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Iron-binding by dissolved organic matter in the Western Tropical South Pacific Ocean (GEOTRACES TONGA cruise GPpr14)

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Iron (Fe) is an essential micronutrient for phytoplankton growth, but its scarcity in seawater limits primary productivity across much of the ocean. Most dissolved Fe (DFe) in seawater is complexed with Fe-binding organic ligands, a poorly constrained fraction of dissolved organic matter (DOM), which increase Fe residence time and impact Fe bioavailability. Here, we present the conditional concentration (L_{Fe}) and binding-strength (log $K_{Fe'L}^{cond}$) of Fe-binding ligands in the Western Tropical South Pacific (WTSP) Ocean during the GEOTRACES TONGA cruise (GPpr14). The transect crossed the Lau basin, a region subject to shallow hydrothermal Fe inputs that fuel intense diazotrophic activity, the oligotrophic South Pacific gyre, and the Melanesian basin. Organic speciation was analyzed by competitive ligand exchange adsorptive cathodic stripping voltammetry (CLE-AdCSV) using salicylaldoxime at 25 μ M. We found a high mean L_{Fe} of 5.2 \pm 1.2 nMeqFe (n = 103) across the entire transect, predominantly consisting of intermediate strength L2 ligands (84%; mean log $K_{Fe'L}^{cond}$ of 11.6 \pm 0.4), consistent with humic-like substances. DFe correlated with the humic-like component of the fluorescent DOM (HS-like FDOM), yet the electroactive Febinding humic-like substances (L_{FeHS}) accounted for only 20 \pm 13% of L_{Fe} in the mixed layer and $8 \pm 6\%$ in deep waters. Ligands were in large excess compared to DFe (mean excess ligand $eL_{Fe} = 4.6 \pm 1.1$ nMeqFe), suggesting poor stabilization of DFe inputs. High L_{Fe} (up to 9 nMeqFe) in samples close to hydrothermal sites could be due to detoxification strategies from plankton communities toward hydrothermally-fueled toxic trace metals other than Fe, with an apparent dilution of the DOM from the Lau basin into neighboring regions. We also observed a different peak potential of the Fe salicylaldoxime complex detected by CLE-AdCSV between the Lau and Melanesian basins, and between surface and deep waters. To our knowledge, this change in potential has not previously been

reported; whether this represents a novel detection of specificities in DOM composition merits further investigation. Competition between Fe and competing metals for ligand binding sites could favor DFe oxidation and precipitation near hydrothermal vents and explain the absence of strong Fe stabilization in the WTSP.

KEYWORDS

iron-binding ligands, hydrothermal, diazotroph, remineralization, humic substances, voltammetry, fluorescence

Introduction

Iron (Fe) is an essential micronutrient for phytoplankton growth because of its role in multiple metabolic processes (Twining and Baines, 2013). Biological Fe demand is such that its availability co-limits primary production across around 40% of the ocean (Moore et al., 2013; Browning and Moore, 2023). Fe scarcity is also explained by its very low solubility in seawater (Liu and Millero, 2002). Fe is quickly hydrolyzed and exported as oxyhydroxide aggregates in oxygenated conditions (Liu and Millero, 2002), and as Fe-sulfide (e.g., pyrite) in more anoxic environments such as hydrothermal vents (Rickard and Luther, 2007). However, despite its low solubility and inorganic scavenging, the Fe concentration found in the dissolved fraction (DFe; defined by the filter of 0.2 μ m or 0.45 μ m pore size) is higher than what is expected from its inorganic speciation (Liu and Millero, 2002). This is partly explained by extensive complexation of DFe by a fraction of the dissolved organic matter (DOM) pool, known as Fe-binding ligands, that enhance Fe solubility and increase its residence time in surface waters, thereby impacting its bioavailability to phytoplankton. It is generally accepted that most DFe (> 99%) is complexed to organic ligands (Gledhill and Buck, 2012). Knowledge of the nature and cycling of these ligands is required to understand and predict Fe distribution and bioavailability in seawater, but obtaining this information is challenging. Characterization of the Fe complexing fraction is difficult because of the sheer chemical diversity of DOM (Mentges et al., 2017; Dittmar et al., 2021) and the relatively low abundance of Fe-binding compounds, only exceeding DFe by around 1 nmol equivalent of Fe per liter (nMeqFe) in the global ocean (Gledhill and Buck, 2012; Boyd and Tagliabue, 2015). Various methods enable measurements of the major individual Febinding ligand groups, namely siderophores (Bundy et al., 2018; Boiteau and Repeta, 2022), humic-like substances (HS-like; Laglera and van den Berg, 2009; Pernet-Coudrier et al., 2013; Whitby and van den Berg, 2015; Sukekava et al., 2018), and exopolymeric substances (EPS; Hassler et al., 2011; Hassler et al., 2015; Norman et al., 2015). Those methods, however, have a limited resolution because of the diversity encountered within each group.

The challenge of identifying individual ligands within the range of metal-binding compounds composing DOM is overcome by

Competitive Ligand Exchange (CLE). This approach is based on the competition between the natural Fe-binding ligands and an added competitive ligand at increasing DFe concentrations. The titration of the natural Fe-binding ligands is monitored by adsorptive cathodic stripping voltammetry (AdCSV), quantifying the electrons needed to reduce the DFe that formed an electroactive complex with the added ligand. This approach allows the determination of the maximum amount of DFe that can be complexed by the natural DOM present within a sample, defined as the Fe-binding ligand concentration (L_{Fe}), and the averaged conditional stability constant (log $K_{Fe'L}^{cond}$) of the Febinding complexes within the detection window ($\alpha_{Fe'AL}$) applied. The detection window is defined as the product of the added ligand concentration ([AL]) forming the electroactive complex FeAL_n and of its binding affinity with DFe ($\beta_{Fe'AL}^{cond}$; $\alpha_{Fe'AL} = [AL]^n * \beta_{Fe'AL}^{cond}$). The log K^{cond}_{Fe'L} values obtained by CLE-AdCSV are commonly divided into classes, with L₁ corresponding to strong Fe-binding ligands (log $K_{Fe'I}^{cond}$ > 12), L₂ to intermediate ligands (10< log $K_{Fe'L}^{cond}$ < 12) and L₃ to weak ligands (log $K_{Fe'L}^{cond}$ < 10; Gledhill and Buck, 2012). These classes roughly correspond to the three major types of ligands identified in the environment, with siderophores falling into the L1 class, HS into L₂, and EPS into L₃ (Hassler et al., 2017), although classes can overlap and remain a simplistic view of the Fe organic speciation. For example, accumulating evidence suggests HS encompass multiple binding sites across the L1 and L2 ligand classes and are able to compete with siderophores (Fourrier et al., 2022; Gledhill et al., 2022; Sukekava et al., 2024). The log $K_{Fe'L}^{cond}$ of specific functional groups has also been determined and their contribution to the pool of Fe-binding ligands highlighted (e.g., Santana-Casiano et al., 2000; González et al., 2019). However, in natural samples, the binding ability of the DOM is a continuum of log $K_{Fe'L}^{cond}$ values rather than a specific value as determined by CLE-AdCSV (Town and Filella, 2000). Indeed, the log $K_{Fe'L}^{cond}$ determined by CLE-AdCSV is an average value of all the titrated ligands, and it is not well understood how the log $K_{Fe'L}^{cond}$ of each compound or functional group is weighted during the titration of a continuum of ligands. In all, while being mindful of limitations, this classification remains a helpful way to interpret and compare Febinding ligand data.

