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Native Mass Spectrometry based Metabolomics Identifies Metal-binding Compounds

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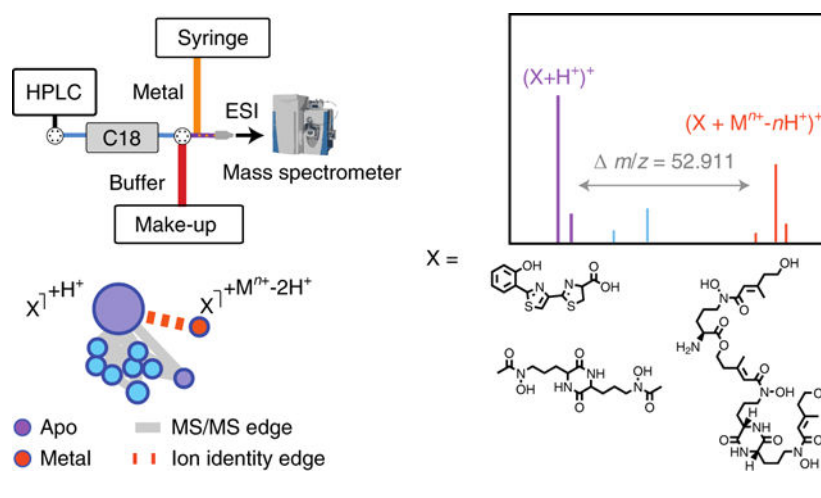
Author contributions: ATA, DP, RS, and PCD developed the concept. DP, JMG, IB, LA, ET, HZ, SPN, CCS, KPM, RJD, and MR cultured organisms and prepared samples. ATA, DP, and JMG ran mass spectrometry experiments. RS wrote code and provided feedback. ATA, DP analysed the data. ET, RJD, LIA, MR and PCD provided supervision and materials. ATA, DP, and PCD wrote the manuscript. All authors edited and approved the manuscript.

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Abstract

Although metals are essential for the molecular machineries of life, systematic methods for discovering metal-small molecule complexes from biological samples are limited. Here we describe a two-step native electrospray ionization-mass spectrometry method, in which post-column pH adjustment and metal-infusion are combined with ion identity molecular networking, a rule-based data analysis workflow. This method enabled the identification of metal-binding compounds in complex samples based on defined mass (m/z) offsets of ion species with the same chromatographic profiles. As this native electrospray metabolomics approach is suited for any liquid chromatography-mass spectrometry system to explore the binding of *any* metal, this method has the potential to become an essential strategy for elucidating metal-binding molecules in biology.

Graphical abstract



Introduction

Life, as we know it, cannot exist without metals. Metals are essential cofactors in critical biochemical reactions such as DNA replication and repair (iron and manganese), respiration (iron and copper), photosynthesis (iron and manganese), and biosynthesis of countless primary and secondary metabolites (e.g., iron, zinc, vanadium, molybdenum, magnesium, and calcium). One common strategy for metal acquisition in microorganisms is through the production and secretion of small molecule ionophores that bind metals and form noncovalent complexes. These complexes bind to specific ionophore receptors for their uptake and release of metal into the cytoplasm. Siderophores are high-affinity chelators of ferric iron (Fe^{3+}). They include ferrioxamines (i.e., desferrioxamines B and E)¹⁻⁴, catecholates (i.e., enterobactin)¹⁻³, and carboxylates (i.e., rhizoferrin and aerobactin)¹⁻⁴, while chalkophores such as methanobactins and SF2768 (a diisonitrile natural product) play the analogous role in binding cuprous copper (Cu^+)^{5,6}. Recently, several zincophores have also been recently elucidated⁷⁻¹⁰. In addition to small molecule ionophores that play a role in microbial metal transport and homeostasis, metals can also serve as cofactors essential for the function of other small molecules and proteins. Small molecule examples include

iron in heme, magnesium in chlorophyll, calcium and magnesium in metal-dependent antibiotics^{11,12}, and cobalt in the vitamin cobalamin (vitamin B12); protein examples range from DNA replication and repair enzymes to oxygen transport proteins and electron transfer enzymes.

Although metal-binding small molecules have various biological functions and many potential biomedical applications, it is currently challenging to find metal-binding compounds present in complex mixtures such as microbial culture extracts¹³, dissolved organic matter in seawater¹⁴, or fecal extracts¹⁵. Additionally, it is challenging to assess metal-binding preferences directly in these mixtures. Predictions of small molecule structures using genome mining strategies have improved tremendously in recent years^{16,17}; however, predicting the selectivity and affinity of metal coordination sites is still not possible, and it is even difficult to predict whether a small molecule contains a metal-binding site^{18,19}. This is because small molecule metal-binding sites are diverse and not conserved; as such, metalbinding must be experimentally established, often relying on methods such as inductively coupled plasma-mass spectrometry (ICP-MS)²⁰, atomic absorption spectroscopy (AAS)²¹, x-ray fluorescence spectroscopy (XRF)²², UV-visible (UV-vis) absorption spectroscopy, and nuclear magnetic resonance (NMR) spectroscopy, which are typically low-throughput, performing best on pure compounds. With each of these methods, it is difficult to understand which molecules form metal complexes within a biological matrix that contains a pool of candidate metal ions, especially when the identities of metal-binding species are not known ahead of time.

Mass spectrometry strategies, such as the use of characteristic metal isotope fingerprints, have been used to discover siderophores from complex samples. This initially facilitated the study of siderophores in seawater²³⁻²⁸ and was subsequently assisted by mass defect/Kendrick analysis²⁹, analysis of hyperfine splitting isotope patterns³⁰, and the development of software packages such as ChelomEx to enable siderophore screening strategies³¹⁻³³. Multimodal approaches have been employed combining matrix-assisted laser desorption/ionization-Fourier transform ion cyclotron resonance mass spectrometry imaging (MALDI-FT-ICR MSI)²⁷ with LC-ESI-MS/LC-ICP-MS^{19,34} based methods. While these methods have been instrumental in the discovery of several siderophores, we hypothesized that many metal-binding compounds (and specifically those binding metals other than iron) remain undiscovered. Our hypothesis is based on a repository scale analysis of the presence of apo (unbound) versus holo (bound) ionophores in public datasets hosted in the Global Natural Products Social (GNPS) Molecular Networking platform (SI Table 1a). To discover new small molecule-metal complexes, we envisioned developing a modular and straightforward non-targeted LC-tandem mass spectrometry (LC-MS/MS) based strategy that is broadly applicable to *any* metal and makes use of a native mass spectrometry approach, which has gained momentum in the field of structural biology and intact protein analysis^{35,36}. In combination with a novel computational tool - Ion Identity Molecular Networking (IIMN) that connects related compounds by correlation analysis, user-defined mass differences, and MS/MS similarity³⁷, this workflow enables screening for metal-binding compounds in complex biological samples. We call the experimental workflow *native electrospray metabolomics*. When combined with IIMN, this strategy facilitates finding metal-binding compounds that may be biologically relevant. A key feature that differentiates native

metabolomics from other LC-based methods is the ability to control pH without altering chromatography. SI Table 1b highlights the critical differences between current MS-based siderophore screening strategies and the native metabolomics approach; the novelty of the native metabolomics approach lies in its broad applicability to find biologically relevant metalcomplexes associated with *any* metal (e.g., Fe, Cu, Zn, Na, and Ca) in a wide range of pH conditions and sample complexities.

Native Metabolomics Concept.

