O₂ present in the sample. Our initial shipboard work indicated that the timescale of oxidation was significantly longer than a few hours and that the potential for O₂ permeation into the bottles was increased, leading to increased H₂O₂ concentrations post sampling. We have found that high concentrations of H₂O₂ (20 nM or more) can cause a small apparent Fe(II) signal (pM level) but this may be related to reactions at the analysis pH between HO₂⁻ and Fe(III) in the sample to generate O₂⁻ and thus unavoidable as the signal will be dependent on the concentration of dissolved and colloidal iron present in the sample. Lastly other potential interferences for both methods that are possible, but difficult to avoid with unfiltered samples, are O₂⁻ production by O₂ sensitive bacteria from the OMZ and/or bioluminescence.

3. Results and Discussions

3.1 Hydrography

The location of the stations sampled during the M77-1 expedition are shown in Figure 1 along with the locations of sampling sites from other studies on Fe(II) in the Peruvian OMZ^{53, 65}. The main hydrographic feature along this part of the Peruvian EBUS is the strong upwelling that occurs year round, which supports some of the highest primary productivity on the globe¹¹². The Peruvian OMZ is created and maintained by a high demand for oxygen in the waters underlying the upwelling zone and weak supply of oxygenated waters¹¹³. The upwelled waters of the OMZ are sourced from nutrient rich equatorial subsurface water (ESSW) transported by the Peru Chile Undercurrent (PCUC), a poleward flowing current¹¹⁴ found at approximately 150 m depth 10° and 15° S on the Peruvian shelf. Highest chlorophyll concentrations are found in the spring and summer, out of phase with the winter upwelling maximum¹¹⁵. Our expedition took place in the austral spring and Figure 2 illustrates the increase in ocean productivity over the shelf immediately preceding and during our work at sea. Long term trends (1996-2011) in the OMZ and nutrients at a monthly time series station off the coast of Callao (Figure 1) reveal a system responding to variability on seasonal to interannual timescales.

3.2 Distribution of dissolved Oxygen (O₂) and Hydrogen Peroxide (H₂O₂)

3.2.1 O₂ distribution

The distribution of dissolved oxygen at 3 selected stations are shown in Figures 3-5 and in the SI (Figures S1-S4). Briefly the surface waters and AML were well oxygenated but O₂ rapidly decreased below that to concentrations below 20 μM. It should

be noted the Seabird SBE43 polarographic membrane oxygen sensor employed on the CTD in this study, can't resolve O₂ concentrations below about 1-2 μM¹¹⁶. Recent studies using the STOX electrode¹¹⁶ have indicated that there is less than 2 nM O₂ in the core of the Peruvian OMZ¹¹⁷. O₂ concentrations increased below 400 m at the deeper stations occupied during M77-1 (Figure S6).

3.2.2 H₂O₂ in the OMZ

H₂O₂ concentrations for 3 stations are shown in Figures 3-5 and the data can also be found in the SI. H₂O₂ concentrations were elevated as expected in the euphotic zone, presumably due to photochemical processes¹¹⁸ and decreased rapidly to very low levels, but still detectable, in the core of the OMZ. H₂O₂ concentrations increased again at most stations near the sediment-water interface. There have only been a few other studies on H₂O₂ in the Peruvian OMZ, and our results are broadly consistent with them for the euphotic zone, though the earliest study by Zika *et al.*¹¹⁹ could not detect H₂O₂ below 5 nM due to the method employed at the time. The most recent study¹¹⁰ only included two stations from the Peru OMZ and reported much higher H₂O₂ concentrations (10 nM) than we observed (0.3-2 nM) in the core of the OMZ and they suggested the high values they observed was due to the presence of nM Fe(II). The methodologies used in our study and theirs are very similar, though they report taking their samples after those drawn for O₂ and that they collected their samples directly from the spigot on the Niskin (to avoid the possible effects of light), this differs from our work where we sampled first and via a sampling tube as for low O₂ measurements to eliminate the introduction of bubbles to the sample.

O₂ is an obligatory precursor for H₂O₂ for all known routes of formation and thus it seems somewhat puzzling at first that H₂O₂ could exist where there is effectively no O₂. However, H₂O₂ could be formed at oxic/anoxic boundaries and migrate into anoxic environments, one such route might be from sinking particles where O₂ is consumed leading to anoxic microenvironments and a flux of H₂O₂ from the sinking particle ^{120, 121}. Further to this a respiratory enzyme in the facultative anaerobe *Escherichia coli*, cytochrome c peroxidase, was recently found to use H₂O₂ as a terminal electron acceptor ¹²² and was only expressed when H₂O₂ is present and molecular oxygen is absent suggesting an unappreciated niche for microbes utilizing this enzyme and substrate.

3.3 Distribution and fluxes of Fe(II) in the OMZ water column

3.3.1 Distribution of Fe(II)

Significant concentrations of Fe(II) were observed in the water column of the Peruvian shelf during both the M77-1. Examples profile are shown from stations 532 (Figure 3), 599 (Figure 4) and 618 (Figure 5). At this site dissolved oxygen was below 2 μM from 20 m, just below the mixed layer, all the way to the bottom. There was no evidence for any sulfide formation at this time, though a later expedition in the same region would detect sulfide and enhanced benthic Fe fluxes ¹²³. Fe(II) was detectable at all stations throughout the water column with highest concentrations in the BBL which decreased with proximity from the sediments consistent with a benthic source of Fe(II). In OMZ waters, the combination of low O₂ and H₂O₂, a low pH (7.6 -7.8) results in a dramatic extension of the half-life for Fe(II) in seawater compared to that in warm well oxygenated surface waters.

Our findings are consistent with earlier work by Hong and Kester on the Peruvian shelf (Figure 1) which also found high concentrations of Fe(II) on the inner shelf close to the sediments⁶⁵, however in that study Fe(II) was measured spectrophotometrically using Ferrozine resulting in a much higher detection limit and many of the features seen in our new data set were not observable. Hong and Kester found Fe(II) and nitrite together suggesting a common source from the sediments and this was also observed by us at station 532 (Figure 3).