Fe-binding ligand distribution is often characterized by high L_{Fe} and $K_{Fe'L}^{cond}$ surface waters with maximum values coinciding with the chlorophyll-a maximum, and progressive decrease from the surface

until the oxycline due to adsorption onto particles (Kondo et al., 2007: Ibisanmi et al., 2011; Gledhill and Buck, 2012; Gerringa et al., 2015; Bundy et al., 2016). Phytoplankton production of L_{Fe} has been widely attested by concomitant L_{Fe} and chlorophyll-a fluorescence (Gledhill and Buck, 2012; Gerringa et al., 2015; Bundy et al., 2016; Hassler et al., 2017). L_{Fe} depletion can sometimes be observed in surface waters due to photochemical breakdown of ligands, but this process is highly variable (Barbeau et al., 2001; Powell and Wilson-Finelli, 2003; Croot et al., 2004), while atmospheric deposition such as dust and rain supply Fe-binding ligands to surface waters (Gerringa et al., 2006; Rijkenberg et al., 2008; Cheize et al., 2012; González et al., 2022). Below the oxycline, L_{Fe} increases through the breakdown of biogenic material by heterotrophic bacteria (Boyd et al., 2010) which produce Fe-binding ligands that are mostly attributed to HS due to their intermediate log $K_{Fe'L}^{cond}$ and relatively homogenous distribution in deep waters (van den Berg, 1995; Witter and Luther, 1998; van den Berg, 2006; Dulaquais et al., 2018; Whitby et al., 2020b). Heterotrophic bacteria produce siderophore compounds, associated with the breakdown of sinking particles (Cordero et al., 2012; Bundy et al., 2016; Velasquez et al., 2016; Hassler et al., 2017) while the resuspension of sediments and hydrothermal activity are important sources of Febinding ligands to bottom waters (Jones et al., 2011; Gerringa et al., 2015; Tagliabue and Resing, 2016. Buck et al., 2018; Sukekava et al., 2023).

The classification of binding strength presented by Gledhill and Buck (2012) has been used to compare basin scale datasets of Febinding ligand characteristics from samples collected in the West and North Atlantic Oceans (Buck et al., 2015; Gerringa et al., 2015), and in the Tropical Pacific Ocean (Buck et al., 2018). In addition to a general trend of decreasing log $K^{cond}_{Fe'L}$ and increasing L_{Fe} with depth and water mass aging, L1 has been shown to correlate with increasing DFe concentrations around hydrothermal vents in the Atlantic and Pacific Oceans (Buck et al., 2015; Buck et al., 2018) and with vent communities supplying siderophores specifically (Hoffman et al., 2023). L1 ligands were also observed in hydrothermal fluid samples collected at the New Hebrides Island arc in the Western Tropical South Pacific (WTSP) Ocean with concentrations reaching the micromolar range for L_{Fe} and DFe (Kleint et al., 2016). These authors also detected weaker L₃ ligands in several WTSP Ocean hydrothermal samples, while intermediate L₂ ligands have been observed in hydrothermal plumes in the Mariana Back Arc of the Western Pacific (Wang et al., 2021) and in the Southern Ocean (Hawkes et al., 2013).

In hydrothermal fluids, concentrations of DFe and Fe-binding ligands can vary from the nM to μ M range depending on the initial fluid chemistry (e.g., Hawkes et al., 2013; Resing et al., 2015; Kleint et al., 2016; Tilliette et al., 2022). The chemistry of hydrothermal fluids is geographically diverse and variable over time, and sampling in such dynamic environments is challenging, making hydrothermal Fe-binding ligand chemistry complicated to constrain (Kleint et al., 2019). In terms of DFe, it is well known that the free ionic Fe²⁺ initially found in hot and acidic hydrothermal fluid is quickly oxidized and precipitated during mixing with seawater by the formation of Fe-sulfides and Fe-oxyhydroxide minerals (Millero et al., 1987; Kleint et al., 2017),

but it can also be adsorbed on organic matrices or stabilized by Febinding ligands (Toner et al., 2009; Sander and Koschinsky, 2011; Kleint et al., 2017; Santana-Casiano et al., 2022). The inorganic transport of DFe as Fe-sulfide nanoparticles and colloidal Feoxyhydroxides has also been observed and the accessibility of these fractions to Fe-binding ligands is controversial (Yücel et al., 2011; Yücel et al., 2021). Recent work showed that electroactive humic component of Fe-binding ligands (L_{FeHS}) can solubilize a large fraction of freshly formed Fe-oxyhydroxides, but this decreases rapidly with aging of the inorganic particles. This experiment was performed at constant temperature and pH (Dulaquais et al., 2023) but is yet to be addressed under variable physicochemical conditions such as during the dilution of hydrothermal plumes. The variable interplay between the inorganic and organic processes impacting the amount of DFe stabilized and transported away from hydrothermal sources is an essential aspect to constrain, as vents could account for an estimated 12-22% of the deep-ocean DFe budget (Bennett et al., 2008).

In the WTSP Ocean, the subduction of the Pacific Plate beneath the Australian Plate has formed the Tonga volcanic arc, one of the most seismically active subduction zones which consequently hosts a high density of submarine volcanoes and associated hydrothermal vents (German et al., 2006; Timm et al., 2013). The submarine hydrothermal activity fuels the area with DFe (Guieu et al., 2018; Tilliette et al., 2022), supporting the growth of nitrogen fixers (diazotroph). Some of the highest rates of dinitrogen (N₂) fixation ever observed in open ocean have been measured in the area (Bonnet et al., 2017; Bonnet et al., 2018), specifically in the Lau basin, but the mechanistic link between shallow hydrothermal Fe inputs and intense N2 fixation has only been recently demonstrated (Bonnet et al., 2023b). The possibility to investigate the connection between hydrothermal sourcing of trace metals and the fueling of diazotroph activity motivated the GEOTRACES process cruise GPpr14 (TONGA cruise; Guieu and Bonnet, 2019, https:// doi.org/10.17600/18000884), which crossed the Melanesian basin, the Lau basin, and the western part of the South Pacific gyre between October 31st and December 5th, 2019.

The TONGA transect presented the opportunity to quantify, for the first time in this region, L_{Fe} and log $K_{Fe'L}^{cond}$, and to explore the relation between Fe-binding ligands and DFe distribution in the area. Several complementary analyses targeting specific fractions of the DOM pool were also conducted, notably regarding HS-like material. HS are of specific interest as they have been estimated to represent around 50% of the DOM (Zigah et al., 2017; Fourrier et al., 2022) with a smaller fraction, around 5% of DOC, able to bind with Fe (Laglera and van den Berg, 2009; Fourrier et al., 2022). The diversity of the HS-like material is responsible for various properties, notably in terms of electroactivity and photoreactivity, the latter in terms of absorbance and fluorescence (CDOM and FDOM, respectively). It has been shown that electroactive Fe-HS complexes $(L_{\mbox{\scriptsize FeHS}})$ are able to control Fe solubility in a variety of systems (Dulaquais et al., 2018; Sukekava et al., 2018; Laglera et al., 2019; Whitby et al., 2020b; Fourrier et al., 2022). Similarly, the HS-like fluorescent component of the DOM (HS-like FDOM) often correlates with DFe distributions (Tani et al., 2003; Ohno et al., 2008; Hioki et al., 2014; Jia et al., 2021).

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The complementarity and/or overlapping of these two different operationally defined fractions of HS is, however, not well constrained. While covariation of L_{FeHS} and HS-like FDOM is observed in estuarine waters (Lee et al., 2023), opposite trends are observed in open ocean surface waters, suggesting that the microbial aerobic respiration of the DOM increases the fluorescent yields and lowers the electroactivity of the HS fraction of the DOM with depth (Fourrier et al., 2022).