Extraction, sample preparation, and chromatography conditions typically employed for non-targeted reversed-phase LC-MS/MS-based analysis of small molecules involve low pH, high percentages of organic solvent, and low metal concentrations, which disfavor metal complexation^{38,39,89}. For example, extraction of marine dissolved organic matter for non-targeted MS/MS analysis, when not explicitly designed for isolation of metal-bound organics, typically employs low pH to protonate carboxylic acids and increase SPE extraction efficiency¹⁴. Based on this, we hypothesized that many metal-binding compounds detected by reversed-phase (RP)-LC-MS/MS are predominantly observed in their protonated (apo) form rather than in a metal-bound form. Investigating experimental non-targeted MS/MS datasets in a repository-scale analysis of MS/MS spectra of proton- and metal-bound siderophores in the GNPS web platform^{40,41} and MetaboLights repository⁴² with the Mass Spectrometry Search Tool (MASST)⁴³ provides evidence for this assumption (SI Table 1a). When querying a list of known siderophores (SI Table 1a) against the GNPS-MassIVE and MetaboLights repository, 73% of files contained the proton-bound form exclusively, with only 27% of files containing the metal-bound form. Notably, even when the metal-bound species is observed, it can be challenging to match it to the corresponding proton-bound species, because proton- and metal-bound species typically elute at different retention times⁴⁴⁻⁴⁶ and exhibit significantly different MS/MS fragmentation behavior³⁷. Therefore, searching for a given (metal-specific) mass (m/z) offset to connect proton- with metal-bound species can be non-trivial with current tools.

Given these inherent limitations of contemporary RP-LC-MS/MS methods, we hypothesized that adjusting pH to physiological values - an approach often applied in intact protein mass spectrometry that has been termed native spray²⁸⁻³⁰ - would help to detect compounds with metal binding affinities in metabolomics experiments. Native electrospray metabolomics facilitates the ability to control pH without altering chromatography, which is not possible with other LC-based methods that aim to identify metal-binding molecules. Preliminary direct injection experiments at acidic, neutral, and basic pH showed higher ratios of metal-adduct to proton-adduct at higher pH values, which is consistent with literature^{38,39,89,90,95}. Thus, we developed a two-step native electrospray ionization (ESI) LC-MS/MS workflow with online post-column pH adjustment is coupled to metal infusion to directly detect candidate small molecules that bind metals. While ion-addition has been utilized for various MS applications⁴⁷⁻⁵⁰, the use of post-LC metal-addition for finding biologically relevant metal-binding molecules and integration of this into a widely used web-based analysis ecosystem is unique.

Our first-generation workflow utilizes a double-barrel syringe pump for infusion of ammonium hydroxide solution and subsequent injection of one or multiple metal salt(s) (SI Figure 1a). As metal-binding kinetics are fast (on the order of milliseconds^{51–54}), we hypothesized that our set-up would allow for sufficient time to establish metal-ligand complexes after chromatography, pH adjustment, and metal infusion, as it takes c.a. 0.6 seconds to reach the entrance of the electrospray inlet after the post-column metal addition. The second-generation of our workflow makes use of a secondary high-performance liquid chromatography (HPLC) pump that provides an ammonium acetate buffered make-up flow and the syringe pump delivers metal solution (Figure 1a, SI Figure 1b). This workflow offers the advantage of a more consistent (and higher percentage aqueous) solvent composition across the entire LC gradient. Based on this, the second-generation workflow was used to assess the effect of pH adjustment on metal-binding. Mixtures of metal salts - FeCl₃, Cu(OAc)₂·(H₂O), Mn(OAc)₂·4(H₂O), Zn(OAc)₂·2(H₂O), Co(OAc)₂·4(H₂O), and NiCl₂ - were infused at two concentrations and at three pH values, namely pH 2, 5, and 9. The ratio of the integrated peak area of Fe-adducts, for example, to the integrated peak area of Hadducts increased at higher pH values (SI Figure 2). Based on this observation, we chose to carry out subsequent studies at pH values greater than 5, and we aimed to operate at the relevant pH value for the biological systems under investigation. While the post-LC pH was verified, ESI-induced changes in pH can occur in the ESI source^{91,92,95}.

Using both first and second-generation set-ups, proton- and metal-bound complexes are observed at the same retention time with the same peak shape; however, one of the critical challenges in developing this method was how to identify metal-binding complexes within large datasets and/or complex mixtures that typically contain thousands of features. The rule-based informatic workflow IIMN was used to overcome this limitation (Figure 1b) by applying MS data processing in MZmine^{41,42} linked to GNPS^{43,44}. Using ion identity networking (IIN) in MZmine 2, LC-MS features, defined here as chromatographic peaks with a specific *m/z*, are grouped based on their retention time and chromatographic feature shape correlation and identified as specific ion types of the same analyte molecule akin to the way it is accomplished by CAMERA⁵⁵ or RAMClust⁵⁶. In this process, all metal-binding small molecules can be identified by linking the metal-bound ions to other adducts and in-source fragments of the apo structures. The connected ion features are then further linked within the molecular network created based on MS/MS similarity. The use of molecular networking based on MS/MS similarity facilitates dereplication of potential metal-binding compounds, i.e., a known siderophore that shares an MS/MS similarity edge with an unknown metal-binding node can be used to dereplicate the unknown metal-binding derivative⁵⁷. Given this, we expect molecular networking will enable dereplication of many metal-binding molecules, as microbial gene clusters can make a series of small molecules in addition to their characterized small molecules product due to multiple substrate pools and promiscuity of the biosynthesis⁵⁸. As such, IIMN is performed by (1) feature finding in MZmine 2^{41,42}, (2) grouping of co-occurring features using Pearson correlation of their corresponding intensity profiles, (3) filtering results based on user-defined ion rules and *m/z* error tolerances, and (4) connection of similar MS/MS spectra in GNPS through the cosine similarity metric. This enables the visualization of apo versus metal-bound compounds, even

when the corresponding ion adducts have different gas-phase fragmentation behaviour in collision induced dissociation (SI Figure 3).

After performing feature finding that considers grouping based on feature shapes correlation in MZmine 2^{41,42}, all grouped features are searched in pairs against a user-defined ion identity library specified by lists of adducts, in-source fragments, and multimers. This approach tests all possible combinations of ion identities that point to the same neutral mass of an ionophore (M). All metal adducts and ion species combinations are searched, which can be crucial if no protonated feature is observed. Consequently, as proton- and metal-bound species have the same retention time (RT) and feature shape in this workflow, IIN enables discovering novel metal-binding molecules by any combination of singly or multiply charged ion species and therefore lifts any restrictions to the most common ion types. This computational workflow facilitates the search for any characteristic m/z delta, defined by the user, to enable finding molecules that form complexes with iron (3+ ion, m/z delta=52.9115), in addition to other metals such as zinc (2+ ion, 62.9208), copper (2+ ion, 61.9212), cobalt (2+ ion, 57.9248), sodium (1+, 22.9892), potassium (1+, 38.9632), calcium (2+, 39.9615), aluminium (3+, 24.9653), etc. Although alkali metals and alkaline earth metals (such as Na⁺, K⁺, and Ca²⁺) are generally considered nuisances in mass spectrometry, these adduction events may also have biological relevance^{59–61} and can serve as a starting point for hypothesis generation.

Results and Discussion

Metal-binding specificity.

To evaluate our native metabolomics workflow, we prepared standard mixtures of commercially available, well-characterized⁹⁴ siderophores, namely yersiniabactin (**1**), vibriobactin (**2**), enterobactin (**3**), ferrichrome (**7**), and rhodotorulic acid (**6**). The standards were separated by HPLC then subjected to post-LC pH adjustment and excess (millimolar) infusion of iron chloride (FeCl₃) via the first-generation set-up. We observed the Fe³⁺-adduct of each siderophore only after iron-infusion (Figure 2a). For initial screening, iron concentrations between 3–5 mM were infused, corresponding to final concentrations of 30–50 μM (SI Table 2) at the electrospray source, a c.a. 10-fold molar excess over siderophore concentration at the electrospray source (SI Table 2). To systematically assess whether iron-binding under these conditions is dependent on concentration and to evaluate the native electrospray metabolomics method under sub-stoichiometric (and physiological) concentrations, iron was titrated into a mixture of standards containing compounds (**3**), (**7**), (**6**), in addition to desferrioxamines E (DFE) and B (DFB). This resulted in a dose-dependent response in the ratio of integrated peak areas of Fe-adduct to H-adduct (SI Figure 4). Sub-stoichiometric iron concentrations at the ESI source (SI Table 2, based on the 51x dilution) resulted in an increase in the ratio of integrated peak areas of Fe-adducts to H-adducts (over background) from which we infer relative binding affinity. This increase is compound-dependent and likely also dependent on solvent composition, particularly the proportion of organic solvent (SI Figure 4). For this reason, we developed the second-generation set-up with an additional make-up flow pump that reduces the variation in solvent composition across the gradient and lowers the organic solvent concentration by ~50%. At

lower metal concentrations, regular cleaning of the source and spray shield (in addition to other mass spectrometer parts) becomes increasingly important as metal salts are infused. Failure to clean the instrument can result in background signals in control (non-infusion) samples, especially when compounds bind with high affinities.