High concentrations of iron in Peruvian shelf waters were also observed by Bruland and coworkers⁵⁹ though they did not measure Fe(II) specifically. More recent work by Vedamati et al.⁵³ off the shelf but in OMZ waters found Fe(II) maxima coincident with the secondary nitrite maxima (SNM), as has been observed in the Eastern Tropical North Pacific⁵¹ and the Indian Ocean⁵². This feature is also present in our data at station 532 (unfortunately for the other stations we don't have nitrite data). Chever et al.⁶⁶ also found significant concentration of Fe(II) to the north of our study site.

3.3.2 Modelling of Fe(II) profiles and estimation of Fluxes

For many of the stations sampled we could apply a simple 1 layer 1D water column model (see the description above) (Figure 3 and 4). While at other stations the Fe(II) distribution was clearly marked by the presence of a well mixed BBL (Figure 5) which required the use of a 2 layer model. Fitting of the profiles was performed in MatlabTM using the lsqcurvefit function with the error of the fit estimated using the Jacobian matrix.

During M77-1, a suite of cores were taken along the same transect as the CTD sampling for which Fe(II) fluxes were estimated from pore water gradients¹⁰⁰. In the present work we use those flux estimates to derive oxidation rates using our model. Alternatively we could have calculated the K_Z from relationships to the buoyancy gradient for BBLs⁹⁶ and then fitted the equations for C_0 and λ only to estimate the flux J (equation 6).

The results from our model simulations are summarized in Table 2. Oxidation rate estimates ranged from $6.7 \times 10^{-8} \text{ s}^{-1}$ to $\sim 9.9 \times 10^{-7} \text{ s}^{-1}$, this is equivalent to a range of half-life values from ~ 200 hours to ~ 2900 hours (8 to 120 days). Our estimates for K_Z range from 2.3 to $28.4 \text{ cm}^2 \text{ s}^{-1}$ for the BBL. These values are reasonably consistent with other measurements from other regions for the BML and BBL though data is sparse³¹.

Cross shelf transport in the Peruvian margin via meso-scale eddies has been identified as a key process in this region⁶⁰. Thus eddies that interacted with the shelf could potentially transport some Fe(II) out across the shelf before it was oxidized and this may represent an important mechanism in the development of INLs⁶⁷ in this region and the supply of Fe to the offshore nutrient rich waters. Alternatively recent 2D modelling work on EBUS regions has indicated that shelf slope and weak stratification can concentrate onshore transport in the BBL¹²⁴ which could help to supply iron to the upwelled waters. Quantifying the cross-shelf and along shore transport of Fe should therefore be a major goal of future work.

3.3.3 Oxidation of Fe(II) in OMZ waters

3.3.3.1 Oxidation kinetics under low O_2 conditions

Comparing our estimates (Table 2) of the *in situ* Fe(II) half life with those estimated from laboratory studies (Table 1) relies on assuming that these rates scalar according to the rate laws, which may not necessarily hold when going from mM to nM concentrations. There is also the question of non-stoichiometry reactions occurring at lower concentrations as other reactions become more important, as seen for Fe(II) oxidation at low O₂ levels, where H₂O₂ can become the major oxidant ⁸⁴. Our data does suggest however that O₂ is not the major oxidant for Fe(II) in the core of the OMZ, as if using the upper bound of 2 nM O₂ given by Revsbech et al. ¹¹⁶ this would results in an apparent rate of $\sim 5 \times 10^{-10} \text{ s}^{-1}$ much slower than what we estimate using our modelling approach.

At these very slow oxidation rates, there is sufficient time for the inorganic Fe(II) diffusing from the sediments to become complexed or adsorbed to particulate matter, thus changing its oxidation rate ²⁸. In many instances it may already be organically complexed when it leaves the sediments. It is however still labile as it reacts in the time frame of our analytical method, while for example the strong complex formed between Fe(II) and Ferrozine does not. Heterogeneous oxidation of Fe(II) on particles is more rapid at ambient O₂ concentrations though the few studies that have been made at low O₂ levels indicate that there is less of an enhancement on the oxidation rate by particles ^{125, 126}. Although there is now data available on the mineralogy and solid phase speciation of particles in the Peruvian OMZ ¹²⁷ we don't have a good indication of how much Fe(II) is adsorbed on them to be able to assess how this might impact the heterogeneous rates of oxidation.

Based on the analysis of the rate data in Table 1, H_2O_2 is likely then to be the key oxidant for Fe(II) but our water column data also strongly suggests that H_2O_2 is not present in excess of Fe(II) often and is thus not capable of completely oxidizing all the Fe(II) coming out of the sediments, so another oxidant more abundant is required.

3.3.3.2 Abiotic and biotic Nitrate/Nitrite reduction – Fe(II) oxidation

Nitrate and Nitrite in OMZ waters clearly fulfil the condition of being an oxidant in excess that could remove Fe(II) from the water column. If we start by examining the abiotic rates from Table 1 we can see that it is unlikely that NO_3^- is the oxidant, though NO_2^- may be important in the secondary nitrite maximum. It is also thought unlikely at this stage that Cu catalyses the NO_2^- reaction⁸⁸ under seawater conditions as Cu is strongly organically complexed in these waters¹²⁸.