We present here the results obtained for Fe-binding ligand characteristics during the TONGA cruise using the same competitive ligand method that was used in both the Equatorial Pacific (Buck et al., 2018) and the North Atlantic Oceans (Buck et al., 2015). We relate the distribution observed in L_{Fe} and log $K_{Fe'L}^{cond}$ to other parameters such as DFe, apparent oxygen utilization (AOU), dissolved organic carbon (DOC), L_{FeHS} and HS-like FDOM to identify the processes impacting Fe-binding ligands and HS-like compounds, and to constrain their impact on DFe distribution in the WTSP Ocean.

Material and methods

Sampling strategy during the **TONGA** cruise

Samples were collected during the TONGA GEOTRACES process study in the WTSP Ocean (GPpr14). The section presented here is composed of 8 main stations between the Melanesian basin and the South Pacific gyre (Figure 1), with additional casts performed around two known hydrothermal sites, LD 5 and LD 10, identified by acoustic detection (Bonnet et al., 2023b). Here, we present data for one of the hydrothermal stations, LD10, for which stations LD 10-T1, T2 and T3 were sampled at 15, 8, and 2 km from the hydrothermal source, respectively. The western part of the Lau basin was sampled at stations 4, 11 and 12. Stations 2 and 3 were located in the Melanesian basin, on the western side of the transect. Stations 6, 7 and 8 were located in the South Pacific gyre for comparison with oligotrophic conditions. Station 6 was located East of the Tonga arc, but on the western side of the Tonga trench, where the topography is expected to limit water exchange between the Lau basin and the South Pacific gyre (Talley et al., 2011).

Sample collection and storage

The seawater samples for DFe and Fe-binding ligand analyses were collected using GO-FLO bottles mounted on a Trace Metal Clean Rosette (TMR; General Oceanics Inc., Model 1,018 Intelligent Rosette) transferred into a trace metal clean van (class-100) and pressurized with 0.2 µm-filtered nitrogen (Air Liquide) for subsampling. GO-FLO bottles and all the sampling material were cleaned before the cruise following the GEOTRACES cookbook (Cutter et al., 2017). Dissolved fractions for the analysis of DFe and Fe-binding ligands were sampled through 0.45 µm acid-cleaned polyethersulfone filters (Supor). For DFe, the cleaning procedure of the bottles used for seawater collection, sample storage, handling and analyses are detailed in Tilliette et al. (2022).

For Fe-binding ligands, 250 mL low-density polyethylene (LDPE; Nalgene) bottles were left to soak for one week in a 2% surfactant bath (Decon), one week in a 1 M HCl bath (laboratory reagent grade 32%, FisherScientific) and one week in a 0.1 M HCl bath, before being left to dry in a laminar flow hood (Class-100) and double-bagged. Between each step, the bottles were rinsed 5-times with Milli-Q water (resistivity > 18.2 M Ω .cm). The samples were collected after rinsing the bottles 3 times with around 20 mL of filtered sample and frozen to -20°C immediately after sampling. Samples were thawed at room temperature in the dark and swirled before analysis.

The seawater samples for dissolved oxygen (O2), DOC and FDOM analyses were collected in glass bottles from a classical rosette equipped with twenty-four 12 L Niskin bottles. The samples were immediately filtered under low vacuum (< 50 mm Hg) through pre-combusted 25-mm fiberglass filters (GF/F, size pore: ~0.7 µm, Whatman) using glass filtration apparatus. For DOC analyses, filtered samples were transferred into 10 mL glass ampoules and acidified with 20 µL of sulfuric acid (H₂SO₄, 95-98%, Sigma-Aldrich). Ampoules were then flame-sealed and stored in the dark at 4°C pending analysis. For FDOM measurements, filtered samples were



Map of the studied area, cruise transect and sampling stations for Fe-binding ligand characteristics during the TONGA cruise (GPpr14; Guieu and Bonnet, 2019, https://doi.org/10.17600/18000884). The small-dashed black line corresponds to the transect drawn to present the data. Colors, symbols and shapes correspond to the splitting of the dataset for presentation and interpretation. Geographic features (in black) include the main islands, the Tonga trench and the location of the Late'iki site.

transferred into 100 mL amber glass bottles with clean Teflon-lined caps and stored at -20°C in the dark before analysis. Before use, all glassware was left to soak for 24 h in a 1 M HCl bath, rinsed thoroughly with Milli-Q water, then combusted at 450°C for 6 h, and finally rinsed three times with the respective sample before filling.

Analyses

Detailed description of the analysis and interpretation for L_{FeHS} , CDOM/FDOM and DFe can be found in connected papers (Tilliette et al., 2022; Dulaquais et al., 2023; Tedetti et al., in prep.¹). For L_{FeHS} , CDOM/FDOM, DFe, dissolved organic carbon and Febinding ligands, the seawater was filtered in-line using an acid-cleaned 0.45 µm polyethersulfone filter (Supor).

Dissolved oxygen

 O_2 concentration was measured following the Winkler method (Winkler, 1888) modified by Carpenter (1965) and Carritt and Carpenter (1966), with potentiometric endpoint detection using a Titrino 716 DMS (Metrohm; Oudot et al., 1988). The recommendations from Langdon (2010) were followed for sampling, reagent preparation and analysis. The thiosulfate solution was calibrated by titrating it against a potassium iodate certified standard solution of 0.0100 N (CSK standard solution; WAKO). The reproducibility, expressed as the standard deviation of replicate samples, was 0.8 μ M (n = 15, mean = 195.4 μ mol kg⁻¹. Apparent Oxygen Utilization (AOU, in μ M) was calculated from the difference between O₂ solubility concentration (at P = 0 dbar) estimated with the Benson and Krause coefficients (Garcia and Gordon, 1992) and *in-situ* O₂.

Dissolved organic carbon

DOC concentration was measured in two ampoule replicates on a Shimadzu TOC-V analyzer according to Sohrin and Sempéré (2005) and Fourrier et al. (2022). Before injection, the GF/F-filtered and acidified samples were bubbled for 2 min with CO_2 -free air to purge inorganic carbon. The accuracy and system blank of the instrument were determined by the analysis of certified water references (batch 19, lot #03–19, Hansell Laboratory, Miami, USA). The nominal precision of the analysis procedure was within 2%.

Iron-binding ligands

The theory of the CLE-AdCSV approach has been thoroughly detailed in previous work (e.g., Gledhill and van den Berg, 1994; Rue and Bruland, 1995; Abualhaija and van den Berg, 2014; Gerringa et al., 2014; Pižeta et al., 2015; Mahieu, 2023). We used the added ligand salicylaldoxime (SA; 98% Acros Organics) at 25 μ M following the procedure described in Mahieu (2023). For data interpretation the side reaction coefficient (β_{FeSA_2}) of 11.1 leading to $\alpha_{Fe'AL}$ of 79 was used, as in previous basin scale work (Buck et al.,

2007; Buck et al., 2015; Buck et al., 2018). A borate buffer (boric acid, analytical reagent grade, Fisher Scientific) at 1 M in 0.35 M of ammonia (NH₄OH; 29% Laporte) was used at 10 mM for a final pH of 8.2 in the sample (Rue and Bruland, 1995; Buck et al., 2007; Abualhaija and van den Berg, 2014; Buck et al., 2015; Buck et al., 2018; Mahieu, 2023). The following equilibration procedure was applied: natural ligands were left to equilibrate with increasing additions of DFe in buffered samples for a minimum of 2 h, with SA added at least 15 min before the start of the analysis. The titrations were performed with 16 aliquots, with DFe additions of 0, 0, 0.75, 1.5, 2.25, 3, 3.5, 4, 4.5, 5, 6, 7, 8, 10, 12 and 15 nM. The DFe additions were adjusted from Buck et al. (2018) after preliminary tests showing emergence of the FeSA₂ reduction current around 3 nM of DFe added (nMeqFe). The sensitivity was estimated using the three last linear additions of the titration of the seawater sample. The apparatus and procedures for the conditioning of cells and tubes (Trace Metal Free, Labcon) are described in Mahieu (2023) and Mahieu et al. (2024), along with a detailed description of the set up and data treatment.