While we observed that metal-binding response is concentration-dependent, we did not observe a strong correlation between relative binding affinity and solution-phase literature K_d values⁶⁶. We speculate this may be due to potential differences in ionization efficiencies for proton- versus metal-bound ionophores, compound-dependent reduction processes that can occur during electrospray ionization, or differences in ionization efficiencies between compounds⁹³. Similar caveats have also been widely described in the context of native protein analysis by mass spectrometry³⁵. Initial proof-of-concept experiments focused on siderophores (versus ionophores for non-iron metals) due to the prevalence of commercially available, well-characterized⁹⁴ iron-binding standards. To assess whether observed iron-binding was specific for known siderophores or whether non-specific adducts form after iron-infusion, we added the siderophore ferrichrome to a matrix of competing molecules without described iron-binding properties, and we found iron-adduction to be specific. This mixture was analysed using the first-generation native electrospray metabolomics workflow in the absence and presence of excess (millimolar) iron. In this experiment, ferrichrome is the only compound connected with an IIMN-edge corresponding to iron-binding and no other molecules were observed to bind iron (Figure 2b).

To compare the effectiveness of native electrospray as an approach to identify small molecule-metal interactions, we compared native electrospray metabolomics to other LCMS-based siderophore screening strategies^{31–33} (SI Table 1b). While these strategies can reproducibly identify iron-binding of strong siderophores, they are less suited to detect the metal adduct of compounds with low metal affinities, missed metal adducts in complex mixtures, and overlooked adducts of non-iron metals (SI Figure 5), as highlighted in the following sections of the text.

Metal-binding selectivity.

In addition to testing specificity, i.e., the ability of native metabolomics to detect only true metal-binding molecules - we next tested whether native metabolomics could be used to understand the selectivity of metal-binding. We assessed this by infusing a mixture of metals at physiological levels (μM), including iron, copper, cobalt, nickel, zinc, and manganese salts, and found that metal selectivity differed among the tested standards tested (SI Figure 2). For example, only in the case of yersiniabactin did copper-binding strongly compete with iron-binding⁶⁷. For metal selectivity, relative binding affinities were consistent with literature that suggests yersiniabactin can bind copper more strongly than iron under certain conditions⁶⁷. Both ferrioxamines showed a higher preference for iron than other infused metal salts; interestingly, the native metabolomics method can differentiate binding preferences of two very similar compounds, namely desferrioxamines E and B. The fact that desferrioxamine B exhibits copper-binding while desferrioxamine E does not is also consistent with literature reports⁶⁸.

Observing Siderophores in Culture Extracts.

To further assess selectivity within biological samples, we applied this workflow on culture extracts from *Glutamicibacter arilaitensis* JB182⁶⁹. This microbe was previously isolated from cheese and grown here in liquid cheese media. Cheese is an iron-depleted environment, as it contains lactoferrin, an iron-binding protein that sequesters free iron⁷⁰. Siderophore production was predicted based on genomic data mining using antiSMASH 4.0⁷¹, which identified biosynthetic gene clusters of ferrioxamine B and fuscachelin with 50% and 44% similarity, respectively. Using the native metabolomics workflow developed here, both apo-desferrioxamine E (DFE) and the Fe³⁺-bound ferrioxamine E were observed from culture extracts and were connected by an iron-binding IIMN edge; DFE was annotated by a spectral match from molecular networking in GNPS^{43,44}. DFE was the only iron-binding connection observed using IIMN (Figure 3a), which illustrates the specificity of iron-binding during the post-LC infusion; none of the other molecules detected in this complex biological sample were iron-bound (Figure 3b–e).

Based on the promiscuous nature of metal-binding by yersiniabactin in both metal-selectivity experiments (Metal-binding selectivity) and in the literature^{67,72}, we were interested in further investigating metal-binding properties of this compound directly in culture extracts of *Escherichia coli* strain Nissle 1917 (*E. coli* Nissle, serotype O6:K5:H1) grown in M9 glucose minimal medium (SI Figure 5). Using the native metabolomics strategy, we ran supernatant extracts with post-LC pH adjustment to ~7 (based on *E. coli* cytosol pH^{73,74}) in the absence and presence of (excess) iron-infusion and could identify a series of iron-binding molecules using IIMN (Figure 4a and SI Figure 6). From those compounds we could annotate yersiniabactin, aerobactin, and HPTzTn-COOH^{75,89,90}, a truncated derivative of yersiniabactin, and each corresponding iron-complex (Figure 4b–d). IIMN reveals several additional yersiniabactin (Figure 4b and c) and aerobactin (Figure 4d) derivatives that also bind iron, suggesting at least fifteen additional siderophores are present in this culture extract. SIRIUS 4.0⁷⁶ was used to predict molecular formulas of these compounds with manual inspection of retention times and MS/MS were manually inspected to exclude in-source fragments and propose putative structures (SI Figure 7). This example illustrates the power of IIMN in the dereplication of siderophore derivatives that also bind iron as the molecules that co-network are structurally related to the siderophores produced by this organism. This indicates an essential aspect of molecular networking – its ability to identify structural similarity, where modifications are often associated with biosynthesis or metabolism⁵⁸. An extension by C₂H₂, for example, cannot be an MS artifact, while an oxidation could be if the retention times are identical – IIMN would uncover this. Based on this, the predicted molecular formulas are tabulated in Table 1a (with retention times and possible in-source fragments tabulated in SI Table 3a).

In addition to investigating iron-binding at physiological pH ~7, we also sought to identify zinc-binding molecules in these culture extracts, as zinc-acquisition mechanisms of *E. coli* Nissle have not been fully elucidated. Evaluating zinc-binding molecules using native metabolomics yielded the novel finding that yersiniabactin along with a number of its derivatives, including HPTzTn-COOH, also bind zinc. Based on this result, we discovered that yersiniabactin facilitates *E. coli* Nissle's ability to flourish in an inflamed gut by

evading zinc sequestration by the antimicrobial protein calprotectin⁸⁸. To confirm the zinc-binding observed by native electrospray metabolomics, we monitored zinc-addition into pure yersiniabactin⁸⁸ and into HPTzTn-COOH (SI Figure 8) by 1D ¹H NMR. Changes in chemical shifts and peak broadening arise from zinc coordination, which changes the electronic environment of these protons; therefore, ¹H NMR studies serve as orthogonal confirmation of zinc-binding.

Given that these culture extracts contain compounds that bind various metals, these samples were valuable for investigating metal-competition at physiologically relevant metal concentrations (μM). The pH and ionic strength of the *E. coli* cytosol have been well characterized, with the intracellular pH ~ 7.5 ^{73,74}. Labile Fe^{2+} and Mn^{2+} concentrations have been estimated to be $10 \mu\text{M}$ ^{77,78}, while labile Cu^+ and Zn^{2+} have been estimated to be $10^{-15} \mu\text{M}$ ⁶¹ and $10^{-9} \mu\text{M}$ ⁶¹, respectively. While alkali and alkaline earth metals (including Na^+ , Ca^+ , and K^+) are present in mM amounts, mM concentrations result in significant ion suppression during mass spectrometry, so only transition metals were infused at the levels listed above.