So are the biotic reactions as has recent studies have suggested that NDFO is the most likely mechanism for the oxidation of Fe(II) in the Peru OMZ?^{127, 129}. Microbial genomics studies indicate that a range of taxa can oxidize Fe(II)¹³⁰, most notably the *Alphaproteobacteria* (phototrophic iron-oxidizing), *Betaproteobacteria* (neutrophilic aerobic and nitrate-dependent), *Gammaproteobacteria* (mostly acidophilic) and *Zetaproteobacteria*. For the anoxic bottom waters on the Peruvian shelf, the current information would suggest that Fe(II) oxidation would be likely carried out by *Betaproteobacteria* and/or *Zetaproteobacteria*. Cellular oxidation rates for Fe(II) by *Zetaproteobacteria* are available from a recent study by¹³¹ and this indicates that oxidizing 1-10 nM of Fe(II) over a time frame of 200 to 2900 hours (see above) would require only ~50 to 7500 cells L^{-1} which would only be a small fraction of the

metabolically active cells present in the water column. As there is no information on the temperature dependence of these organisms it is likely at colder temperatures the rates may be slower than reported here. However it does appear that NDFO bacteria are the most likely controls on Fe(II) in the Peru OMZ, if this is the case and the bacteria were uniformly distributed in the BBL and/or core of the OMZ this would then be some support for our assumption in equation 2 that the rate of Fe(II) oxidation was constant (e.g. $\lambda = k_{NDFO}[cell]$), this however awaits quantification of Fe(II) oxidizing bacteria and measurement of these rates under ambient conditions.

One key aspect of Fe cycling that still remains unclear at present however is the kinetics of formation and oxidation of Fe(II) at the SNM (Figure 3) as the presence of NDFO bacteria and/or abiotic NO_2^- oxidation helps explain the Fe(II) loss terms, but for this feature to form it requires a faster reductive process to be occurring so that significant concentrations of Fe(II) can be built up. This is an interesting puzzle that requires further investigation.

3.4 Residence time of Fe(II) in the Peru OMZ

Previously Scholz *et al.*¹²⁹ estimated the residence time of Fe in the Peruvian OMZ to be 16-18 h based solely on Fe(II) fluxes and the total water column inventory for dissolved iron, our data for the same region indicate a longer residence time for Fe(II) based of 200 to 2900 hours. In the case of Scholz et al. they made the assumption the dissolved iron loss term was all due to NDFO. A further difference between our estimate and theirs is also in the inventories, as we only considered Fe(II) and their estimate is made using the total dissolved iron pool (Fe(II) and Fe(III)). Our model only examines

Fe(II) oxidation losses, while their loss term would be scavenging of dissolved Fe(III) by particulate phases which is likely to be the main driver of the short residence time for dissolved iron in these waters. This is analogous to aerosol deposition in the upper ocean under the Saharan dust plume, where dissolved iron has a significantly longer residence time than total iron ¹³².

4. Conclusions and Future work

We have shown that combined measurements of Fe(II) and H₂O₂ provide valuable information on redox processes and mixing in the BBL of the Peruvian OMZ and can be combined with benthic flux estimates to improve modelling of iron fluxes to EBUS regions. Our data also indicate that H₂O₂ may be an important transient species in OMZ, but it is unlikely to be the controlling oxidant for Fe(II) as it appears at present that NDFO bacteria are capable of this. Overall it is expected that this new approach to iron redox processes will help to significantly improve our understanding of redox cycling in the ocean and on mixing processes at ocean boundaries.

Associated Content:

The supporting information is available free of charge on the ACS Publications website at DOI: 10.1021/acsearthspac-chem.xxxxxxx

Location of stations sampled during M77-1 for Fe(II) (Table S1); Fe(II) and H₂O₂ at S532 (Table S2); Fe(II) and H₂O₂ at S547 (Table S3); Fe(II) and H₂O₂ at S556 (Table S4); Vertical profiles for hydrographic and redox parameters at S556 (Figure S1); Fe(II) and H₂O₂ at S569 (Table S5); Vertical profiles for hydrographic and redox parameters at

S569 (Figure S2); Fe(II) and H₂O₂ at S575 (Table S6); Vertical profiles for hydrographic and redox parameters at S575 (Figure S3); Fe(II) and H₂O₂ at S596 (Table S7); Fe(II) and H₂O₂ at S599 (Table S8); Fe(II) and H₂O₂ at S609 (Table S9); Vertical profiles for hydrographic and redox parameters at S609 (Figure S4); Fe(II) and H₂O₂ at S618 (Table S10); O₂ concentrations (as measured by a SBE-43 sensor) along for the transect along 11° S during M77-1 (Figure S5); Fe(II) oxidation half-life comparison (Figure S6) (PDF)

Acknowledgements:

We kindly thank the government of Peru for allowing us access to their territorial waters and to the officers and crew of RV Meteor for their help and cooperation during the research expedition M77-1 described in this work. Special thanks are also due to our colleagues at GEOMAR for their support of our shipboard and shore-based laboratory work. We gratefully acknowledge the authorities in Peru for their kind permission to work in their territorial waters. This work is a contribution of the Sonderforschungsbereich 754 “Climate – Biogeochemistry Interactions in the Tropical Ocean” (www.sfb754.de). Financial support for this work was provided by the Deutsche Forschungsgemeinschaft (DFG) via grants to PLC (CR145/15-1 and SFB754 B5). Aspects of this work were also supported by SOPRAN, a BMBF Verbundprojekt (FKZ 03F0462A and 03F0611A) which forms part of the German contribution to SOLAS. PLC acknowledges that this publication has emanated from research supported in part by a research grant from Science Foundation Ireland (SFI) under Grant Number 13/RC/2092 and co-funded under the European Regional Development Fund and by iCRAG industry partners. This work is also a contribution to SCOR working group 151: Iron Model Intercomparison Project (FeMIP).

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1 Table 1: abiotic and biotic Fe(II) oxidation rates (in seawater unless noted)

