Reviews and syntheses: Use and misuse Opportunities for robust

<u>use</u> of peak intensities from high resolution mass spectrometry in organic matter studies: opportunities for robust usage

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Abstract Earth's biogeochemical cycles are intimately tied to the biotic and abiotic processing of organic matter (OM). Spatial and temporal variation in OM chemistry is often studied using direct infusion, high resolution Fourier transform mass spectrometry (HRMSFTMS). An increasingly common approach is to use ecological metrics (e.g., within-sample diversity) to summarize high-dimensional HRMSFTMS data, notably Fourier transform ion cyclotron resonance MS (FTICR MS). However, problems can arise when HRMSFTMS peak intensity data are used in a way that is analogous to abundances in ecological analyses (e.g., species abundance distributions). Using peak intensity data in this way requires the assumption that intensities act as direct proxies for concentrations, which is often invalid. Here we discuss theoretical expectations and provide empirical evidence why concentrations do not map to HRMS peak intensities. The theory and data. Here we show that comparisons of the same peak across samples (within-peak) may carry information regarding variation in relative concentration, but comparing different peaks (between-peak) within or between samples does not. We further developed a simulation model to study the quantitative implications of using peak intensities to compute ecological metrics that rely on information about both within-peak and between-peak errors that decouple concentration from intensity. These implications are studiedshifts in terms of commonly used ecological metrics that quantify different aspects of diversity and functional trait values relative abundance. We showfound that despite the poor linkages between analytical limitations of linking concentration and to intensity, the ecological metrics often perform well in terms of providing robust qualitative inferences and sometimes quantitatively-accurate estimates of diversity and trait values molecular characteristics. We conclude with recommendations for using robust use of peak intensities in an informed and robust way for natural organic matter studies. A primary recommendation is the use and extension of the simulation model to provide objective, quantitative guidance on the degree to which conceptual and quantitative inferences can be made for a given analysis of a given dataset. Without objective guidance, researchers that Broad use peak intensities are doing so with unknown levels of uncertainty and bias, potentially leading to spuriousthis approach can help ensure rigorous scientific outcomes from the use of FTMS peak intensities in environmental applications.

# 1 Introduction

Organic matter (OM) plays a central role in Earth's biogeochemical cycles, and is both a resource for and product of metabolism. The detailed chemistry of OM (e.g., nominal oxidation state) can modulate and reflect biogeochemical rates and fluxes within and across ecosystems (e.g., LaRowe and Van Cappellen, 2011; Boye et al., 2017;

Garayburu-Caruso et al., 2020), yet our understanding of this complexity is limited by our analytical abilities to

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view it (Steen et al., 2020; Hedges et al., 2000; Hawkes and Kew, 2020a). Given the importance of OM chemistry to biogeochemical cycling, there is a need to understand how and why that chemistry varies through space and time. To help meet this need, there has been growing interest in using concepts and methods from ecology to study the chemogeography and chemodiversity of OM in a variety of ecosystems (e.g., Kujawinski et al., 2009; Kellerman et al., 2014; Tanentzap et al., 2019; Danczak et al., 2021). This is a promising approach as there are many conceptual parallels between the chemical species that comprise OM and the biological species that comprise ecological communities (Danczak et al., 2020).

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The most fundamental ecological data type is the species-by-site matrix. This matrix indicates how many individuals of each species occur in each sampled community. Ecologists use species-by-site matrices to ask myriad questions related to biological diversity. Two common analyses are known as  $\alpha$ -diversity and  $\beta$ -diversity, and there are acch with numerous metrics for each (Whittaker, 1972; Anderson et al., 2011).  $\alpha$ -diversity measures the diversity within a given community.  $\beta$ -diversity has been variously defined, but essentially measures variation in composition across communities. Both  $\alpha$ -diversity and  $\beta$ -diversity can be quantified using presence-absence data or they can include estimates of each species' relative abundance within and between communities (Fig. 1).

The chemistry of OM is commonly studied using high resolution mass spectrometry (HRMS) techniques (e.g., Hawkes and Kew, 2020b). Specifically, Fourier transform mass spectrometry (FTMS) techniques are predominantly used, i.e., (e.g., Hawkes and Kew, 2020b), such as Orbitrap or Ion Cyclotron Resonance (ICR) MS, via direct infusion of samples. At present, the highest resolution approach for untargeted analysis of OM is via a 21 Tesla FTICR MS (Marshall et al., 1998; Shaw et al., 2016; Smith et al., 2018; Bahureksa et al., 2021). The output data produced is a spectrum containing peaks represented by a signal intensity (Fig. 2 y-axis) and a mass-to-charge ratio (m/z) (Fig. 2 x-axis), which is equivalent to the mass for singly charged ions as routinely detected in natural organic matter (NOM) measurements. In turn, regardless of the type of MS instrument used, the MS data inherently lead to an OM peak-by-sample data matrix, akin to an ecological species-by-site data matrix. The high resolution data from MS often results in a large matrix, wherein a single sample may contain thousands to tens of thousands of peaks. To take advantage of these rich data, HRMSFTMS data have been analyzed using the same  $\alpha$ -diversity and  $\beta$ -diversity metrics that are commonly used by ecologists to study biological diversity (e.g., Kellerman et al., 2014). This isSuch analyses are exciting, as it allowsthey enable the same conceptual questions and quantitative frameworks to be applied to biological (e.g., microbial communities) and chemical (i.e., OM) components that directly interact with each other within ecosystems (Lucas et al., 2016; Osterholz et al., 2016; Li et al., 2018; Tanentzap et al., 2019; Danczak et al., 2020, 2021).