We found that metal adducts were still observed when physiologically relevant levels were infused. However, the increase in feature intensity of metal adduct-features is significantly less than the observed increase with excess iron infusion (SI Figure 9a–b). Interestingly, yersiniabactin exhibits both iron and copper-binding, even when only trace levels of copper are infused (SI Figure 9b–d). Additionally, we observe that metal reduction can occur during ESI⁷⁹ (Gianelli et al. 2001), as we detected small amounts of $[\text{M}+\text{Fe}^{2+}]^{2+}$ and $[\text{M}+\text{Fe}^{2+}-\text{H}^+]^+$ adducts in addition to the expected $[\text{M}+\text{Fe}^{3+}-2\text{H}^+]^+$ adducts (SI Figure 10).

After illustrating the ability of native metabolomics to elucidate metal-binding molecules present in *E. coli* Nissle extracts specifically, we next tested this method on increasingly complex samples. We applied our method to fungal extracts from the wine fungus *Eutypa lata*. The *Eutypa lata* genome⁸⁰ is approximately ten times the size of the *E. coli* Nissle genome⁸¹. Moreover, the published *Eutypa lata* strain UCREL1 contains a predicted domain structure that matches the coprogen synthetase SSM2 from *Magnaporthe oryzae*^{82,83}. Applying the post-LC pH adjustment and Fe^{3+} -infusion mass spectrometry workflow enabled the detection of apo-coprogen B and apo-hydroxydimethyl coprogen B in addition to the iron-bound forms of both siderophores. Apo and Fe^{3+} -bound forms were each connected through an Fe^{3+} -binding IIMN edge (Figure 4e). We additionally identified several structurally related derivatives in the same molecular family that bind iron (Figure 4f). As in the *E. coli* Nissle example, we used SIRIUS 4.0⁷⁶ to elucidate molecular formulas and have putatively annotated six structurally related novel siderophores, again underscoring the potential of this method for the discovery of new analogues^{58,84}. These siderophores are tabulated in Table 1b (with retention times tabulated in SI Table 3b), and MS/MS spectra are annotated (SI Figure 11).

Observing Metal-Binding Molecules in Complex Environmental Samples.

Finally, we applied the native metabolomics workflow to environmental samples to test whether this workflow could identify metal-binding compounds in ultracomplex samples. We analysed solid-phase extracted surface ocean samples collected in the California Current

Ecosystem during a phytoplankton bloom in June 2017 with our native metabolomics approach. Dissolved organic matter (DOM) from surface seawater is among the most complex samples analysed by UHPLC-MS/MS in our labs (more complex than fecal and on par with crude oil samples). Metals were infused post-column after pH adjustment to ~8 (average pH of the ocean water in California Current Ecosystem⁸⁵). The native metabolomics method identifies domoic acid (DA) as a copper-binding molecule, which is consistent with the literature^{86,87}. Furthermore, we confirmed the identity of DA as Metabolomics Standards Initiative level 1 annotation⁹⁶ with an authentic standard. Interestingly, IIMN indicated that DA binds Cu^{2+} as a dimer (Figure 5), perhaps coordinating copper in a similar binding configuration to ethylenediaminetetraacetic acid (EDTA)⁹¹, which also contains four carboxylic acids and two amino groups that form a hexadentate complex. This example illustrates how native metabolomics can be applied to environmental samples, identifying metal-binding compounds in ultra-complex matrices.

Conclusion

The native metabolomics workflow described here can identify known and novel ionophores from complex samples, such as culture extracts and DOM. Currently, the screening for metal-binding compounds is difficult to perform on complex mixtures, as the majority of techniques used to assess metal-binding (including NMR, UV-vis, EPR) are usually performed on purified compounds. ICP-MS, ICP-OES and AAS can be used to assess bulk metal content of complex samples. However, they cannot provide molecular detail about which compounds present in a complex sample actually bind metals. Split-flow LC approaches offer innovative strategies for simultaneously analysing metal content and structures but require expensive and specialized equipment and customized set-ups²⁸. Based on the minimal technical adjustments, we foresee the routine implementation of this native metal metabolomic approach into non-targeted LC-MS/MS metabolomics workflows for elucidating metal-binding small molecules in biology. Additionally, this workflow can be adapted to give detailed insights into metal-binding selectivity. Based on our results, we suspect that molecules traditionally described as iron-binding siderophores may also be responsible for binding and uptake of other metals, as we recently discovered in the case of yersiniabactin and its derivative HPTzTn-COOH; yersiniabactin has been shown to play a biological role in microbial zinc-acquisition⁸⁸. The method outlined here is uniquely set up to identify novel metal-binding molecules in complex samples. We anticipate this will significantly improve our understanding of the role of metals across diverse fields ranging from human health to agriculture to the environment.

Online Methods

Sample Preparation.

Commercial siderophore standards were purchased from EMC Microcollections (<https://www.microcollections.de/product-services/siderophores.html>). Domoic acid standards were purchased from Sigma Aldrich. Standards were diluted in either water or methanol to a concentration of 1 mM and diluted further from the stock solution.

Direct Injection MS data acquisition.

For MS analysis, 5 μL were directly infused into a Q-Exactive orbitrap mass spectrometer via flow injections through a Vanquish UHPLC system (Thermo Fisher Scientific, Bremen, Germany). A flow rate between 0.2 mL/min and 0.4 mL/min was used. MS¹ data acquisition was performed in positive mode. Electrospray ionization (ESI) parameters were set to 53 L/min sheath gas flow, 14 L/min auxiliary gas flow, 0 L/min sweep gas flow, and 400°C auxiliary gas temperature. The spray voltage was set to 3.5 kV and the inlet capillary to 320°C. 50 V S-lens level was applied. MS scan range was set to 150–1500 m/z with a resolution at m/z 200 ($R_{m/z\ 200}$) of 35,000 with one micro-scan. The maximum ion injection time was set to 100 ms with an automated gain control (AGC) target of 1.0E6.

Standard-flow LC-MS/MS data acquisition.

2–5 μL were injected into a Vanquish HPLC system coupled to a Q-Exactive orbitrap mass spectrometer (Thermo Fisher Scientific, Bremen, Germany). A C18 porous core column (Kinetex C18, 50 \times 2 mm, 1.8 μm particle size, 100 Å pore size, Phenomenex, Torrance, USA) was used for chromatography. For gradient elution, a high-pressure binary gradient system was used. The mobile phase consisted of solvent A H₂O + 0.1 % formic acid (FA) and solvent B acetonitrile (ACN) + 0.1 % FA, unless otherwise specified. The flow rate was set to 0.5 mL/min, unless otherwise specified. One of the following two methods (*method 1* or *method 2*) was used for analysis. After injection, the samples were eluted with one of the following linear gradients: 0–0.5 min, 5% B, 0.5–5 min 5–99% B, followed by a 2 min washout phase at 99% B and a 3 min re-equilibration phase at 5% B (*method 1*) or 0–0.5 min, 5 % B, 0.5–9 min 5–100% B, followed by a 2 min washout phase at 99% B and a 3 min re-equilibration phase at 5% B (*method 2*). Data-dependent acquisition (DDA) of MS/MS spectra was performed in positive mode. ESI parameters were set to 53 L/min sheath gas flow, 14 L/min auxiliary gas flow, 0 L/min sweep gas flow, and 400°C auxiliary gas temperature, the spray voltage was set to 3.5 kV and the inlet capillary to 320°C and 50 V S-lens level was applied (*method 1*); or 60 L/min sheath gas flow, 20 L/min auxiliary gas flow, 3 L/min sweep gas flow, and 300°C auxiliary gas temperature, the spray voltage was set to 3.5 kV and the inlet capillary to 380°C and 60 V Slens level was applied (*method 2*). MS scan range was set to 150–1500 m/z with a resolution at m/z 200 ($R_{m/z\ 200}$) of 35,000 with one micro-scan. The maximum ion injection time was set to 100 ms with an automated gain control (AGC) target of 1.0E6. Up to 5 MS/MS spectra per MS¹ survey scan were recorded DDA mode with $R_{m/z\ 200}$ of 17,500 with one micro-scan. The maximum ion injection time for MS/MS scans was set to 100 ms with an AGC target of 1E6 ions (*method 1*) or 5E5 ions (*method 2*). The MS/MS precursor isolation window was set to m/z 1 (*method 1*) or 2 (*method 2*). The normalized collision energy was set to a stepwise increase from 20 to 30 to 40% with $z = 1$ as default charge state. MS/MS scans were triggered at the apex of chromatographic peaks within 2 to 15 s from their first occurrence. Dynamic precursor exclusion was set to 5 s (*method 1*) or 30 s (*method 2*). Ions with unassigned charge states were excluded from MS/MS acquisition as well as isotope peaks.