Oxidant	Stoichiometry	Rate Law	Temperature	pH	Rate (k_{app}) Fe(II)	Reference
O ₂	~4:1	$-k[\text{Fe(II)}]_d[\text{O}_2]$ $-k[\text{Fe(II)}]_d[\text{O}_2][\text{OH}^-]^2$	13° C, 34.9 S	7.6	$1.2 \times 10^{-8} \text{ s}^{-1}$ @ 30 nM O ₂ $\sim 1 \times 10^{-8} \text{ s}^{-1}$ @ 30 nM O ₂	42 83 84
H ₂ O ₂	~2:1	$-k[\text{Fe(II)}]_d[\text{H}_2\text{O}_2]$	13° C	7.6	$2.74 \times 10^{-5} \text{ s}^{-1}$ @ 1 nM H ₂ O ₂	41
O ₂ ⁻	1:1	$-k[\text{Fe(II)}]_d[\text{O}_2^-]$	25° C	7.6	$1 \times 10^7 \text{ M}^{-1} \text{ s}^{-1}$ (a)	133, 134
OH	1:1	$-k[\text{Fe(II)}]_d[\text{OH}]$	25° C	7.6	$3 \times 10^8 \text{ M}^{-1} \text{ s}^{-1}$ (b)	135
[NO ₃ ⁻]	~2:1	$-k[\text{Fe(II)}]_d[\text{NO}_3^-]$	20° C	8	$1.5 \times 10^{-9} \text{ s}^{-1}$ @ 30 μM NO ₃ ⁻	88
[NO ₂ ⁻]	~2:1	$-2k[\text{Fe(II)}]_d[\text{NO}_2^-]$	25° C	7	$6.67 \times 10^{-9} \text{ s}^{-1}$ @ 1 μM NO ₂ ⁻	136
[NO ₂ ⁻], HFO	~2:1	$-k[\text{Fe(II)}]_d[\text{Fe(II)}]_s[\text{NO}_2^-]$	26° C	6.8	$\sim 4 \times 10^{-15} \text{ M s}^{-1}$	137
ZPB CP8 ^(c)	1:1	$-k[\text{cell}]$	22° C	7.1	$1.84 \times 10^{-18} \text{ mol cell}^{-1} \text{ s}^{-1}$	131

(a) Bimolecular rate constant k , there are no estimates for steady state O₂⁻ in the core of an OMZ at present, presumably less than 1 pM.
(b) Bimolecular rate constant k , there are no estimates for steady state OH in the core of an OMZ at present, presumably less than 1 fM.
(c) (c) ZPB CP8 = Zetaproteobacteria, *Mariprofundus ferrinatatus* CP-8 ¹³⁸

6 Table 2: Fe(II) sediment fluxes estimated from M77/1 water column data using a single layer 1D model.

Station	Water Depth (m)	Sediment ^a flux (mmol m ⁻² y ⁻¹)	Integrated Fe(II) ^b Water Column (μmol m ⁻²)	C ₀ ^c [Fe(II)] nM	Oxidation ^d Rate (s ⁻¹)	K _z (cm ² s ⁻¹)
532/MUC 47&48	143	n.d.	219.4 (to 60 m)	13.5 ± 1.2	n.d.	n.d.
556/MUC 54&55 (deeper)	315	0.59	281.2 (to 130 m)	2.4 ± 0.4	6.7 ± 0.7 x 10 ⁻⁸	9.0 ± 1.1
569/MUC 66	185	3.48	1383.5 (to 50 m)	18 ± 2.1	8.0 ± 0.8 x 10 ⁻⁸	4.7 ± 1.4
575/BIGO T4 & T5	304	8.44 (T4) 12.95 (T5)	1374.4	21.3 ± 2.0	1.9 ± 0.2 x 10 ⁻⁷	8.1 ± 1.8
599/MUC 61	295.5	5.39 ^f	198.1 (to 60 m)	8.7 ± 0.4	8.6 ± 0.8 x 10 ⁻⁷	4.5 ± 0.4
618/MUC 83	153.8	59.2 ^f	1904.8	35.5 ± 4.0	9.9 ± 1 x 10 ⁻⁷	28.4 ± 7.2

^aData from Noffke (2012), from flux chamber or porewater diffusive flux estimates from a MUC (Multicorer) or BIGO (Biogeochemical Observatory) Lander. ^bTrapezoidal integration. ^cUsing the 1-layer model as described above. ^dOxidation rate calculated using equation x with the flux as estimated from the sediment core data. ^eUsing the 2-layer model as described in section x.y.z ^fChristian Hensen (GEOMAR) personal communication.

Figure Legends

Figure 1: Station locations for Fe(II) water column profiles occupied during M77-1 (red triangles). The location of stations where Fe(II) data was reported in earlier works are also shown: purple hexagons – Hong and Kester⁶⁵, white diamonds – Vedamati *et al.*⁵³. The location of the IMARPE time series station S4 is also shown in gray¹¹⁵. Bathymetry contours are shown for 50, 100, 200, 300 (blue), 400, 500, 1000, 1500 and 2000 m.

Figure 2: Monthly Chlorophyll – CMEMS (top, left) August 2008, (top, right) September 2008, (bottom, left) October 2008 and (bottom, right) November 2008. [Data product Global Ocean Chlorophyll (Copernicus-GlobColour) from Satellite Observations – Reprocessed]

Figure 3: Station 532 (left) Potential temperature (θ , red) and salinity (S, blue), (centre) Nitrite (brown squares), Fe(II) (black circles), the 1layer fit to the Fe(II) profile is shown in green, (right) Dissolved Oxygen (red) and H₂O₂ (blue circles).

Figure 4: Station 599 (left) Potential temperature (θ , red) and salinity (S, blue), (centre) Fe(II) (black circles), the 1layer fit to the Fe(II) profile is shown in green, (right) Dissolved Oxygen (red) and H₂O₂ (blue circles).

Figure 5: Station 618 (left) Potential temperature (θ , red) and salinity (S, blue), (centre) Fe(II) (black circles), the 2layer fit to the Fe(II) profile is shown in green, (right) Dissolved Oxygen (red) and H₂O₂ (blue circles).