Statistical treatments

The variables considered in this study did not follow a normal distribution, and were inter-dependent and non-linear, calling for the application of non-parametric statistical methods. The significance of the difference between water masses and regions was tested using the Wilcoxon-Mann-Whitney test, using the Python function 'scipy.stats.mannwhitneyu'. The difference between two samples is defined by p values< 0.05. Full p value matrices can be found in Supplementary Information 2, 3. The Spearman rank correlation (p; Spearman, 1904) analysis was chosen to compare the Fe-binding characteristics and the other parameters described hereafter as it has been previously used for the investigation of Fe-binding ligands and FDOM (e.g., Heller et al., 2013; Genovese et al., 2018). The calculation was performed using the Python function 'scipy.stats.spearmanr' (version 3.9). For the Spearman test, p from 0.00 to 0.19 were attributed to very weak correlation, ρ from 0.20 to 0.39 to weak correlation, ρ from 0.40 to 0.59 to moderate correlation, ρ from 0.60 to 0.79 to strong correlation, and ρ from 0.80 to 0.99 to very strong correlation ($\rho = 1.00$ being a perfectly monotonic correlation). We considered the correlation to be significant for p< 0.0001, at least one order of magnitude lower than in other studies discussing Spearman correlation matrices including Fe-binding ligands data to ensure a strong level of confidence in the interpretation of the dataset (e.g., Heller et al., 2013; Genovese et al., 2018; bold values in Table 1).

Results

Hydrography

The water mass composition and circulation have been thoroughly described by Tilliette et al. (2022). We present here a simplistic description of the water column based on O_2 fluctuations (Figures 2, 3A). During the TONGA transect, surface waters were characterized by high O_2 due to primary production and

¹ Tedetti, M., Guigue, C., Mahieu, L., Fourrier, P., Dulaquais, G., Benavides, M., et al. Chromophoric and fluorescent dissolved organic matter in the Western Tropical South Pacific.

Considered data	DFe (nM)	L_{Fe}	L _{FeHS}		DOC (µgC L ⁻¹)	HS-like FDOM (QSU)
		(nMeqFe)	(nMeqFe)	(µgeqC L ⁻¹)		
Mixed layer	0.45 ± 0.50	5.3 ± 1.4	1.3 ± 0.7	40.2 ± 23.1	883 ± 53	1.7 ± 0.5
	n = 60	n = 24	n = 31		n = 40	n = 36
Deep waters	0.51 ± 0.30	5.2 ± 1.1	0.5 ± 0.3	12.5 ± 10.1	580 ± 126	2.5 ± 0.7
	n = 178	n = 79	n = 75		n = 129	n = 94

TABLE 1 Mean values for DFe, L_{Fer}, L_{Fer}, DOC and HS-like FDOM for the mixed layer and deep waters.

Estimates of Fe-binding equivalent concentrations for L_{FeHS} were calculated using 16.4 nmolFe.mgFA⁻¹ from Sukekava et al. (2018). Estimates of carbon equivalent were calculated using the official value of 0.5244% gC/gSRFA (International Humic Substances Society).

atmospheric exchange enhanced by physical mixing in the mixed layer, which ranged from 70 to 140 m deep during the cruise. The water mass below the mixed layer corresponds to the Subtropical Underwater (STUW) in the South Pacific gyre and Lau basin, and to the Western South Pacific Central Water (WSPCW) in the Melanesian and Lau basins. Both STUW and WSPCW are characterized by a maximum of salinity (S) and minimum of O₂ around 200 m, the latter being due to microbial respiration (e.g., remineralization). Below these two water masses is the Antarctic Intermediate Water (AAIW), with a S minimum and an O₂ maximum (Figure 2) also corresponding to the thermocline at around 650 m depth (Kawabe and Fujio, 2010; Talley et al., 2011). Below the AAIW, S increases and temperature decreases. O2 concentrations below 150 µmol kg-1 is characteristic of the Pacific Deep Water (PDW), originating from the upwelling of Antarctic Bottom Water (AABW) in the middle of the Pacific Ocean, mixing with the O2 depleted waters from the Pacific oxygen minimum zone and with the deep waters from the oxygen minimum zone off Chile in the Eastern South Pacific Ocean



FIGURE 2

Profile of σ_0 (black full line), O_2 (red full line, diamond), T (temperature, orange dash line), and S (salinity, blue dot line) at station 7 with the water mass separations applied in this study, namely the mixed layer (ML), the Subtropical Underwater (STUW), the Western South Pacific Central Water (WSPCW), the Antarctic Intermediate Water (AAIW), the Pacific Deep Water (PDW) and the Lower Circumpolar Deep Water (LCDW). The WSPCW, not present at ST7, is also mentioned to not omit any water mass investigated in this work.

(Silva et al., 2009). AAIW and PDW supposedly reach the Lau and Melanesian basins from a branch of the South Pacific gyre flowing from the North of the Lau basin following south-westward currents (Summers and Watling, 2021; Tilliette et al., 2022). However, the circulation in and between the Lau and Melanesian basins is variable, and northeast currents have been observed in the Melanesian basin (Komaki and Kawabe, 2007). In the South Pacific gyre, deeper than 3500 m, O₂ increases to around 190 μ mol kg⁻¹ due to the intrusion of Lower Circumpolar Deep Water (LCDW) from the southeast (Kawabe and Fujio, 2010). Over the whole transect, O₂ concentrations range from 121 μ mol kg⁻¹ in the PDW of the Lau basin, to 211 μ mol kg⁻¹ in the AAIW of the South Pacific gyre (Figure 3A).

DOC

The distribution and range of DOC concentrations measured in this study are typical to what has been previously reported in the Tropical South Pacific Ocean (e.g., Buck et al., 2018; Fourrier et al., 2022). DOC concentrations ranged from 35 to 80 μ M (Figure 3B), with the highest values observed in the mixed layer typical of primary production, ranging from 60 to 80 μ M. DOC concentrations decreased with depth due to microbial remineralization to reach typical deep-water values, ranging from 35 to 45 μ M in the PDW (and LCDW at stations 7 and 8). Typical deep values are reached within the STUW/WSPCW in line with microbial aerobic respiration at most stations (Figures 2, 3), except at stations 3, 4 and 7, where the concentrations gradient reaches depths of up to 1000 m.

DFe

DFe distribution has been thoroughly described in Tilliette et al. (2022). For the dataset considered in this work, DFe concentrations ranged from 0.13 nM in the mixed layer of the Western Lau basin to 3.13 nM in the mixed layer of the Eastern Lau basin (Figure 4A). Higher DFe concentrations, up to 50 nM, were observed at another location (station 5; Tilliette et al., 2022), however, no data is available for Fe-binding ligand characteristics at this station. The Melanesian basin and the South Pacific gyre show DFe depletion in surface waters attributed to predominant removal processes and limited supply



and 3), the Western Lau basin (stations 12, 4 and 11), the Eastern Lau basin (stations LD10-T1, LD10-T2 and LD10-T3) and the South Pacific gyre

(stations 6, 7, and 8). See Figure 1 for more details on the sampling locations.