Micro-flow LC-MS/MS data acquisition.

For micro-flow HPLC-MS/MS analysis extracts and DOM samples were re-dissolved in methanol/water/ formic acid (80:19:1, Fisher Scientific, San Diego, USA) and 2–10 μL were injected into vanquish UHPLC system coupled to a Q-Exactive quadrupole orbitrap mass spectrometer (Thermo Fisher Scientific, Bremen, Germany) with an Agilent 1260 quaternary HPLC pump (Agilent, Santa Clara, USA) as a make-up pump. For reversed-phase chromatographic, a C18 core-shell microflow column (Kinetex C18, 150×1 mm, 1.8 μm particle size, 100 \AA pore size, Phenomenex, Torrance, USA) was used. The mobile phase consisted of solvent A ($\text{H}_2\text{O} + 0.1\%$ formic acid (FA)) and solvent B (acetonitrile (ACN) + 0.1 % FA). The flow rate was set to 100 $\mu\text{L}/\text{min}$. A linear gradient from 5–50 % B between 0–8 min and 50–99 % B between 8 and 10 min, followed by a 2 min washout phase at 99% B and a 6 min re-equilibration phase at 5 % B. Data-dependent acquisition (DDA) of MS/MS spectra was performed in positive mode. Make-up flow of ammonium acetate buffer (10 mM) was set to 100 $\mu\text{L}/\text{min}$ and infused post-column through a peak T-splitter. Metal-mixture (at the concentration specified in SI Methods) was infused post-column and post-make-up through a second T-splitter at a flow rate of 5 $\mu\text{L}/\text{min}$. ESI parameters were set to 40 arbitrary units (AU) sheath gas flow, auxiliary gas flow was set to 10 AU and sweep gas flow was set to 0 AU. Auxiliary gas temperature was set to 400 $^\circ\text{C}$. The spray voltage was set to 3.5 kV and the inlet capillary was heated to 320 $^\circ\text{C}$. S-lens level was set to 70 V applied. MS scan range was set to 2002000 m/z with a resolution at m/z 200 ($R_{m/z\ 200}$) of 70,000 with one micro-scan. The maximum ion injection time was set to 100 ms with automatic gain control (AGC) target of $5\text{E}5$. Up to two MS/MS spectra per duty cycle were acquired at $R_{m/z\ 200}$ 17,000 with one micro-scan. The maximum ion injection time for MS/MS scans was set to 100 ms with an AGC target of $5.0\text{E}5$ ions and a minimum 5% AGC. The MS/MS precursor isolation window was set to m/z 1. The normalized collision energy was stepped from 20 to 30 to 40% with $z = 1$ as the default charge state. MS/MS scans were triggered at the apex of chromatographic peaks within 2 to 15 s from their first occurrence. Dynamic precursor exclusion was set to 5 s. Ions with unassigned charge states were excluded from MS/MS acquisition as well as isotope peaks.

Data analysis.

Feature finding and ion identity networking were performed using an in-House modified version of MZmine2.37^{40–42}, corr.17.7 available at https://mzmine.github.io/iin_fbmn. Feature tables, MS/MS spectra files (mgf), and ion identity networking results were exported, uploaded to MassIVE, and submitted to GNPS for feature-based molecular networking analysis. Details of the analysis are provided in the Supplementary Methods.

Data and software availability.

The data supporting the findings of this study are available within the paper and its Supplementary Information. All mass spectrometry .raw and centroided .mzXML or .mzML files are publicly available in the mass spectrometry interactive virtual environment (MassIVE) under massive.ucsd.edu with project identifier [MSV000084237](https://massive.ucsd.edu/MSV000084237), [MSV000085669](https://massive.ucsd.edu/MSV000085669), [MSV000085206](https://massive.ucsd.edu/MSV000085206) (Standards), [MSV000084289](https://massive.ucsd.edu/MSV000084289) (Cheese siderophores), [MSV000082999](https://massive.ucsd.edu/MSV000082999) and [MSV000084030](https://massive.ucsd.edu/MSV000084030) (Fungal siderophores),

MSV000083387 (*E.coli* Nissle siderophores), MSV000085554 (California Current Ecosystem phytoplankton bloom samples), MSV000086744 (Chelomics vs Native Metabolomics Comparison). Ion Identity Molecular Networks (IIMN) can be accessed through gnps.ucsd.edu under direct links: <https://gnps.ucsd.edu/ProteoSAFe/status.jsp?task=79d0f380b4814ff9a720836c5570036f>, <https://gnps.ucsd.edu/ProteoSAFe/status.jsp?task=ad4b2665dfb744d09a9d2445f1213720>, <https://gnps.ucsd.edu/ProteoSAFe/status.jsp?task=5459d22126e843a3a1449f8362cd267f>, <http://gnps.ucsd.edu/ProteoSAFe/status.jsp?task=1c3e79f0ab984386bd468e2d163281e0>, <https://gnps.ucsd.edu/ProteoSAFe/index.jsp?task=196a29a94c2f4c788e204b9934ea4d9b>, <http://gnps.ucsd.edu/ProteoSAFe/status.jsp?task=256ba734f4334c1c90f65ffbd9141d0e>, <https://gnps.ucsd.edu/ProteoSAFe/status.jsp?task=042939579da64e029a8b5caef8f7f2a8>.