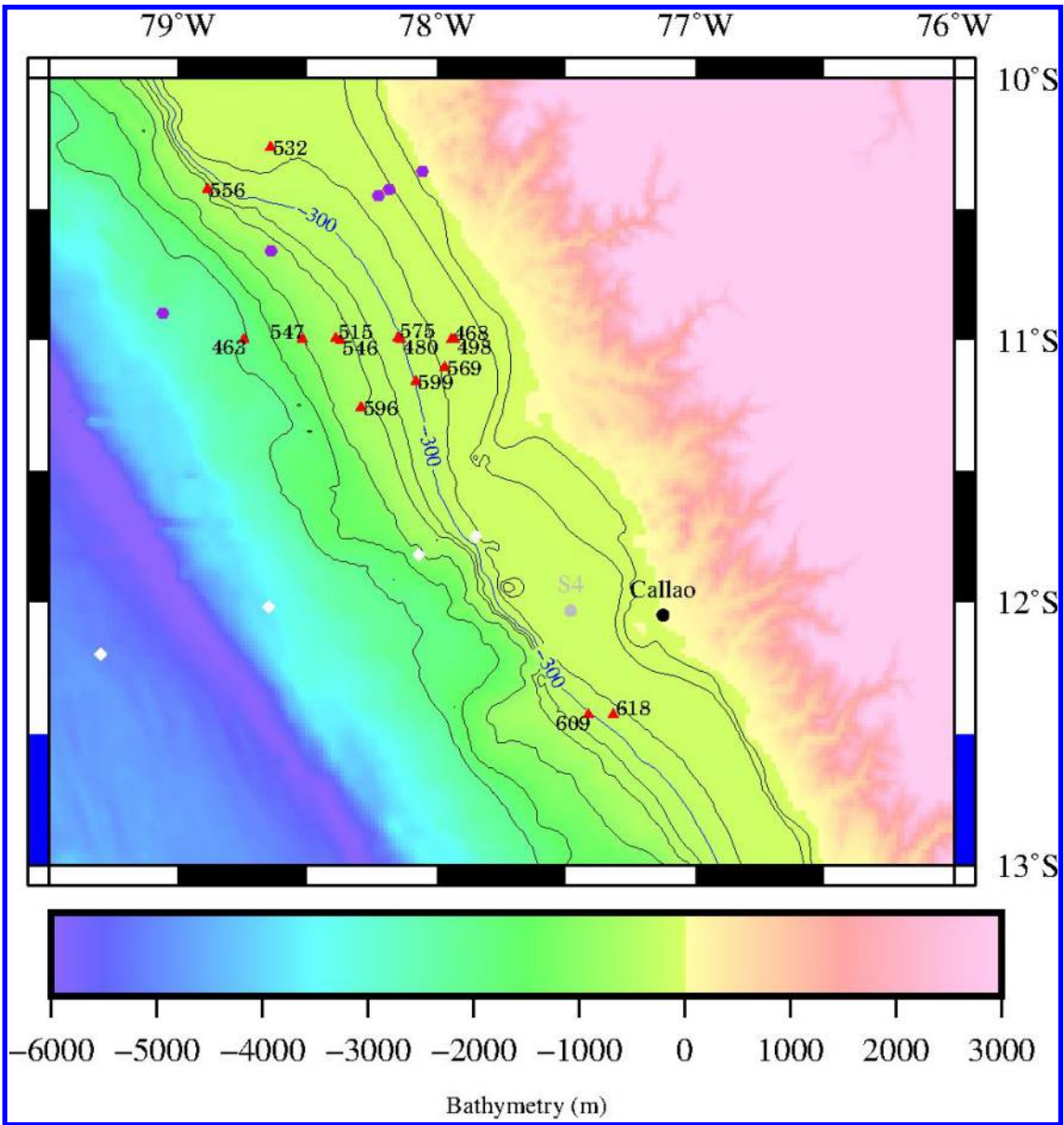


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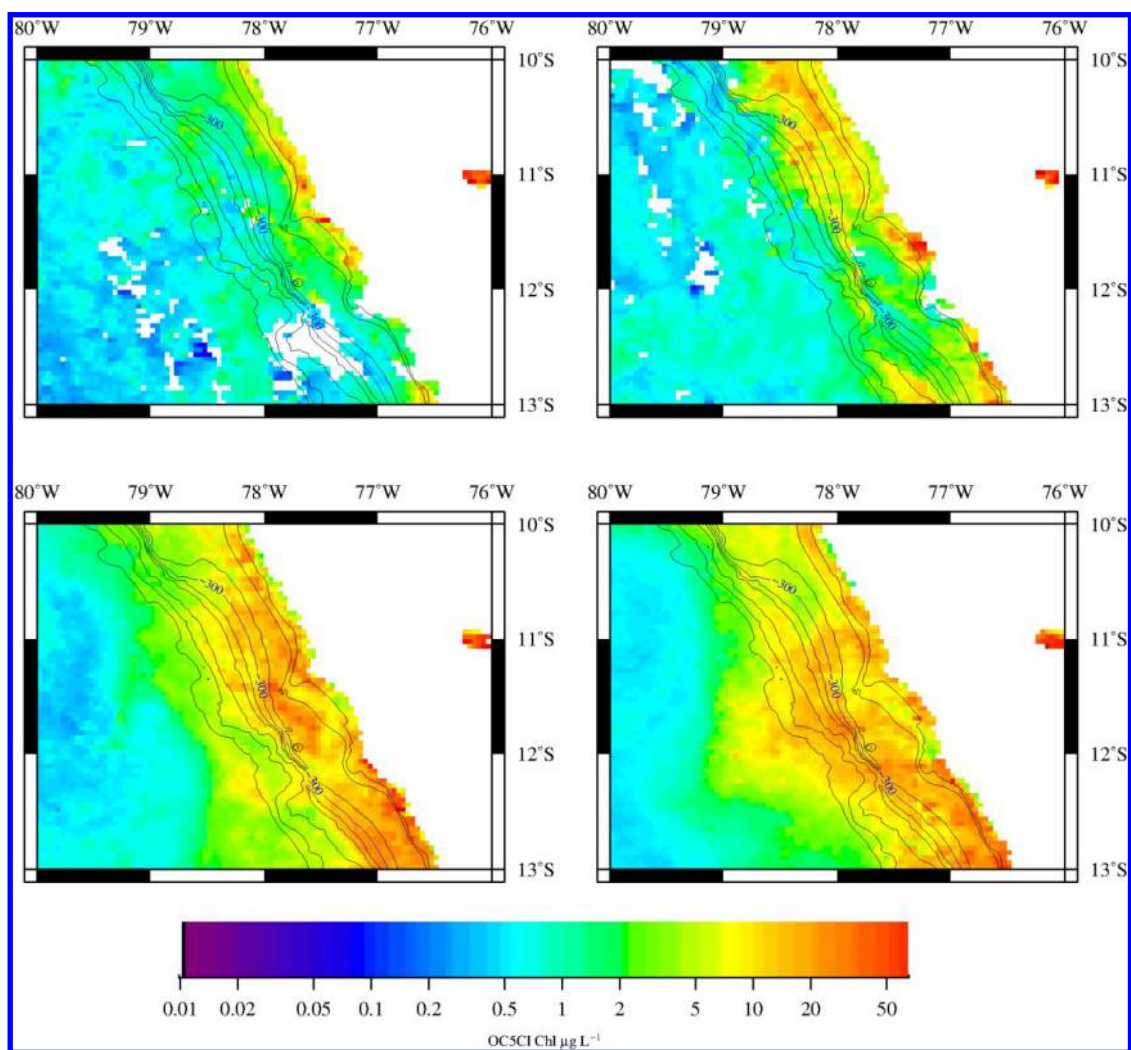