The use of ecological metrics with MS data is particularly common with FTMS datasets, and eontainsthere is great potential to continue leveraging concepts from ecology in high-resolution OM analyses. However, when Care is required, however, in using FTMS peak intensity data are used in the estimations ofto estimate α-diversity, βdiversity, and related ecological analyses (e.g., 'species' abundance distributions), potential problems can arise. At the root of). Key to these problems lies ecological analyses is the assumption that within complex NOM samples, differences in peak intensity are proportional to differences in concentrations of the associated molecules. Consequently, the Studies using FTMS often avoid using peak intensities due to uncertainties in whether it is valid to assume proportionality between peak intensities and concentrations within and across NOM samples (Kujawinski, 2002). These studies may be discarding useful information, though it is unclear what biases and uncertainties are introduced by relying on this assumption are unclear. In certain situations, however, peak intensity-based ecological analyses of MS data can provide valid information—even when the underlying assumption is invalid—and the extent to which such situations exist is likewise unclear into ecological metrics when using FTMS peak intensities. To help advance the robust use of FTMS datasets that has been emerging in environmental science studies over the last few decades for NOM studies, we review the theoretical reasons why between-peak intensities domay not correspond to differences in concentrationreflect true concentrations, provide empirical support for our assertions, use evaluation of this theory, and invoke in silico studies simulation to quantify the associated impacts on ecological

analyses, provide. While theory and empirical analyses demonstrate disconnects between peak intensities and concentrations in FTMS data, the simulations show that ecological metrics are often still robust. We end with practical recommendations, and propose a path forward that may eventually enable improved usage for increasing robust use of FTMS peak intensities for quantification NOM studies.

## 2 Theoretical Limitations Foundations

To address Here we provide a review of the theoretical foundations behind why assuming proportionality between peak intensities and concentrations in FTMS eannot be used to infer between-peak changescan be challenging. This section will be of most value to FTMS data users that are not formally trained in concentration, we review critical theoretical concepts about mass spectrometry, and serves as a review of mass spectrometry principles (see also Kujawinski, 2002; Urban, 2016; Bahureksa et al., 2021). We focus on FTMS (i.e., FTICR and Orbitrap), but many of the principles and limitations—especially ionization and ion transmission—are applicable across all MS platforms. In this section, weWe highlight three main mass spectrometry considerations: ionization, ion transfer, and ion signal detection in the context of a generalized commercial FTICR mass spectrometer. Theoretical limitations have two mainFTMS instruments. These considerations have practical implications tied to within-peak and betweenpeak comparisons (Fig. 2). Here, we define 'within-peak comparison peak intensities of the same feature (i.e., m/z or molecular formula) across different sample spectra (i.e., within two or more and 'between-peak' as comparing peak intensities across different features. As discussed below, within-peak comparisons can be robust under certain situations, but there are limitations with between-peak comparisons that may be unavoidable. The following discussion is not an exhaustive treatment of all decisions associated with a complete FTMS experiment, and we do not deeply address factors such as sample spectra), whereas between peak comparison occurs between different features (*m/z* or molecular formulas) across the same spectra.

The first implication is that if preparation, choice of ionization mode, and instrument parameters are kept consistent, within-peak/between-sample biases are minimized, though between-peak/within-sample biases are inherently unavoidable. The second implication is that because of inherent sample and matrix variation and subsequent effects, between-peak/between-sample biases can be significant and may be indeterminable. specific parameter optimization. These topics have been discussed in a recent review (Bahureksa et al., 2021).

# 2.1 Ionization Biases Efficiency and Isomers

Electrospray ionization (ESI); is the most commonly usedcommon technique for generating ions from NOM samples, is a 'soft' ionization technique that predominantly yields intact molecular ions. Generally, When using ESI of NOM samples produces protonated or deprotonated ion types (positive or negative polarity, respectively), which can only be formed if some pre-existing basic or acidic functionality is available in the molecule to support this. ESI also commonly produces adduct ions, such as sodium adducts [M+Na]\* in positive mode and chloride adducts [M+CI]\* in negative mode. The ionization efficiency of the peak intensity for any given molecule dependsmolecular mass (or molecular formula) will depend on its both concentration and ionization efficiency, the latter of which is dependent on structure, pKa, and the other molecules in the sample matrix and composition (Kruve et al., 2014). Ionization suppression occurs when multiple species are present in a sample, and the ionization efficiency of one analyte is altered by the presence of another (Ruddy et al., 2018). These issues all confound when dealing with complex samples with unknown compositions. While users can apply controls to account for some matrix differences (concentration, solvent, pH), the unknown (and unknowable) differences in molecular composition of complex mixtures cannot be accounted for, and therefore comparison of peak intensities in different samples is prone to uncertainty.