Code availability statement

The modified version of MZmine 2 (2.37, corr.17.7)^{40–42} can be found here https://mzmine.github.io/iin_fbm. GNPS^{43,44} can be accessed here <https://gnps.ucsd.edu/ProteoSAFe/static/gnps-splash.jsp> Cytoscape⁵⁹ version 3.7.1 can be accessed here <https://cytoscape.org/>.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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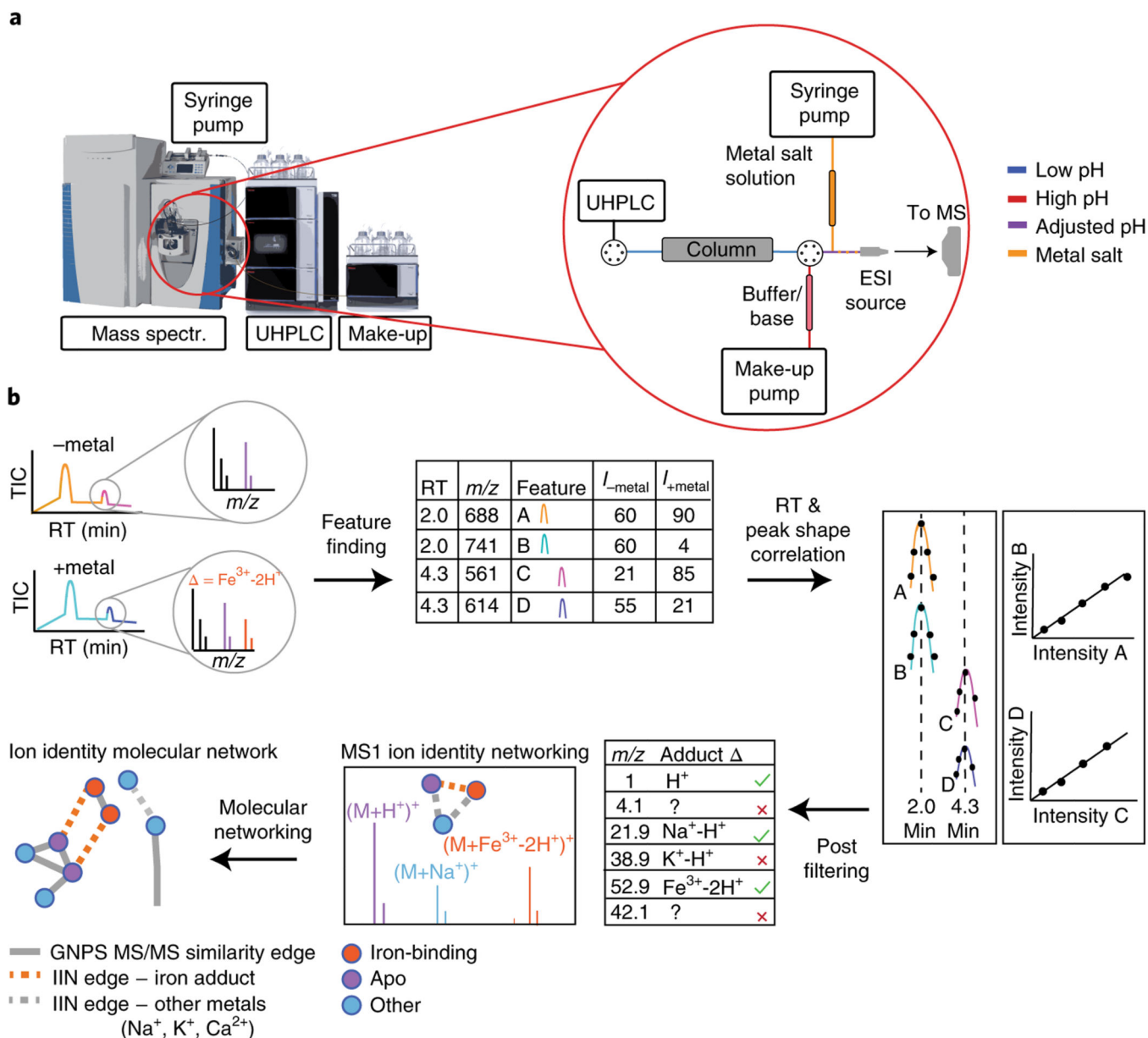


Figure 1. Overview of the native spray small molecule binding experiment.

(a) The two-step native ESI-MS/MS workflow utilizes a post-column infusion of ammonium hydroxide solution or ammonium acetate buffer, followed by infusion of metal salt solution (second-generation workflow shown). (b) Data can be analysed using a computational ion identity molecular networking (IIMN) workflow in MZmine and GNPS.

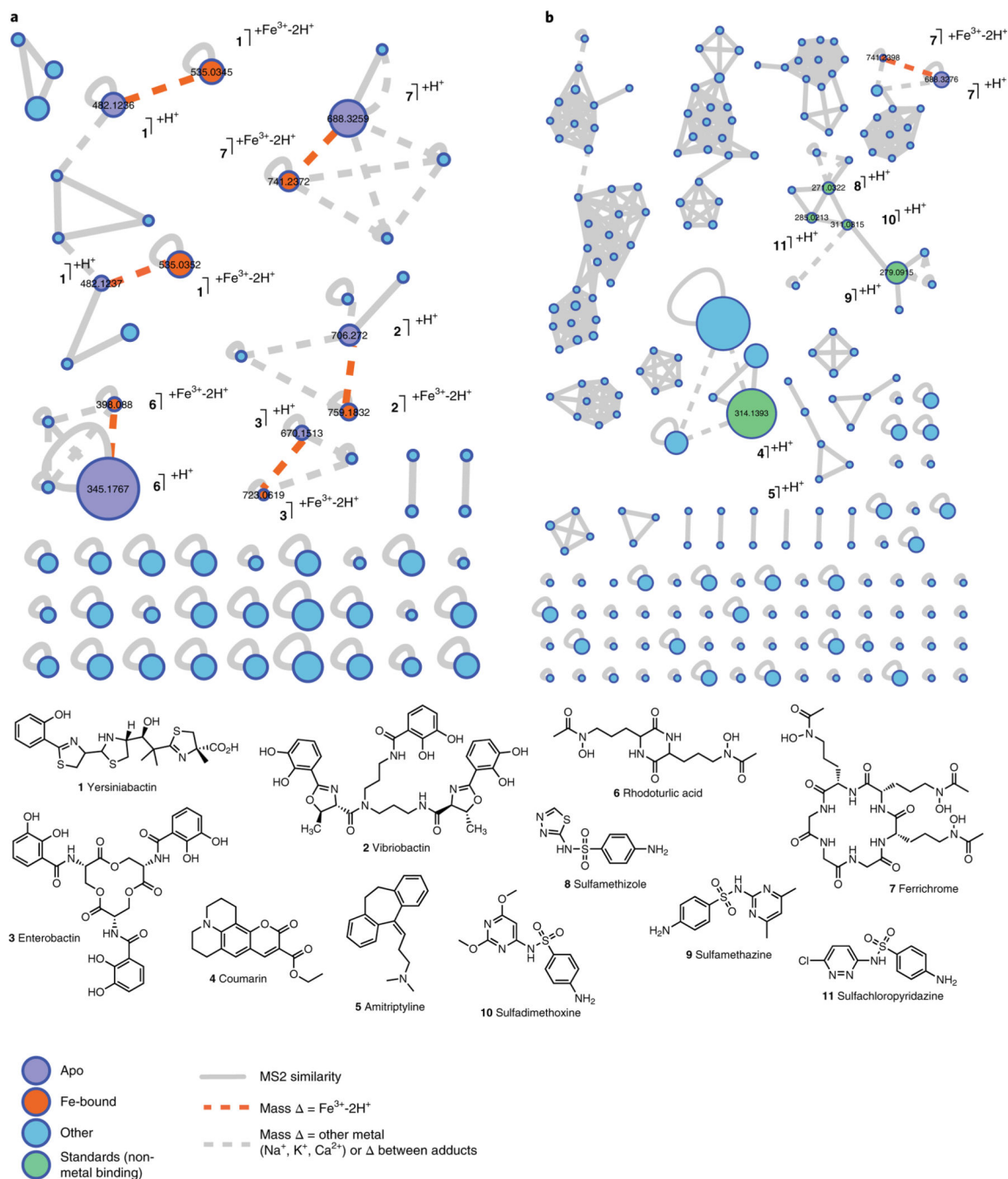


Figure 2. Post LC metal-infusion into mixtures of standards.

(a) Post-LC infusion of Fe^{3+} and post-pH adjustment of a mixture containing ionophore standards (1,2,3,6 and 7). Each of the five ionophore standards is complexed with iron (orange dashed line). (b) Post-LC infusion of Fe^{3+} and post-pH adjustment can be utilized to selectively observe the iron-bound adduct ferrichrome (7) in the presence of molecules (4,5,8,9,10, and 11), that have no reported iron-binding behaviour. Nodes corresponding to an $[\text{M}+\text{H}]^+$ ion of apo siderophores are coloured in purple and nodes corresponding to iron-bound siderophores are coloured in orange. MS/MS similarity edges are solid grey

lines, IIN edges corresponding to an iron m/z delta are dashed orange lines, and IIN edges corresponding to other metals (such as Na^+ , Ca^{2+} , and K^+) or m/z deltas between different combinations of metals are dashed grey lines. The size of the nodes corresponds to the relative area of the feature.