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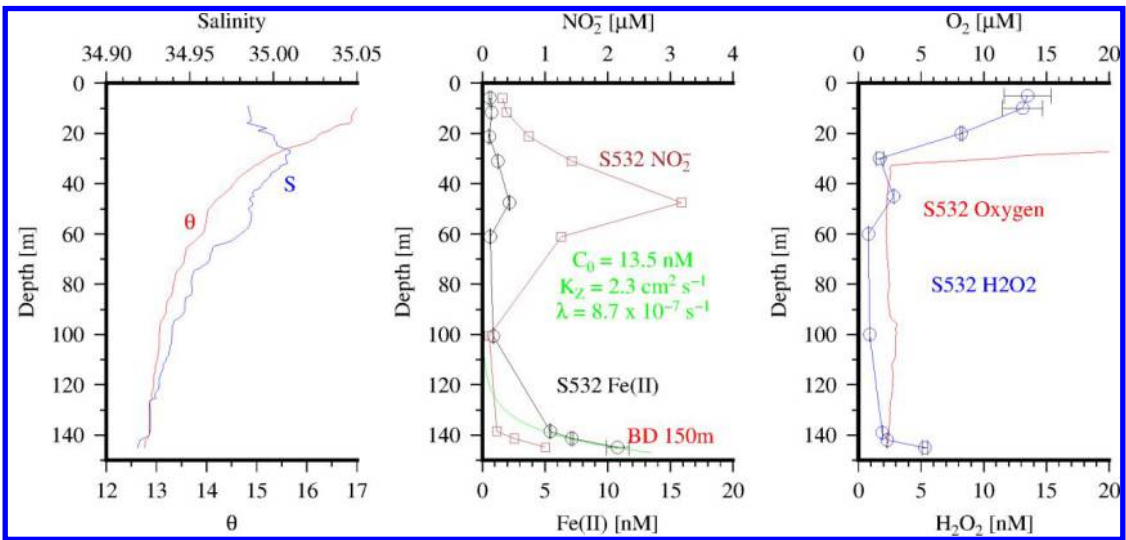


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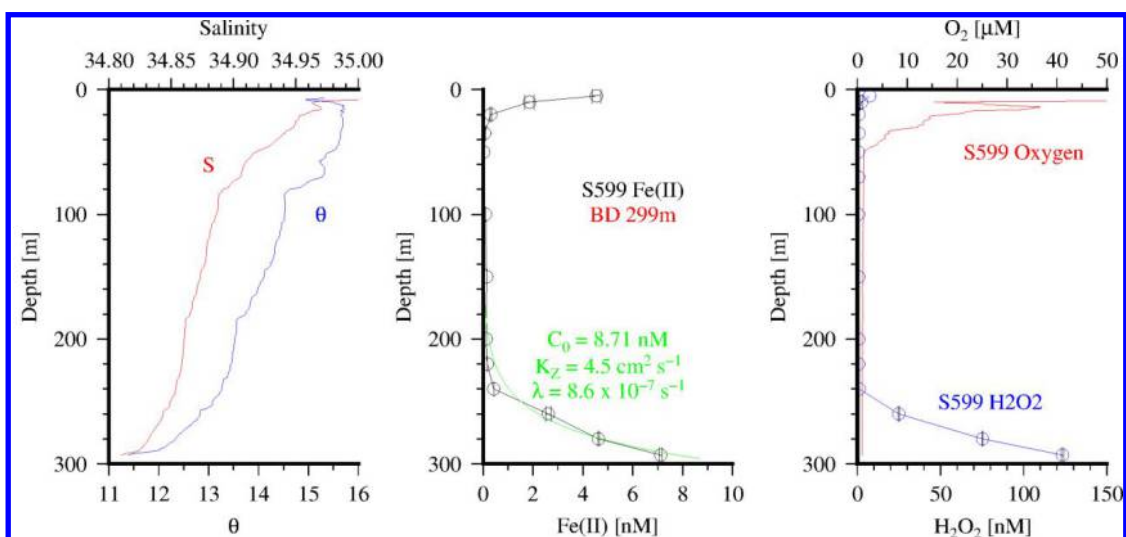