Importantly, in these highly complexIn NOM samples, one detected mass or peak commonly combines signals from multiple different isomers, i.e., which all have the same molecular formula but with a different structure. While

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the structures. The different structures impact the ionization efficiency of a given molecule, the recorded spectrum shows the superposition of these isomers. To, but FTMS data contains no information about this structural variation. Unfortunately, to date, no liquid chromatography (Kim et al., 2019; Han et al., 2021) or ion mobility separation (Tose et al., 2018; Leyva et al., 2020) technique has yet demonstrated sufficient resolution for the most complex of samples (such as NOM), instead yielding broad distributions in these orthogonal dimensions. Therefore, not only must we apply extreme caution in inferring chemical properties from to completely infer structural variation among isomers within complex NOM samples. Unknown variation in structure can, therefore, lead to unknown variation in peak intensities. This challenge can be compounded by ionization suppression that occurs when the ionization efficiency of one type of molecule (i.e., peak) is altered by the presence of other types of molecules (Ruddy et al., 2018). Ionization suppression can be mitigated by online separation whereby non-targeted LC-MS approaches may yield more quantitative data (Kruve, 2020), but matrix effects remain a significant issue even for LC-MS (Trufelli et al., 2011). In NOM samples with thousands of types of organic molecules, the molecular formula alone, but we must also be aware of underlying subtleties that may distort comparisons ofinteractions likely have complex influences over realized ionization efficiencies. While it is possible to control for some of these challenges (e.g., using consistent sample concentrations and preparations), many additional factors (e.g. molecular formula and structures, pKas, and interactions among molecules in NOM samples) cannot yet be accounted for. Interpretation of peak intensities within samples such as the presence of isomersas proxies for concentrations in FTMS datastreams may, therefore, be prone to uncertainty.

#### 2.2 Ion transmission and collection

Before ions can be detected in the (ion) trap, they must be transmitted from the instrument source to the trap. Ion transmission, including ion accumulation, is not unbiased. Ions are manipulated through the instrument ion optics, across differential pressure regimes, using radiofrequency (RF) and direct current (DC) potentials to guide, focus, and accumulate the ions. The specific values of these parameters have effects on the mass ranges transmitted. Further, the specific timings, geometries, and vacuum regimes all have effects upon ion transmission efficiency and biases. For this reason, quantitative comparison of intensities across widely differing m/z is not directly possible.

In FTMS, packets of ions are accumulated and 'cooled' in a trap prior to their transmission to the analyzer cell (Fig. 3 Panel A section d; Senko et al., 1997; Makarov et al., 2006). The duration of time in which ions are accumulated is often varied to yield an optimal ion population for the analyzer cell, which has a finite charge capacity. The duration of this event has been directly observed to can change the relative ion populationsabundance, and thus observed peak intensities of different ions (Cao et al., 2016). Thus, when balancing Increases in the need for controlled true abundance of other ions can decrease the measured peak intensity of a given ion due to a dilution effect resulting from a finite number of ions that can fit within the ion populations—critical for a high resolution, high fidelity measurement—and minimaltrap. Additional challenges arise due to variation in ion-the speed at which different ions move from the accumulation trap and into the analysis cell. Smaller ions move more quickly and therefore reach the analysis cell sooner than larger ions. Variation in the accumulation time, there is a risk of further biasing the relative ion intensities across samples and FTMS instruments, combined with among-ion variation in transmission speed, can introduce additional uncertainty in the relationship between peak intensities and true concentrations.

Finally, time of flight biases come into play in FTICR MS. Ions are transmitted from the ion accumulation trap to the ICR cell along one or more transfer multipoles (Fig. 3 Panel A section e). The distance between ion accumulation trap and ICR cell may be quite long, e.g., 2.4 meters on a 21 Tesla instrument (Shaw et al., 2016), and therefore the time required for ions to travel this distance (the 'time-of flight', a millisecond or longer) may cause dispersion in the ion packet (Fig. 3 Panel B). While the packet of ions may leave the accumulation trap simultaneously, because smaller ions travel faster the packet arrives at the analyzer cell as a dispersed distribution of ions. Therefore, only a subset of this population, with regards m/z-range and ion energies, is optimally trapped in the

ICR cell. Thus, these biases in ion transmission do not allow for quantitative comparison of peak intensities between ions with differing m/z ratios.