(Tilliette et al., 2022), and an increase at depth with mineralization in the range of previously reported open-ocean values (e.g., Tagliabue et al., 2012; Resing et al., 2015; Tonnard et al., 2020). The Western Lau basin shows similar DFe depletion in subsurface waters except for an enhancement at station 11 in the WSPCW. Another strong DFe enhancement is observed in the mixed layer of the Eastern Lau basin. The highest values observed in this study are considered to be solely hydrothermal, while deep waters of the Eastern and Western Lau basin show anomalies of DFe enrichment related to either remineralization, island effect, benthic fluxes and/or hydrothermal activity (up to 1.5 nM; Tilliette et al., 2022). Similar concentrations were previously reported for bottom enrichment related to hydrothermal and shelf inputs (e.g. Klunder et al., 2011; Klunder et al., 2012; Resing et al., 2015; Tonnard et al., 2020).

HS-like FDOM

The HS-like FDOM component identified here through EEM/ PARAFAC (λ_{Ex1} , $\lambda_{Ex2}/\lambda_{Em}$: 235, 315/436 nm), which are referred to as peaks A + M/C (Coble, 1996; Hudson et al., 2007; Coble et al., 2014) or component 3 (Ishii and Boyer, 2012) in different DOM fluorophore classifications, has been reported in various coastal and marine environments (Ferretto et al., 2017; Tedetti et al., 2020). HS-like FDOM content ranged from 1 QSU in the mixed layer of the Western Lau basin to 4 QSU in the PDW of the Melanesian basin (Figure 4B). HS-like FDOM systematically increased with depth, reflecting its loss by photobleaching or extinction of its fluorescence in the photic layer, and its production in deep waters through the remineralization of organic matter (Nelson et al., 2010; Heller et al., 2013; Yamashita et al., 2017). The depth profiles of HS-like FDOM along the transect (i.e., lower values in surface waters, higher at depth) were thus opposite to those of DOC, a phenomenon already reported for the Atlantic and North Pacific Oceans, as well as at our station 8 in the South Pacific gyre (Omori et al., 2010; Stedmon and Alvarez-Salgado, 2011; Yamashita et al., 2017; Cao et al., 2020; Fourrier et al., 2022). High HS-like FDOM signals were observed in the deep waters at stations 2, 3, 4, LD10-T1, LD10-T2, the highest signal being observed at station 8.



L_{FeHS}

Details on the distribution and determination of electroactive HS (eHS) in the area during the same cruise are fully described elsewhere (Dulaquais et al., 2023). We show here the eHS determined by complexation with Fe (Dulaquais et al., 2023), referred to as L_{FeHS} , for comparison with HS-like FDOM and Febinding ligands (Figure 4C). L_{FeHS} are expressed in nMeqFe following the estimated Fe binding capacity of the fulvic acid standard used (16.4 nmolFe mgFA⁻¹; Sukekava et al., 2018). For the dataset considered in this work, L_{FeHS} concentrations ranged from 0.2 to 2.4 nMeqFe with a mean value of 0.6 ± 0.5 nMeqFe (n = 106). The highest values were observed in the mixed layer at all stations and concentrations decreased with depth down to relative minima (between 0.2 and 0.4 nMeqFe) in intermediate waters. Below, in the abyssal waters, concentrations increased again to values higher than 0.6 nMeqFe. Comparatively, high L_{FeHS} concentrations were detected through the entire water column in the oligotrophic waters of the South Pacific gyre collected at station 8 (Fourrier et al., 2022).

Fe-binding ligands

A single class of Fe-binding ligand was detected in all samples, with a mean L_{Fe} of 5.2 ± 1.2 nMeqFe and mean log $K_{Fe'L}^{cond}$ of 11.6 ± 0.4 (n = 103; Figures 5A, C, respectively). Log $K_{Fe'L}^{cond}$ ranged from 10.5 ± 0.2 in the AAIW of the Melanesian basin (station 2) to 12.7 ± 0.3 in the PDW of the Western Lau basin (station 11), encompassing the three ligand classes defined by Gledhill and Buck (2012). As such, 84% of our samples fall in the L₂ class, 13% in the L₁ class, and 4% in the L₃ class.

 $L_{\rm Fe}$ ranged from 2.8 \pm 0.4 nMeqFe in the mixed layer of the Melanesian basin (station 2) to 9.3 \pm 1.0 nMeqFe in the PDW of the Melanesian basin (also station 2). The co-occurrence of the maximal and minimal values at the same station illustrates the variability of the processes impacting the DOM in the area, discussed hereafter. High values of $L_{\rm Fe}$ reaching up to 9.0 nMeqFe were observed at two other locations, in the AAIW above the Tonga ridge in the South Pacific gyre (station 6), and in the mixed layer in the Lau basin (station LD 10-T3). At stations 2 and 6, higher $L_{\rm Fe}$ coincided with increased HS-like FDOM (Figure 4B).



Top 600 m and full depth distribution of (A) L_{Fe} , (B) eL_{Fe} and (C) $\log K_{Fe'L}^{cond}$ for the TONGA cruise (GEOTRACES GPpr14). The approximative water mass boundaries are defined by O₂ (see Figures 2, 3A). The section is separated following the description of the studied area (Figure 1).

In all samples, L_{Fe} was present in unusually large excess compared to DFe, with excess ligand ($eL_{Fe} = L_{Fe}$ - DFe) ranging from 2.6 to 8.6 nMeqFe (Figure 5B) and a mean eL_{Fe} of 4.6 ± 1.1 nMeqFe (n = 103).

Fe-binding ligands show relatively constant L_{Fe} and log $K_{Fe'L}^{cond}$ values along water masses and regions over the whole transect (Figure 6), but a few significant differences were observed (Mann-Whitney test, p< 0.05; Supplementary Material 2, 3). The significant differences for L_{Fe} were observed between the low values of the mixed layer of the Melanesian basin and the other regions, and between the high values of the PDW and both the mixed layer and the STUW/WSPCW in the Melanesian basin. Log $K_{Fe'L}^{cond}$ was significantly lower in the LCDW in the South Pacific gyre and all other water masses, suggesting specific Fe-binding characteristics of the DOM in this deep water mass.

Discussion

L_{Fe} and log $K_{Fe'L}^{cond}$ distribution in the WTSP

Our eL_{Fe} values (mean = 4.6 \pm 1.1 nMeqFe) are c.a. 2.5 times higher than the average reported in the Eastern and Central

Tropical South Pacific Oceans (Buck et al., 2018) using exactly the same experimental procedure. The large fraction of eL_{Fe} belonging to the L₂ class throughout the water column suggests a rather homogenous Fe-binding ability of the DOM, while the decreasing DOC concentration along the water column infers a refractory nature of the binding sites and an increase in binding site density with depth. We find that the ligand pool is dominated by L₂ ligands even though we use a high detection window, which can favor detection of the strongest ligands and miss contributions from weaker ligands. In comparison, it was found in the Eastern and Central Tropical South Pacific Oceans that L1 ligands were generally in excess compared to DFe in the first 1000 m, while excess L₂ and L₃ increased in deeper waters (Buck et al., 2018). The ligand distribution observed in these environments reflects the biological production of strong L1 in upper waters, and the loss and saturation of these L1 ligands as DOM is remineralized, resulting in increasing DFe and weaker L2 and L3 ligand concentrations in deep waters. In our study, a decrease in log $K_{Fe'L}^{cond}$ with depth was observed in the South Pacific gyre, but without change of ligand class, and no change in log $K_{Fe'L}^{cond}$ in the Melanesian and Lau basins (Figures 5, 6).



FIGURE 6

Box and Whiskers plots of L_{Fe} and log $K_{Fe'L}^{cond}$ split by the regions and water masses. The mean values of the entire dataset are represented by black dot lines. The hatched boxes correspond to water mass being significantly different compared to the same water mass in other regions (Wilcoxon-Mann-Whitney test, p < 0.05). Water mass acronyms can be found in Figures 2, 3A.