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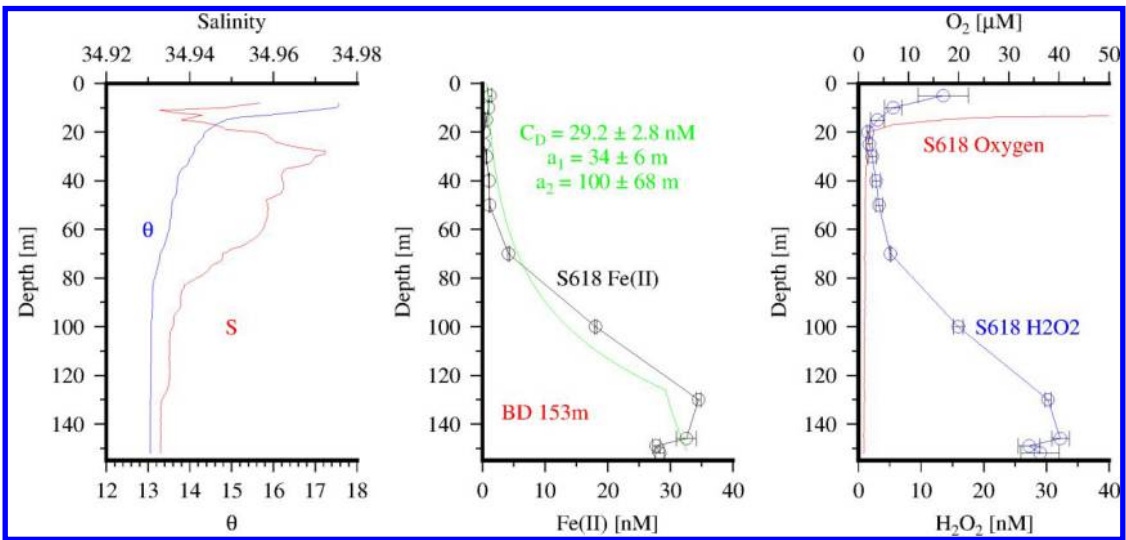
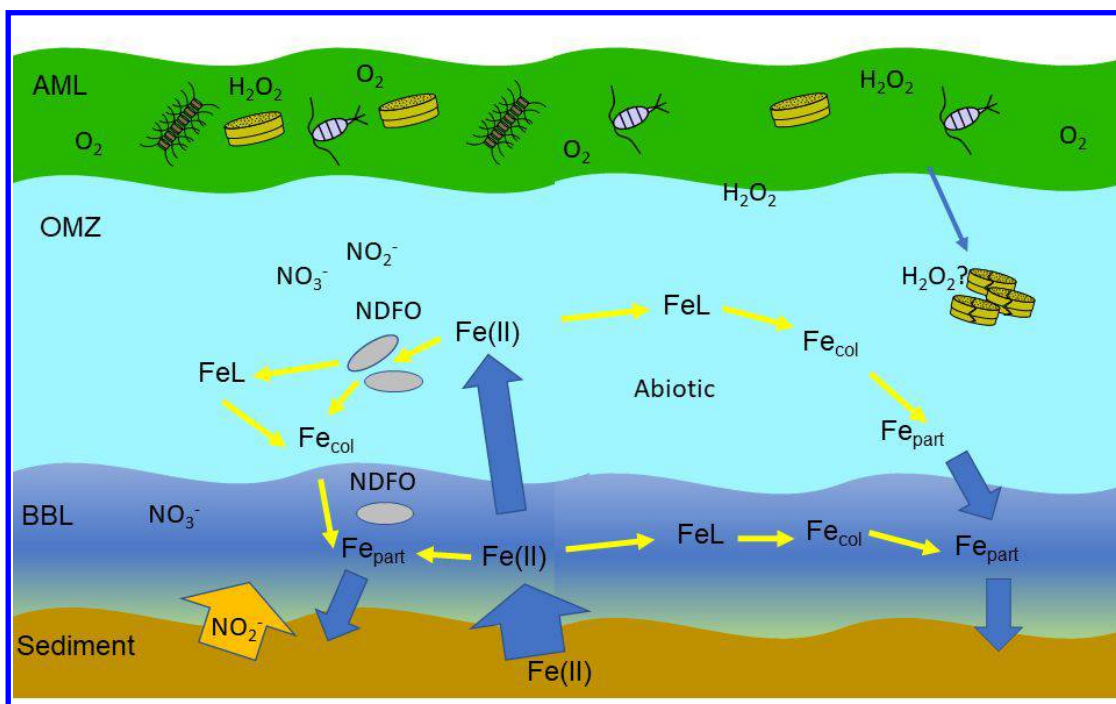
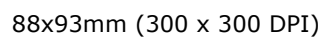
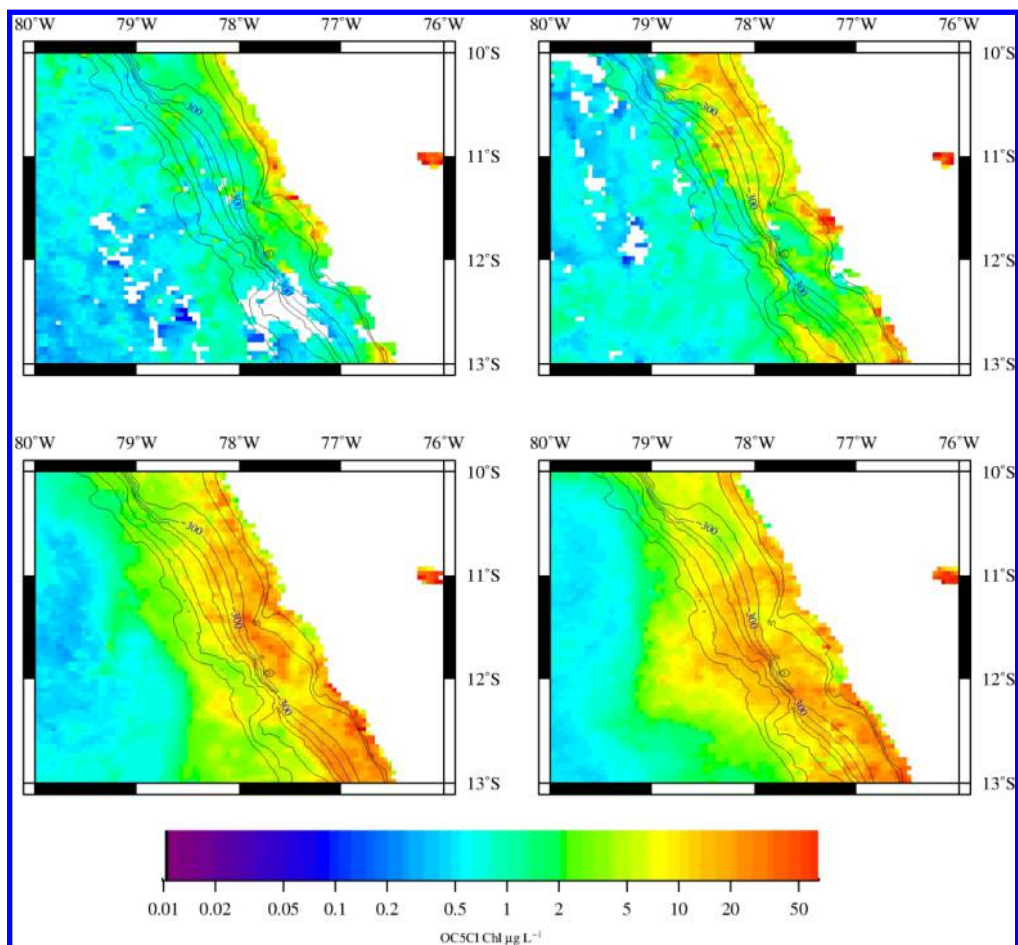