#### 2.3 Ion signal detection

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To directly use FTMS peak intensities quantitatively, we must first understand how those intensities arise and the biases which can affect them. In ion trapping measurements, such as FTICR and Orbitrap MS, the motion of ions within a static magnetic (ICR) or electric (Orbitrap) field induces an image current upon The final step in data collection via FTMS is signal detection-electrodes. The frequency of this motion is proportional to the mass-tocharge ratio (m/z) of the ion, while the. The intensity of the signal is proportional to the abundance of thea given ion in the trapanalysis cell, the proximity of the ionions to the electrodedetector (Kaiser et al., 2013), and the ion charge state of the ion (Wörner et al., 2020). Thus, at a first approximation, the signal intensity between Similar to molecular interactions impacting ionization efficiencies, different m/z ions could be compared provided they are excited to the same radius (ICR) and have the same charge state (Fig. 3 Panels B, C). However, these provisions are not always met. Ions with very close frequencies, which are not fully resolved, may types of ions can interact to affect each other's signal intensity, and the. The Fourier transform does not allow for applied to the data also complicates extremely accurate relative quantification of ion abundance between peaks (Makarov et al., 2019). With FTICR, most commercial (e.g., Bruker) instruments use a CHIRP, or frequency-swept, excitation pulse which does not excite all ions to exactly the same radii (Kaiser et al., 2013). In addition, while most ions in NOM mass spectra are singly charged, some mass spectra contain multiply charged interferences (Smith et al., 2018; Patriarca and Hawkes, 2021). Still, in both instrument types, signal intensities may be used to describe the ion populations quantitatively provided that the charge states are the same, a flat excitation profile is used (or the ions are sufficiently close in frequency space such that they are excited to the same radii), and the user clearly understands that the ion population in the trap may not accurately reflect the molecular composition of the sample.

Within a well-designed experiment and a constrained sample set, many of these points may be mitigated. However, objectively proving the degree of mitigation is non-trivial, and there remains great These challenges at the detection stage can add more uncertainty aboutto the relationship between peak intensity and molecular concentrations, particularly for complex matrices such as NOM. Furthermore, as shown in a recent interlaboratory study (Hawkes et al., 2020), measuring the same samples with different instrumentation can lead to differing results, thus further highlighting potential pitfalls in quantitative analysis of these dataNOM samples.

# 3 Empirical Limitations Evaluations

Despite the aforementioned fundamental and

In this section, we move beyond theoretical limitations and uncertainties in using peak intensity data, it is still helpfulconsiderations to demonstrate these limitations with empirical evaluations of the real-world empirical measurements. In this section, we demonstrate, with ideal and non-ideal samples, the non-quantitative nature of these measurements-relationships between peak intensities and concentrations. Similar to above, this section will be of primary value to those without formal training as mass spectrometrists, but who use FTMS data to study NOM.

### 3.1 Direct comparison of signalpeak intensities in idealized samples is problematic

In the ideal case, samples are analyzed with identical matrices, equivalent concentrations for each compound, and free from competitive ionization/ionization suppression (Ruddy et al., 2018). However, even in this ideal case, different molecules ionize with different efficiencies, and thus their signal intensities are not equal. To demonstrate the non-equal response for various analytes in various conditions, we acquired a series of contemporaneous mass spectra of several compounds in different conditions. First, in Fig 4A, we prepared three dilution ladders of three pure compounds—analyzed separately—in pure methanol. Clearly, these three molecules yield starkly different signal intensities for otherwise identical conditions, and thus directly comparing their intensities would not be a valid means to infer their relative concentrations in solution. At an extreme, trehalose, a carbohydrate, yields nearly

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as little signal at 500ppb as sinapic acid does at 200ppb. Even between the two structures containing a carboxylic acid moiety—a typical indicator of good negative mode ESI response—there is a significant difference in signal intensity. Thus, directly comparing the signal intensities of different ions—even in idealized situations—cannot be used as a proxy for concentration or abundance determination absent a calibration curve.

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Subsequently, we highlight the challenge of comparing ions of the same exact mass. Here, in Fig 4B, we again prepared dilution ladders of three pure compounds in methanol, however these are all structural isomers with the same molecular formula and thus exact mass. Again, a stark difference in signal intensity is observed, even between nominally similar structures. This issue is particularly troubling for direct infusion measurements of complex mixtures, where we do not, and cannot, know the structural identity of individual peaks, and instead are limited to molecular formulas. Thus, if we compare peaks with the same exact mass, same molecular formula, between different samples, we cannot be sure that they are the same molecule, and thus again comparing their signal intensities as a proxy for abundance is problematic. Additionally, structural isomers can have vastly different ecological/biogeochemical function, and therefore this consideration is important to note for subsequent interpretations of NOM samples. Further complicating this issue is the known fact that in highly complex mixtures like organic matter, most - or all - peaks are actually the superposition of multiple different isomeric compounds. Demonstrated by chromatography (Kim et al., 2019) or ion mobility separations (Leyva et al., 2019), or by statistical inference of tandem mass spectrometry (Zark et al., 2017), each peak may be several isomers of various relative intensities. Thus, even if the same isomers were present across samples, it cannot be known that their relative abundances are the same - and again, it is problematic to directly compare the intensities of signal corresponding to nominally the same molecular formula across different mass spectra.