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The progressive decrease in log $K_{Fe'L}^{cond}$ with increasing depth and increasing distance from the Lau basin gives the impression of a dilution gradient from stronger to weaker ligands into the South Pacific gyre and within the mixed layer of the Melanesian basin (Figures 5, 6, Supplementary Information 3), even though the South Pacific gyre is largely isolated from the Lau basin waters by the Tonga ridge, as seen in DFe and diazotroph abundance (Lory et al., 2022; Tilliette et al., 2022; Figure 4A). Assuming that the DOM produced in the Lau basin has specific Fe-binding ligand properties and it disperses into the Melanesian basin and the South Pacific gyre, it would have to be decoupled from diazotrophic activity and DFe dispersion. This would be in line with previous studies suggesting faster turnover of DFe than Fe-binding ligands, the latter having a much longer residence time (Gerringa et al., 2015; Tagliabue et al., 2023). However, higher resolution is needed to better constrain the spreading of the DOM thought to originate from the recycling of the diazotroph that are thriving in the Lau basin (Bonnet et al., 2023a).

L_{Fe} production for trace metal detoxification

A strong production of L_{Fe} was observed in the surface waters at LD10-T3, related to a hydrothermal input of DFe (Tilliette et al., 2022; Figures 4A, 5B; Table 2). Interestingly, the L_{Fe} enhancement was only detected at the maximum DFe value and did not follow the linear dilution pattern showed by DFe with distance from the hydrothermal vent, suggesting that the amount of hydrothermally-sourced trace metals reached a concentration threshold triggering the production of Fe-binding ligands. The DFe enhancement did not reach toxic levels, therefore, the Febinding ligands detected could have been produced by the local plankton communities, including abundant diazotrophs, to neutralize toxic elements and not be Fe-specific/specifically produced to bind Fe. While HS can bind a range of metals, L_{FeHS} and HS-like FDOM did not increase at LD10-T3 (Table 2), and siderophores alone cannot be responsible for the increase in LFe and DFe since they are usually found at pM concentrations and have higher log $K_{Fe'L}^{cond}$ (Mawji et al., 2008; Bundy et al., 2018; Boiteau et al., 2019). On the other hand, bacterial production of EPS by species collected near hydrothermal vents has been reported as a

TABLE 2 Values in surface samples (depth < 20 m) with distance from the hydrothermal site.

	LD10-T1 – 15 km	LD10-T2 – 8 km	LD10-T3 – 2 km
DFe (nM)	0.38	1.52	3.20
L _{Fe} (nMeqFe)	4.2 ± 0.6	4.7 ± 0.9	8.8 ± 0.7
$\logK^{cond}_{Fe\}L}$	11.8 ± 0.2	11.4 ± 0.3	11.9 ± 0.2
L _{FeHS} (nMeqFe)	1.2	2.2	1.1
HS-like FDOM (QSU)	1.8	2.1	1.3

mechanism for slowing the diffusion of toxic elements into the organism and participating in the scavenging of toxic trace metals (Rougeaux et al., 1996; Nichols et al., 2005; Moppert et al., 2009; Deschatre et al., 2013). Production of EPS showing Fe binding properties was also observed for the nitrogen fixer Crocosphaera watsonii collected during the TONGA cruise (C. Lory, unpublished). Our results call for further consideration of the potential implication of EPS for regulating the distribution of the hydrothermally-sourced Fe and other toxic elements in the surface waters of the WTSP Ocean. One must also keep in mind the possible implication of small inorganic forms of Fe to explain the short-distance transport of the DFe enhancement, as highlighted by Dulaquais et al. (2023) and Tagliabue et al. (2023), as well as enhanced scavenging and precipitation in hydrothermallyinfluenced waters, which may result in ligand unsaturation (Gerringa et al., 2015; Thuroczy et al., 2011).

Both the Melanesian basin and the South Pacific gyre show strong L_{Fe} production in several deep-water samples (Figures 5A, B). On the eastern side of the Tonga ridge, L_{Fe} production coincides with relatively high log $K_{Fe'L}^{cond}$ (Figure 5), high HS-like FDOM (Figure 4) and high tryptophan-like FDOM (Tedetti et al., in prep), while no change in L_{FeHS} nor DFe is apparent (Figure 4). In the Melanesian basin (particularly at station 2), increased L_{Fe} coincides with relatively low log $K_{Fe'L}^{cond}$ (Figure 5), high HS-like FDOM (Figure 4) and high thiol-like compounds (Portlock et al., this issue²) and increased L_{FeHS} and DFe (Figure 4). The increased LFe, thiol-like compounds and tryptophan-like FDOM could be related to the toxicity of hydrothermally sourced metals. Indeed, the production of thiol-like compounds was recently associated with the detoxification of hydrothermal fluid in a ship-board incubation experiment performed with WTSP waters (Tilliette et al., 2023). Regarding the tryptophan-like signal, abiotic processes can lead to the production of tryptophan in hydrothermal systems (Ménez et al., 2018), and benthic interaction could also be involved. More information is required to disentangle the sources of Fe-binding ligands and tryptophan-like FDOM at this location, and efforts should focus on identifying the metals preferentially bound by the DOM associated with hydrothermal activity.

Humic contribution to DFe stabilization and to the Fe-binding ligand pool

Previous studies have also found L₂ ligands to dominate the water column in the South Tropical Pacific (e.g., Buck et al., 2018; Cabanes et al., 2020). The L₂ ligand class is often considered to be humic-like (Gledhill and Buck, 2012; Hassler et al., 2017), and our averaged log $K_{Fe'L}^{cond}$ of 11.6 ± 0.4 corresponds to log $K_{Fe'L}^{cond}$ values reported for Suwannee River humic-acid standards (SRHA; 11.1-11.6; Laglera and van den Berg, 2009; Abualhaija and van den Berg, 2014; Abualhaija et al., 2015). While there was no correlation of Fe-

² Portlock, G., Tilliette, C., Bonnet, S., Guieu, C., Whitby, H., and Salaun, P. Distribution and behaviour of reduced sulphur substances in the oligotrophic and hydrothermal waters of the Western South Tropical Pacific.

binding ligands with any parameters considered in this study, DFe showed the strongest correlation with HS-like FDOM (Table 3). Both DFe and HS-like FDOM increased in deep waters, with DFe being possibly complexed by HS-like FDOM as Fe(II) and/or Fe (III). Indeed, HS-like FDOM have been reported in hydrothermal fluids and are potentially capable of stabilizing Fe(II) (Yang et al., 2012; Sarma et al., 2018). However, it is not possible to estimate the proportion of L_{Fe} represented by the HS-like FDOM; for this, fluorescence quenching experiments, consisting of titrating the fluorophore with DFe, could be considered in future studies (Chen et al., 1994; Ohno et al., 2008).

The fluorescent and electroactive components of HS are thought to overlap, but the electroactive fraction of humic material, L_{FeHS} which comprises around 5% of the DOC in deep waters (Laglera and van den Berg, 2009; Dulaquais et al., 2018; Fourrier et al., 2022; Dulaquais et al., 2023), is thought to represent the metal-binding fraction specifically and can be compared to ligand concentrations directly. LFeHS have been shown to compose a large fraction of the Fe-binding ligands supplied during remineralisation (Whitby et al. 2020a). DFe has been shown to correlate with L_{FeHS} in some oceanic regions (Dulaquais et al., 2018; Laglera et al., 2019; Whitby et al., 2020b), and with AOU and HSlike FDOM in the Arctic and North Pacific Oceans (Tani et al., 2003; Hioki et al., 2014; Yamashita et al., 2017; Cao et al., 2020). On the other hand, DFe, LFe, LFeHS and HS-like FDOM tend not to correlate in the Atlantic Ocean (Heller et al., 2013; Whitby et al., 2020b). The electroactive and fluorescent properties of HS have been thoroughly investigated at station 8 of the TONGA cruise (Fourrier et al., 2022), but in hydrothermal systems, the contribution of L_{FeHS} and HS-like FDOM to the Fe-binding ligand pool and DFe transport is currently unknown.