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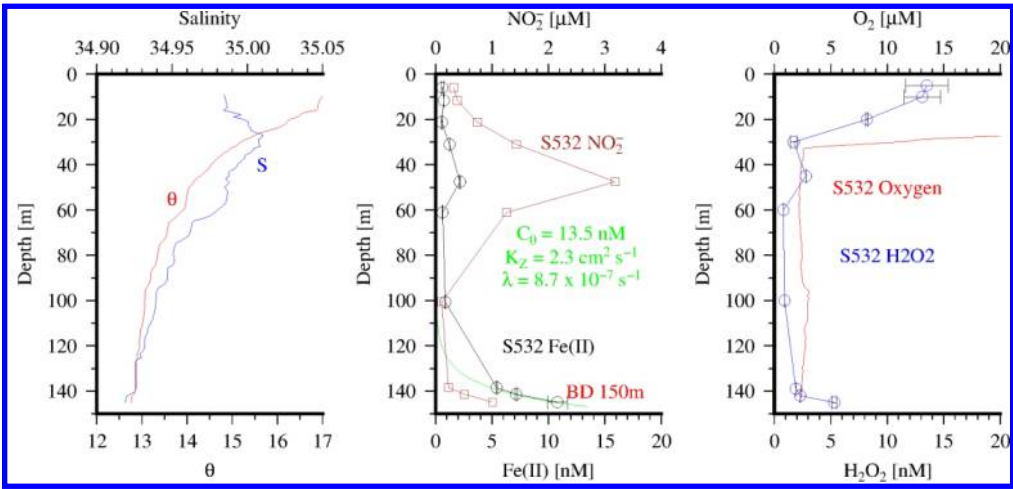


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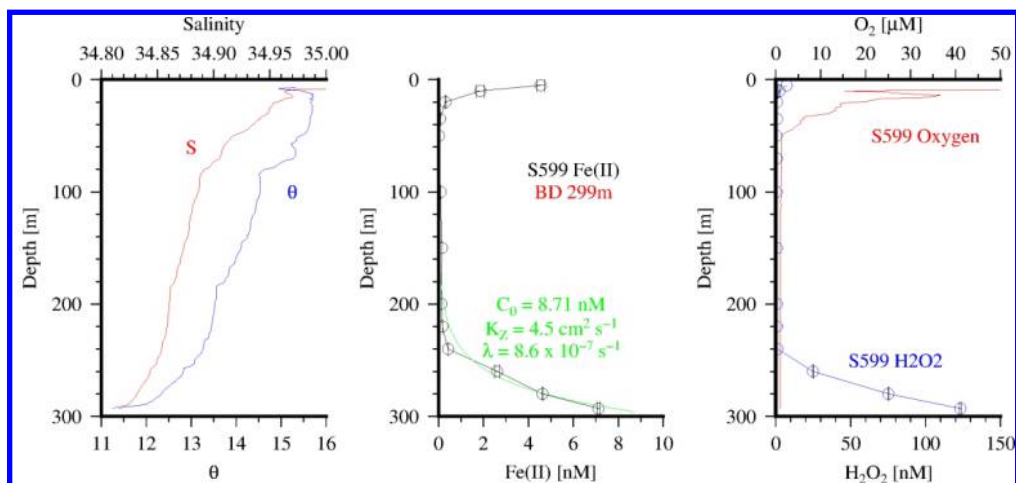




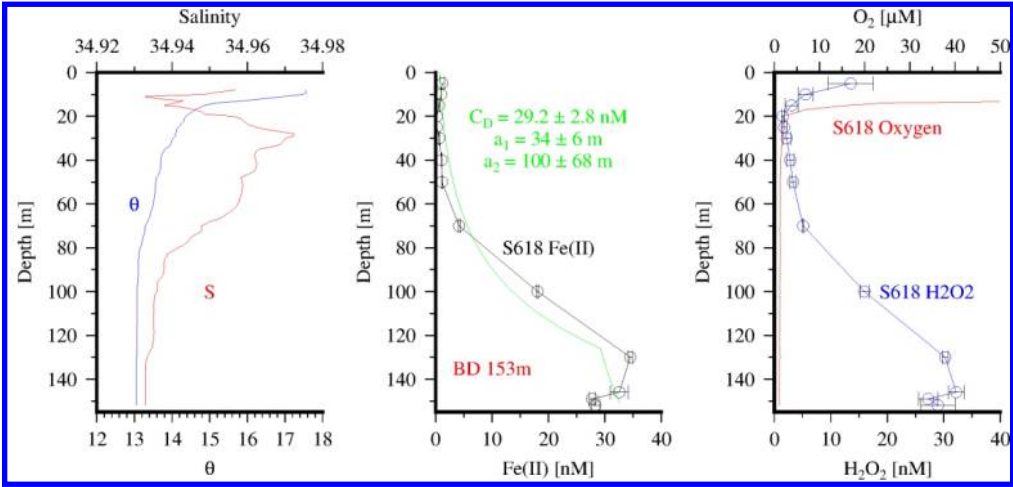
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