One caveat with the above experiments, of course, is that it is a direct infusion measurement. The chemicals used were nominally pure, but any trace impurity—either from their production and isolation, or from sample preparation—may impact the measured signal intensity. Which leads us to the next point—matrix effects are intrinsically challenging to control for, and have significant impacts on mass spectra.

### 3.2 Matrix effects substantially impact signal intensities in complex mixtures

Of course, analyses are often performed on complex mixtures, containing As discussed above, different organic compounds ionize with different efficiencies. In theory, this may lead to variation in observed peak intensities even when all organic compounds have the same true concentration. To evaluate this theoretical expectation, we analyzed several different types of organic compounds in different conditions via FTICR-MS. We selected chemical standards which are natural products with molecular formula and chemistries typical of compounds commonly observed in organic matter, and were amenable to negative mode ESI analysis. First, we analyzed three separate dilution ladders of individual pure compounds dissolved in pure methanol. These standards were analyzed at higher concentrations than typically observed for NOM because they were single compounds rather than formula-summed features (with multiple isomers) within a NOM spectrum; higher concentrations were required to compensate for lower isomeric diversity. These three compounds gave rise to different peak intensities under otherwise identical conditions (Fig. 4A). Trehalose, for example, had much lower peak intensity than sinapic acid at the same actual concentration. The difference in signal intensity was also apparent amongst compounds that ionize well under negative mode ESI; for example, two different structures containing the same number of carboxylic acid units exhibited differences in signal intensity. We also observed differences in peak intensities amongst structural isomers (i.e., same molecular formula and mass) (Fig 4B). Each peak observed via direct infusion FTICR-MS may be several isomers. These isomers may be observable through chromatographic separation (Kim et al., 2019), ion mobility separations (Leyva et al., 2019), or by statistical inference of tandem mass spectrometry (Zark et al., 2017), but not via direct infusion FTICR-MS. We note that absolute differences in signal intensity may be smaller between molecules at lower concentrations, but this does not necessarily mean that low intensity signals consistently indicate low concentrations and this does not aid in quantitatively interpreting higher intensity signals. In summary, differences in peak intensities across organic

compounds do not necessarily equate to differences in concentration, unless assessed via a calibration curve for each compound.

### 3.2 Comparison of peak intensities in in real world samples

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Routine NOM samples contain a diverse range of thousands of molecules of unknown structures and relative concentrations. Furthermore, samples\_and\_often contain 'inorganic' interferences, such as salts. Routinely, scientists will desalt samples with solid phase extraction, but such processes can leach impurities into the sample, don't necessarily remove all interferences, and can remove select pools of NOM due to their functionalities, depending on the sorbent or resinSample clean up that focuses on pre-concentration and desalting is imperfect (Racke et al., 2016; Li et al., 2017). As such, real world non-ideal samples contain a multitude of ), but is commonly used to minimize inorganic interferences. Interactions among molecules remains a challenge, however, as discussed above. The collection of molecules in a sample is referred to here as the 'matrix.' To explore matrix effects and sources for ionization suppression, or adduct formation, which yield spectra that are even more challenging to quantitatively interpret.

To explore the impacts of matrix effects (Fig. 4C-E), on peak intensities, we prepared solutions of six different pure compounds at a fixed concentration (100ppb100 ppb) in three different solvent systems - pure methanol, methanol eluted from elution off of a BondElut SPE cartridge, and methanol from elution off of a BondElut SPE cartridge which had been loaded with artificial river water (ARW). Additionally, we added a complex mixture -that is often used as a NOM standard, Suwannee River Fulvic Acid (SRFA), at six different concentrations, to each sample. Again, samples Samples were analyzed independently but contemporaneously on the same instrument to mirror a real study.

In methanol\_only solvent, with no addition of added SRFA, the six compounds —as expected—yieldyielded different signalpeak intensities (Fig. 4C), further confirming what was previously observed which is consistent with results from the previous subsection. As the concentration of SRFA iswas increased to 2 ppm, the relative signal intensity increasesincreased for some of these analytes—possibly as a function of endogenous molecules with the same molecular formula as those spiked in—the six compounds, but decreased creased for others. Above 2 ppm of SRFA, however, all signalspeak intensities for our referenceall six compounds arewere substantially decreased, most likely as a result of competitive ionization effects of the addition of the complex mixture.