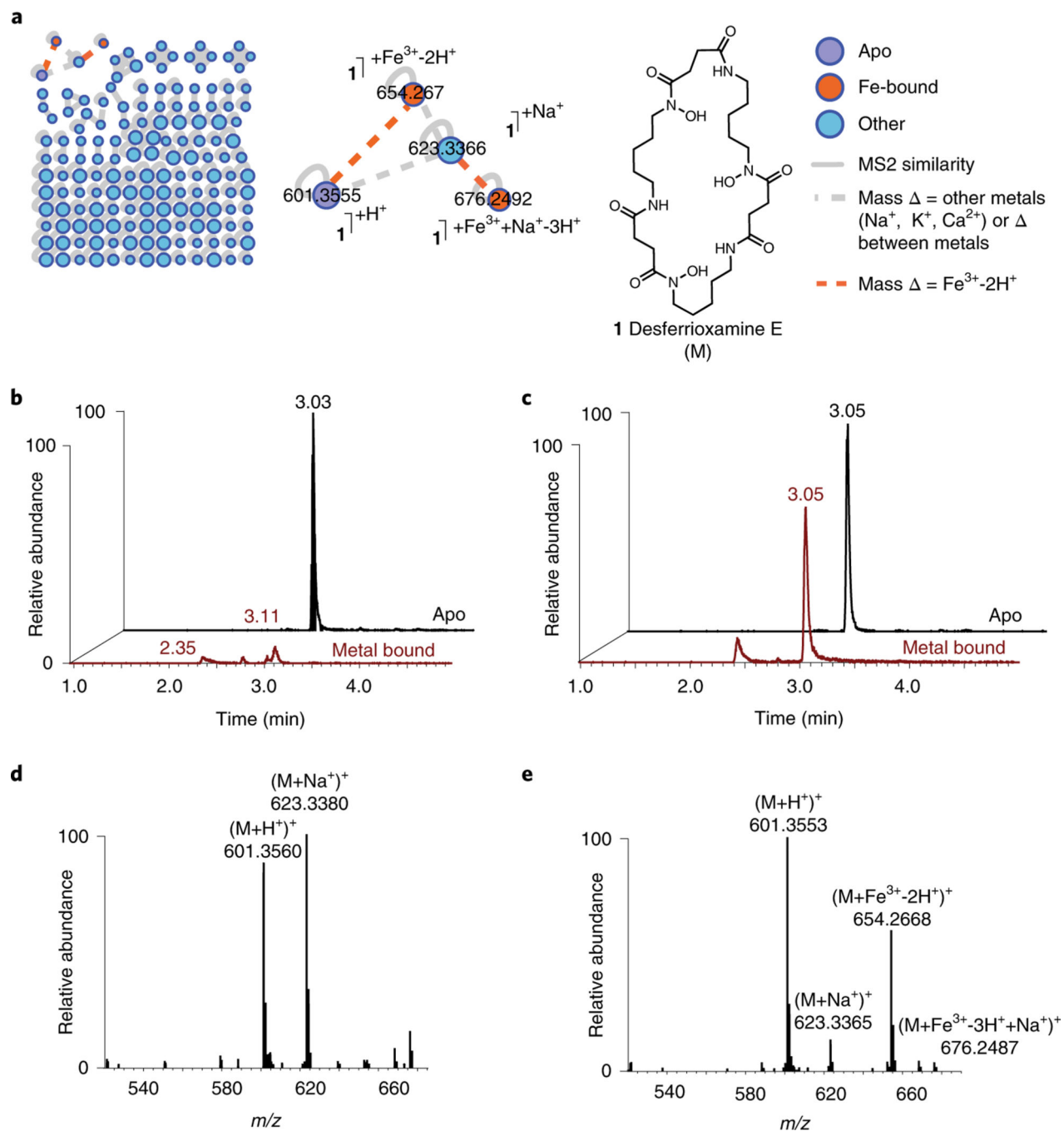


Figure 3. Native spray metal metabolomics is used to identify siderophores in bacterial culture extracts.

(a) Desferrioxamine E (DFE) is identified as the only siderophore present in *Glutamicibacter arilaitensis* JB182 extracts. The only Fe^{3+} -binding IIN edges observed in the entire network correspond to DFE and the sodium adduct of DFE. (b) Extracted Ion Chromatogram of apo-DFE (m/z 601.3560, black) and Fe^{3+} -bound DFE (m/z 654.2668, red) when analysis is performed using standard LC-MS/MS conditions. (c) Extracted Ion Chromatogram of apo-DFE (m/z 601.3560, black) and Fe^{3+} -bound DFE (m/z 654.2668, red) when the analysis is performed using Fe^{3+} -infusion LC-MS/MS conditions. (d) MS^1 of the DFE peak (at

retention time 3.03 min) in (b). Only the apo- and Na⁺-adduct are observed, with an m/z delta of 21.9819, but no Fe³⁺-bound peak is observed. (e) MS¹ of the DFE peak (at retention time 3.05 min) in (c). The apo- and the Fe³⁺-bound peak are observed, with an m/z delta of 52.9115.

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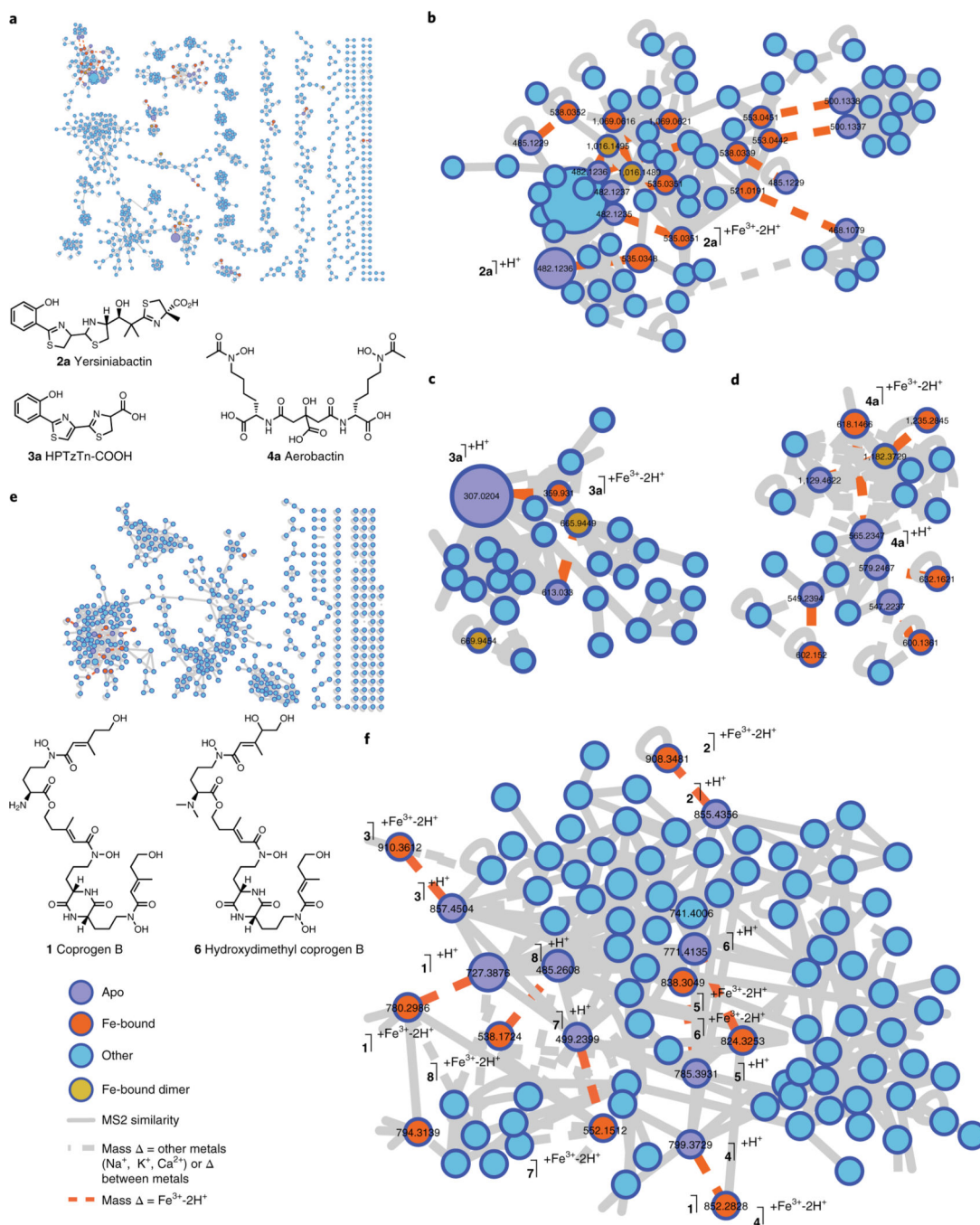


Figure 4. Native spray metal metabolomics is used to identify known and novel siderophores in bacterial and fungal culture extracts.