Here, L_{FeHS} represented 20 ± 13% of L_{Fe} in the mixed layer, and 8 ± 6% in deep waters (Table 1; error calculated with L_{FeHS} standard deviation); therefore, while the average log $K_{Fe'L}^{cond}$ across the samples is consistent with humic complexation, and DFe correlates well with humic-like FDOM, electroactive L_{FeHS} specifically do not dominate the Fe-binding ligand pool. While this may seem unexpected, this %

contribution to L_{Fe} is similar to previous studies in the Pacific (2-51%; Cabanes et al., 2020; Whitby et al., 2020b), which tends to have a lower contribution of L_{FeHS} to the total ligand pool than in other ocean basins [e.g. L_{FeHS} comprise ~20-60% of the ligand pool in the Atlantic and Mediterranean (Dulaquais et al., 2018; Whitby et al., 2020b) and almost all of the ligand pool in parts of the Arctic Ocean (Slagter et al., 2017)]. While a large fraction of the ligand pool is not associated with electroactive HS, the L_{FeHS} that are present play an important role in Fe complexation, actively complexing around 30% of the DFe in these samples (Dulaquais et al., 2023).

Comparing the carbon content of the FA standard to DOC concentrations, L_{FeHS} represented 4.6 ± 2.6% of the DOC in the mixed layer (n = 17), and $2.2 \pm 1.7\%$ in deep waters (n = 60), similar to previously reported ranges (2 to 5%; Laglera and van den Berg, 2009; Dulaquais et al., 2018). While other, non-humic ligand groups may be contributing to the high L_{Fe} , the low correlation of L_{FeHS} with DFe, DOC, and AOU could be an indication of intense in-situ production of non-electroactive HS, corroborated by the low negative correlation between L_{FeHS} and HS-like FDOM (Table 3), characteristic of microbial respiration (Fourrier et al., 2022). Other metals can compete with Fe for L_{FeHS} complexation (e.g., Yang and van den Berg, 2009; Abualhaija and van den Berg, 2014; Whitby and van den Berg, 2015) and can also impact the HS-like FDOM (Zhao and Nelson, 2005). Non-specific ligands which contribute to L_{Fe} but can complex other metals could explain the unsaturation of the Febinding ligands produced in the Lau basin. The effect of competitive metals on Fe-binding ability of ligands, L_{FeHS} and HS-like FDOM would need to be investigated to better understand their potential impact on Fe speciation and cycling in hydrothermal environments.

Suggestions for future CLE-AdCSV studies

Toward tracing DOM characteristics with the reduction potential of the FeSA₂ complex

During AdCSV analyses, a variation in the peak potential of the reduction of the Fe-added ligand complex ($FeSA_2$) was

TABLE 3 Table of the Spearman rank correlation (ρ) of investigated parameters in this study.

	AOU (µM)	DOC (µM)	DFe (µM)	eL _{Fe} (nMeqFe)	$\log K^{cond}_{Fe''L}$	L _{FeHS} (nMeqFe)
DOC (µM)	-0.93 n = 161					
DFe (µM)	0.52 n = 238	-0.50 n = 151				
eL _{Fe} (nMeqFe)	0.07 n = 103	0.00 n = 66	0.09 n = 103			
$\logK^{cond}_{Fe]L}$	-0.18 n = 103	0.12 n = 66	-0.08 n = 103	-0.18 n = 103		
L _{FeHS} (nMeqFe)	-0.47 n = 120	0.46 n = 79	-0.23 n = 119	-0.05 n = 94	0.11 n = 94	
HS-like FDOM (QSU)	0.71 n = 130	-0.73 n = 124	0.45 n = 116	-0.03 n = 50	-0.17 n = 50	-0.26 n = 61

Blue corresponds to negative correlation, red to positive correlation, and white to absence of correlation. The intensity of the color is proportional to the strength of the correlation. Bold values correspond to p< 0.0001.

observed along the water column. The FeSA₂ peak potential ranged from -413 mV to -445 mV within the first 200 m (well-oxygenated mixed layer and most of the STUW/WSPCW), then decreased with depth to 1200 m where it reached a stable mean value of -473 mV down to the most abyssal waters (Figure 7). Also, in the mixed layer, the peak potential was significantly different between the Melanesian basin, the Lau basin, and the South Pacific gyre (Mann-Whitney test, p< 0.05; Supplementary Information 3). We hypothesize that the shift in reduction potential was due to an electroactive fraction of the DOM produced in the surface waters in the area. Indeed, the reduction of the metal complexed by the added ligand during AdCSV analysis happens at a specific potential following the Nernst equation (Nernst, 1889). The value of this potential is driven by i) the conditions of temperature, pH, and salinity, ii) the activity of the redox species, iii) the experimental conditions such as stripping scan rate, pulse time and pulse amplitude and iv) the adsorbed layer of organics accumulated on the electrode surface during the deposition step. Here, the temperature in the laboratory was controlled, the pH fixed by the addition of a buffer, and no correlation was found between the reduction potential and the salinity. The experimental measuring conditions were kept the same. The CLE-AdCSV method using SA takes advantage of a catalytic loop involving the reduction of the dissolved O₂ to hydrogen peroxide which immediately re-oxidize Fe(II) back into Fe(III), allowing for multiple reductions of the same ion and enhancing the signal (Laglera et al., 2016; Mahieu, 2023). The samples were equilibrated with air in the voltametric cell, ensuring a constant oxygen concentration (Mahieu, 2023) and a similar catalytic effect, regardless of the sample being analyzed. AdCSV analysis includes an accumulation step at a positive potential where the electroactive

complex FeSA₂ adsorbs at the electrode surface prior to its stripping. During this stripping, other compounds, organic and inorganic, may adsorb at the electrode surface and may impact the stripping peak potential. The influence of the DOM composition on the peak potential of FeSA₂ is yet to be addressed, but the ubiquity of the peak shift suggests similar electroactive DOM in the surface waters of all basins of the WTSP. It will be interesting to assess whether it is possible to identify specific components of DOM by the shift in the peak potential. Overall, this observation could highlight a novel way to identify changes in DOM composition, but it requires further investigation.

Analytical constraints for Fe-binding ligand detection by CLE-AdCSV

There are different CLE-AdCSV methodologies currently in use to evaluate the role of DOM complexation in DFe stabilization and transport. Although all are valuable, they are difficult to compare because of the specificities (e.g. detection window) of each method. Several basin scale investigations have presented Fe organic speciation in the WTSP, but they were obtained using different experimental procedure which might explain some of the observed differences. Using TAC as added ligand, Kondo et al. (2012) reported in most samples a low eL_{Fe} (< 1 nMeqFe) in the L₁ ligand class. In contrast, other studies in the Pacific Ocean using the added ligand SA showed ubiquitous, large eL_{Fe} (> 1 nMeqFe) falling in the L₂ class (Buck et al., 2018; Cabanes et al., 2020; this study). The difference can be explained by the analytical conditions with a higher detection window ($\alpha_{Fe'AL}$) for TAC than for SA, meaning that titrations with TAC focus on fewer Fe-binding ligands of



higher log $K_{Fe'L}^{cond}$ than the titrations with SA. The difference in detection window could lead to the underestimation of the HS-like contribution to the ligand pool because of the absence of competition with TAC (Laglera et al., 2011).