\_Use of an 'impure' methanol solvent, i.e., the eluent from a SPE blank (Fig. 4D) or from an SPE of artificial river water (Fig. 4E), resultsresulted in even more ionization suppression and differential signal response.further decreases in peak intensities. In both cases, the maximum signalpeak intensity is only-was ~20% of what was seen in pure methanol (Fig. 4C), indicating that and some of the leachate or residual salts from the SPE protocol impacted sensitivity. Further, here only two analytes (asseulin and chlorogenic acid) ionize well at all, with the other 4 yielding poor orsix compounds were no signal-longer observed. Addition of SRFA to these samples with 'impure' solvents, again, decreases signal intensity, though at 40 ppm SRFA some minor features increase, likely due to endogenous features with the same molecular formula as our standardsgenerally, decreased peak intensities.

Cumulatively, Combining the empirical evidenceresults from this subsection and instrumental the previous subsection with instrument theory demonstrate that it is not possible—with discussed above suggests significant uncertainty in relationships between true concentrations and peak intensities from direct infusion measurements of complex mixtures—to directly compare signal intensities as a proxy for molecular abundance between different peaks within a spectrum, or between the same peak across spectra, even in idealized cases. Strategies to use calibration FTICR-MS. Calibration curves will fail due to unknowncan be used in the simplest of situations, but may be challenging when there are structural isomers and sample-to-sample variation in matrix composition, and established normalization techniques cannot factor in the large range of sources of experimental variation. That said,

there may be cases where a high level comparison of trends can yield valid semi-quantitative comparisons between spectra, relying on a statistical aggregation of individually unreliable trends. Additionally, modeling. Modeling of constrained systems may, however, allow for improved, data-driven and mechanistic based machine learning data normalization strategies for enhanced use of peak intensity data.

### 4 Conceptual implications for use of ecological metrics

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The preceding sections have shown both theoretically and empirically that there are indicate challenges towhen using HRMSFTMS peak intensities as proxies for relative changes in concentrations of organic molecules. The implication is that there may be specific kinds of some ecologically-inspired analyses (e.g., Fig. 1) that are or are not appropriate1) may be challenging to use with HRMSFTMS peak intensity data. To understand what may or may notwhich analyses could be a valid analysis, it is critical to impacted, we differentiate analyses into two classes: those based on within-peak intensity comparisons and those based on between-peak intensity comparisons (Fig. 2). As noted above, within-peak is based on comparing the same feature (m/z or molecular formula) across spectra/samples, whereas between-peak compares different features (m/z or molecular formulas) across and within spectra/samples.

AnalysesWe posit that analyses using FTMS between-peak intensity comparisons are could have the most likely to be problematic. To help clarify why this is, consider greatest uncertainty. Consider an ecological setting in which a researcher aims to quantify within-sample diversity ( $\alpha$ -diversity) and among-sample diversity ( $\beta$ -diversity) (Fig. 1) of tree communities (Fig. 5, left-side). The researcher will likely set up a plot of a given size and then directly count the number of each tree species in each plot. This generates, thus generating the species-by-site matrix filled with directly observed abundance counts for each species. In such a situation, the The ability of the researcher to observe individuals of each species does not vary appreciably across species because each tree is not moving and our ability to see ita static object is not influenced by environmental factors. In turnThus, the number of individuals observed for a given tree species is quantitatively comparable to the number of individuals observed for all other tree species in the plot. The assumption that differences in observed abundances carry robust information about differences in actual abundances is thus supported, in this example. In turn, it is valid to use relative abundances to compute  $\alpha$ -diversity such as via Shannon evenness (Elliott et al., 1997; Mouillot and Leprêtre, 1999; Redowan, 2015). Furthermore, because the ability to observe each tree species is the same across communities. In turn, it is valid to use relative abundances to compute  $\beta$ -diversity (e.g., via Bray-Curtis; Anderson et al., 2011) or conduct any other ecological analysis analyses that uses abundance data (e.g., species abundance distributions McGill et al., 2007).

We contrast this tree community example with another ecological setting. Consider a researcher studying bird communities (Fig. 5, right side) that estimated species abundances solely based on the number of times an observer hears the call of a given species. In this case, those species that call more frequently and/or more loudly (will be more likely to be heard), and thus an observer will be inferred to haveinfer a higher abundance even if all species in the community have the same abundance. That is, such a method generates data that may indicate which species are present, but the 'call counts' do not carry reliable information regarding absolute or between-species relative abundances. Follow-on analyses of  $\alpha$ -diversity and  $\beta$ -diversity should, therefore, be limited to approaches that use presence/absence data, and species abundance distributions cannot be quantified.