(a) Several siderophores are observed in *E. coli* Nissle culture extracts when a post-LC infusion of Fe^{3+} and post-pH adjustment is performed. Singleton nodes have been removed from this network. A number of (b) yersiniabactin (**2a**) derivatives, including (c) the truncated biosynthetic intermediate of yersiniabactin, HPTzTn (**3a**), in addition to (h) aerobactin (**4a**) derivatives are identified using native metal metabolomics. These derivatives are tabulated in Table 1a. (e) Several siderophores are also observed in *Eutypa lata* culture

extracts when native metabolomics is used, (f) corresponding to coprogen B (**1**) and hydroxydimethyl coprogen B (**6**) derivatives (**2–5**, **7–8**), which are tabulated in Table 1b.

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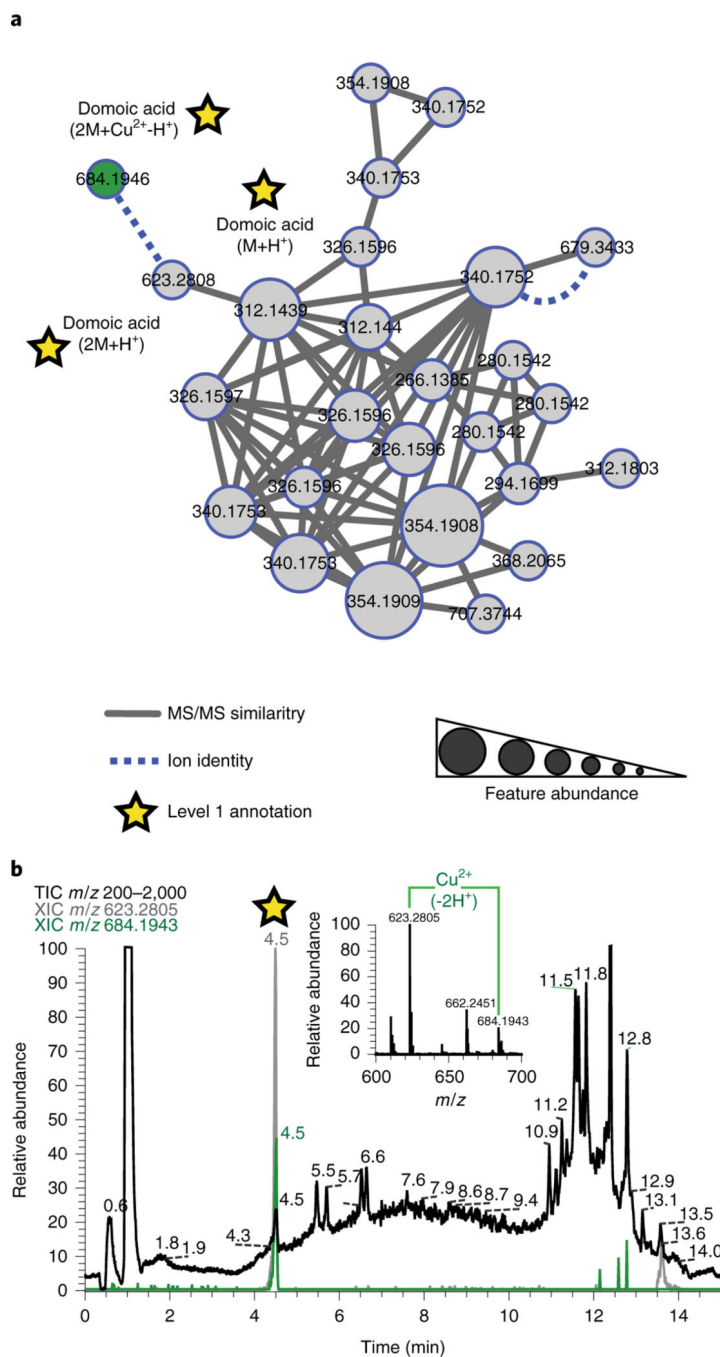


Figure 5. Native spray metal metabolomics of dissolved organic matter samples. Surface Seawater Samples were collected in the California Current Ecosystem during a phytoplankton bloom in June 2017. (a) Ion-Identity Molecular Network of the domoic acid (DA) molecular family shows a DA dimer as well as this dimer bound to Cu^{2+} . The identity of DA was confirmed as level 1 with an authentic standard in its apo, apo-dimer and the dimerholo form. (b) Total ion chromatograms (TIC) and extracted ion chromatograms (XIC) of DA dimer and the copper-bound dimer.

Table 1.

Iron-binding derivatives identified in *E. coli* Nissle (a) and *Eutypa lata* (b) cultures have been tabulated above.

	m/z [M+H ⁺] ⁺	Mol. Formula
1a	325.0311	C ₁₃ H ₁₂ N ₂ O ₄ S ₂
2a	482.124	C ₂₁ H ₂₇ N ₃ O ₄ S ₃
3a	307.021	C ₁₃ H ₁₀ N ₂ O ₃ S ₂
4a	565.229	C ₂₂ H ₃₆ N ₄ O ₁₃
5a	478.0921	C ₂₁ H ₂₃ N ₃ O ₄ S ₃
6a	468.108	C ₂₀ H ₂₅ N ₃ O ₄ S ₃
7a	485.116	*
8a	500.136	C ₂₁ H ₂₉ N ₃ O ₅ S ₃
9a	547.226	C ₂₂ H ₃₄ N ₄ O ₁₂
10a	549.246	C ₂₂ H ₃₆ N ₄ O ₁₂
11a	579.302	C ₂₃ H ₃₈ N ₄ O ₁₃
12a	1129.462	*
13a	297.0762	C ₁₃ H ₁₆ N ₂ O ₂ S ₂
14a	295.0569	C ₁₃ H ₁₄ N ₂ O ₂ S ₂
15a	449.1187	C ₂₀ H ₂₀ N ₂ O ₁₀
16a	465.1137	C ₂₀ H ₂₀ N ₂ O ₁₁
17a	327.0466	C ₁₃ H ₁₄ N ₂ O ₄ S ₂
18a	627.167	C ₂₆ H ₃₀ N ₂ O ₁₆

	m/z [M+H ⁺] ⁺	Mol. Formula
1	727.381	C ₃₃ H ₅₅ N ₆ O ₁₂
2	855.437	C ₃₉ H ₆₂ N ₆ O ₁₅
3	857.456	C ₃₉ H ₆₄ N ₆ O ₁₅
4	799.372	C ₃₅ H ₅₄ N ₆ O ₁₅
5	785.393	C ₃₅ H ₅₆ N ₆ O ₁₄
6	771.41	C ₃₅ H ₅₈ N ₆ O ₁₃
7	499.24	C ₂₂ H ₃₄ N ₄ O ₉
8	485.26	C ₂₂ H ₃₆ N ₄ O ₈

^(*)The molecular (mol.) formula corresponds to the top-scoring formula with the top score assigned in SIRIUS 4.0¹, and those compounds with one or less matched peaks in SIRIUS 4.0 have been labelled with an asterisk.