The analyses presented in this study were all performed in the dissolved fraction, without discrimination of the soluble and colloidal size fraction. Metal-binding compounds in the soluble Fe fraction (< 0.02 μ m) are assumed to be mostly organic, while the colloidal fraction (0.02 to 0.2 μ m, or 0.45 μ m here) may be composed of a wider mix of organic and inorganic compounds. It was suggested that this colloidal fraction of Fe was essential for DFe transport in the Subarctic Pacific (Kondo et al., 2021). Similarly, knowledge of the size fractionation of DFe, Fe-binding ligands, and HS partitioning in the Lau basin would greatly help better understand DFe speciation in this very dynamic region.

Accurate DFe concentrations are required in the calculation of L_{Fe} and log $K_{Fe/I}^{cond}$. Compared to mid-ocean ridges, volcanic arcs are characterized by higher carbon dioxide (CO_2) , sulfur dioxide (SO_2) and hydrogen sulfide (H₂S) enrichments, responsible for increased DFe removal by the formation of Fe-sulfide and Fe-oxyhydroxides along the dilution of hydrothermal plumes (e.g., Field and Sherrell, 2000). Such gas enrichment and DFe removal have been observed at several locations along the Tonga-Kermadec volcanic arc (Massoth et al., 2003; de Ronde et al., 2007; Resing et al., 2011; Neuholz et al., 2020; Kleint et al., 2022). The presence of such inorganic colloidal material can be problematic because DFe may be released under the acidic conditions used for DFe determination while being possibly inert species for CLE-AdCSV, which is performed at natural pH. The net effect is therefore an overestimation of the effective DFe concentration, leading to biased L_{Fe} and log $K_{Fe'L}^{cond}$. It is known for instance that the crystalline structure of some Fe-oxyhydroxides can sterically isolate Fe from metal-binding ligands (Kraemer et al., 2005; Dulaquais et al., 2023) while dissolving in acidic conditions (Liu and Millero, 2002). This effect has been observed in hydrothermal plumes by Kleint et al. (2016). They showed that up to 90% of DFe determined in acidified samples was non-labile to their added ligand in the buoyant plume, but this value was down to 15% in the non-buoyant plume. It is worth noting that the hydrothermal fluids investigated by Kleint et al. (2016) had DFe concentrations in the µM range, so the overestimation of DFe had a considerable impact on L_{Fe} and log $K_{Fe'L}^{cond}$ calculations. In the transect presented here, the most hydrothermally-influenced station was still 2km away from the vent and 90% of DFe concentrations were< 1 nM, which represents no more than 20% of the average L_{Fe} measured for our dataset. Thus, the error related to the potential overestimation of DFe is negligible for most of our samples. Nevertheless, the impact on CLE-AdCSV titrations of sulfides, oxyhydroxides and other aggregate materials filtered and classified as dissolved by the traditional operational definition is not known, and the competition between binding with the added ligand and adsorption/binding with inorganic and colloidal material is unknown.

The increasing number of Fe-binding ligand investigations in hydrothermal systems have encompassed a variety of techniques, each with advantages and disadvantages. Kleint et al. (2016) highlighted the issue related to the accurate definition of DFe concentration with SA within the hydrothermal plume in the Lau basin, and Wang et al. (2022) presented the correlation of the Fe isotopic ratios with the log $K_{Fe'L}^{cond}$ defined by reversed CLE-AdCSV using 1-nitroso-2-napthol (NN). Here, we present the distribution of the Fe-binding ligands in the Lau Basin and adjacent waters using SA. The next step forward in the characterization of hydrothermally derived Fe-binding ligands could be to directly compare the different available electroanalytical methods on the same water samples.

Conclusion

The TONGA transect (GPpr14) revealed intense production and accumulation of Fe-binding ligands predominantly of intermediate log $K_{Fe'L}^{cond}$ at all stations. The high L_{Fe} and eL_{Fe} observed in the area contrast with similar regions and may originate from the unusually high diazotroph activity known to occur in the Lau basin. Limited stabilization of the hydrothermallysourced DFe is most likely explained by competition from other metals and non-specificity of the ligands being produced, further supported by companion studies that report high levels of reduced sulfur substances and humic-like FDOM that coincide with our high L_{Fe} (in prep.). These non-specific ligands could be a detoxification response to hydrothermal waters, a hypothesis supported by ship-board incubation experiments (Tilliette et al., 2023).

Around 30% of DFe in these samples is within humic complexes (Dulaquais et al., 2023) and good agreement between FDOM and DFe suggest possible stabilization of a fraction of DFe as Fe(II) by HS-like FDOM. However, while the average log $K_{Fe'L}^{cond}$ was consistent with humic complexation across the ligand pool, ligands were in great excess of DFe and electroactive $L_{\rm FeHS}$ accounted for only 20 \pm 13% of L_{Fe} in the mixed layer and 8 \pm 6% in deep waters. Different trends in $L_{\mbox{\scriptsize FeHS}}$ and HS-like FDOM distributions confirm that HS-like FDOM and L_{FeHS} account for overlapping yet distinct fractions of the humic pool, and that the electroactive fraction of humics may underestimate the role of humic material in metal complexation. We recommend the use of quenching experiments in samples collected in the WTSP to evaluate the role of the HS-like FDOM in DFe transport and in Fe-binding ligand composition in future studies. Other ligands such as EPS and siderophores likely also contributed to Fe complexation, although the average log $K_{Fe'L}^{cond}$ values suggest strong ligands such as siderophores were also not a dominant component of the ligand pool. Our results may also be explained by competition between metals for non-specific ligands, and a multi-metal approach to examine the affinity of the binding sites of humic nature toward hydrothermally sourced trace metals would provide further insight. Ultimately, the partitioning of DFe, Fe-binding ligands, L_{FeHS} and HS-like FDOM should be investigated in both the soluble and colloidal fractions, and include the characterization of EPS, siderophores and other potential ligands.

The Tonga ridge effectively isolates the South Pacific gyre from the Lau basin, and yet there was an apparent dilution gradient from stronger to weaker ligands from the Lau basin into the South Pacific gyre and within the mixed layer of the Melanesian basin. This gradient was also seen in the electrochemical potential of the FeSA₂ peak in the mixed layer, which was highest in the Lau basin and decreased in adjacent surface waters of the Melanesian basin and South Pacific gyre. To our knowledge, this is the first time that a change in peak potential associated with changes in sample composition has been reported. We hypothesize that the peak shift could present a novel way to identify specific components of DOM, but this requires further investigation. In view of the existing studies harnessing a combination of forward and reverse titrations and different added ligands to determine Fe speciation in the WTSP, this region could present an ideal location for further intercomparison efforts to evaluate the ability of CLE-AdCSV in characterizing Fe-binding ligands of hydrothermal origin.

Data availability statement

The original contributions presented in the study are included in the article/Supplementary Material. Further inquiries can be directed to the corresponding author.

Author contributions

LM: Formal Analysis, Investigation, Writing – original draft. HW: Supervision, Writing – review & editing. GD: Formal Analysis, Writing – review & editing. CT: Formal Analysis, Writing – review & editing. CaG: Formal Analysis, Writing – review & editing. MT: Formal Analysis, Writing – review & editing. DL: Formal Analysis, Writing – review & editing. PF: Formal Analysis, Writing – review & editing. MB: Writing – review & editing. GS: Writing – review & editing. SB: Writing – review & editing. CéG: Writing – review & editing. PS: Supervision, Writing – review & editing.

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Conflict of interest

The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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Supplementary material

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fmars.2024.1304118/ full#supplementary-material

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