If we continue with the bird community example and assume that the detectability of a given bird species is consistent across sampled locations (or times); then it would be appropriate to examine variation in within-species call counts. This within-species analysis is directly analogous to the HRMSFTMS within-peak time series analysis in Merder et al. (2021), discussed below. However, if call counts of a given species are suppressed by the presence/or abundance of other species, then call counts of a given species domay not indicate an increasechanges in its abundance. ThisThe call count example is directly analogous to influences of the OMNOM matrix: if the

presence/abundance of a given organic molecule modifies the ionization of other molecules, then within-peak changes in intensity domay not indicate changes in their concentrations concentration. In turn, analyses based on within-peak intensity comparisons are not always valideould lead to error and uncertainty in values of computed ecological metrics, especially if there are significant cross-sample changes in the OMNOM matrix.

Unfortunately, as demonstrated in the previous sections, HRMS data align with the bird community examples and never reflect the tree community example. The the unique chemistry of every molecule fundamentally results in differentin a NOM sample can influence ionization properties for other molecules. In the sample. Thus, FTMS data align with the bird community example rather than the tree community example, with the differing physics of each molecule strongly influences influencing between-peak differences in peak intensity. Those molecules Molecules that more readily ionize more easily result inwill produce higher peak intensities, which is akin to bird species that call more frequently with noisier or more loudly resulting in a numerous calls producing a larger number of 'call counts.' In turn that do not accurately represent the underlying population distribution. Similarly, between-peak differences in intensity as observed via FTMS cannot be directly used as a proxy to indicate between-peak differences in concentration. This could invalidate the application of ecological metrics that use between-peak differences in intensity.

In contrast to between-peak comparisons, within-peak comparisons examine changes in <a href="mailto:the">the</a> relative intensity of a single peak across samples. Such within-peak comparisons may be repeated independently for each peak of interest in a given dataset. For example, Merder et al. (2021) quantified temporal dynamics of individual <a href="https://linearchys.org/li

#### **5 Quantitative impacts**

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### 5 Ecological metrics using peak intensities are often robust

The previous sections show that highlight challenges in connecting between-peak changes in peak intensity do not accurately reflectto between-peak changes in abundance (Fig. 4). This violates a fundamental These challenges violate an assumption of abundance-based ecological analyses: proxies offor abundance (e.g., peak intensity) must reflect actual abundance. In turn, it is tempting to infer that mass spectrometry peak intensities cannotshould be used at all in ecological analyses proportional to true abundances. However, the impacts of violating the assumption have not been directly quantified. This is a significant gap considering quantitative impacts of this situation likely vary across ecological metrics and with study details. There may be certain metrics or situations in which robust inferences can be made despite poor linkages between peak intensities and true abundances. These cases are important to understand, especially given the growing number of publications that useusing peak intensities to compute abundance-based ecological metrics—over the last couple decades.

Therefore, there is a need to quantitatively understand biases and uncertainties introduced in ecological metrics  $(e.g., \alpha \text{-and }\beta \text{-diversities}) \text{ and/or models when peak intensity does not reflect abundance or concentration.} To$ provide an initial evaluationguidance on best practices for using FTMS peak intensities with ecological metrics, we developed an in silico simulation model that. This model generates synthetic data, introduces specific kinds of error commonly found with HRMS datasets (discussed above in detail), errors that degrade the linkage between peak intensity and true abundance, and computes within-sample (e.g., Shannon diversity) and between-sample (e.g., Bray-Curtis) ecological metrics (Fig. 6). This The model allows for comparisonus to probe how the introduction of each type of error impacts the relationship between true values of the metrics and the values observed after each type of error is introduced, which is impossible to do with non-simulated datasets values. To generate synthetic data, we randomly assigned abundances to either 100 or 1000 peaks. Abundances were sampled with replacement from a Gaussian distribution that varied in mean and standard deviation across synthetic samples and across simulation iterations. Abundances were drawn twice to generate two independent samples per simulation, and the simulation was run 100 times for each number-of-peaks (100 or 1000 peaks per sample; referred to below as 'peak richness'). The reason for variation in We varied the Gaussian distributions was to generate synthetic samples that variedvarying in composition within and across simulations to ensure that the ecological metrics (see below) would vary across simulations. This step was necessary to evaluate how biases in the metrics varied metric performance across a broad range of metric values.

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We simulated two types of error, and which can both ean be representative of variation in ionization efficiency. The goal was to generate synthetic data that mimicked our empirical and theoretical observations that indicate noise in the sense that relationships between observed peak intensities did not reflectand true abundances. For each type of error and within each iteration of the simulation, the error was introduced 100 times (i.e., 100 error iterations were nested within each sample-generation iteration). The first type of error was designed to diminish the between-peak relationship between observed peak intensity and true abundance. For To introduce this error, we multiplied the true abundance of each peak by a random number drawn from a uniform distribution ranging from 0 to 100. The inclusion of 0 indicates situations in which a given peak (i.e., ion) does not ionize well enough to be observed. The results should not be sensitive to the selected range, but as a sensitivity analysis, we also used a distribution of errors ranging from 0 to 8. Our empirical data suggest that this narrower range is appropriate (Fig. 4B), but simulation results were not affected by the selected error range (Supplementary Figs. S3-S8). For each peak we multiplied the same random error toby its abundance in each of the two synthetic samples within each iteration. The This errormodified abundance of each peak in each synthetic sample was considered to be the observed peak intensity. We recognize that randomized errors do not perfectly reflect real-world variation in ionization efficiency. However, because the true impacts of matrix effects and individual molecular chemistries in complex mixtures are currently not known, the errors introduced in the model are simply used to diminish the relationship between observed peak intensities and true abundances.

As expected, introducing error resulted in a relatively weak relationship between observed peak intensity and true abundance; (median  $R^2 = \sim 0.5$ ; see black line in Figure 7), with the amount of error increasing with true abundance (Fig. S1)S1). This relationship additionally supports our inclusion of error into the model as a means to simulate relatively weak relationships between observed peak intensity and a median  $R^2$  of  $\sim 0.5$  (see black line in Figure 7). true abundance. Between-peak differences in observed intensity were also weakly related to between-peak differences in true abundance (Fig. 8A), with a median  $R^2$  of  $\sim 0.5$  (see blue line in Figure 7). Because the same peak-level error-factor was used across both synthetic samples within a given simulation iteration, the within-peak between-sample differences in observed intensity were relatively strongly correlated to within-peak between-sample differences in true abundance (Fig. 8B8C), with a median  $R^2$  of  $\sim 0.75$  (see the gray line in Figure 7). As seen in Figure 8C, the differences collapse when near zero. This phenomenon can be explained by the fact that when two samples have essentially the same peak intensity for a given peak, introducing the same error to that peak in both samples has little influence on the between-sample difference in peak intensity.

The second type of error we introduced represents situations in which there is variation in-ionization efficiency varies across molecules – as in the first type of error – but that ionization efficiency also varies well as across samples. Molecules may varyexhibit variations in their ionization efficiency across samples due to changes in the composition of organic molecules and/or changes in inorganic solutes. In this case, ionization efficiency of any given molecule is due to interactions with other organic and inorganic molecules within a given sample. For this in the matrix (see above). To account for these effects, we multiplied the true abundance of each peak by a random number drawn from a uniform distribution ranging from 0 to 100-; for sensitivity analysis, we also used an error distribution ranging from 0 to 8, which did not have meaningful influences on the results. For each iteration of the simulation this was done, we introduced errors independently for boththe two synthetic samples. In this way, the simulated ionization efficiency for a given peak in a given synthetic sample was independent of its ionization efficiency in the other synthetic sample. In turn, the The error-modified abundance of each peak in each synthetic sample was considered to be the observed peak intensity.

We observed a relatively large influence of on observed peak intensities when allowing ionization efficiency to vary randomly across samples. That is, the within-peak between-sample differences in observed intensity were relatively weakly correlated to within-peak between-sample differences in true abundance (Fig. 8B8C), with a median R<sup>2</sup> of ~0.5 (see the red line in Figure 7). Comparing this Compared to the same relationship that emerged under the first type of error-shows, our results show a much weaker relationship between peak intensity and true abundance when ionization efficiency varies between samples (compare the gray and red lines in Figure 7). This result is expected, as variation variations in ionization efficiency will add random noise to the within-peak between-sample differences in observed peak intensity. We note that the variation in ionization efficiency is independent between peaks for both the first and second types of error. The between-peak relationship summarized in Figure 7 (blue line) is, therefore, equivalent for both types of error, which is also shownfurther supported by the strong similarity between FigureFigures 8A and 8C8B.

To examine influences ofhow both types of error oninfluence ecological metrics, we used the initial true abundances and the error-modified abundances (i.e., observed peak intensity values) to calculate true and 'observed' values of within-sample Shannon diversity and between-sample Bray-Curtis. We also assigned an arbitrary trait value to each peak and calculated true and observed sample-level mean trait values; the mean values for each sample were weighted by true abundance (true mean) or observed peak intensity (observed mean). To evaluate biases This analysis is analogous to the approach commonly used in ecological studies for computing community-level abundance-weighted trait values, such as plant leaf area index or animal body size (Muscarella and uncertainty introduced by both types Uriarte, 2016). This approach is also commonly used with FTMS data, such as sample-level peak-intensity-weighted values of error wehydrogen-to-carbon ratios and molecular weight (Roth et al., 2019; Wen et al., 2021). We regressed observed values for each metrie Shannon diversity, Bray-Curtis, and mean traits against their true values. This was done, and performed this process independently for each level of peak richness to evaluate how bias and uncertainty vary with the number of peaks contained within a sample.

Relating 'observed' values of each metric to their true values revealed that the patterns observed in peak-intensity-based ecological metrics are actually likely to be qualitatively robust, even though despite the existence of quantitative biases do exist (Figs. 9-11). All three ecological metrics showed monotonic relationships between observed and true values. Uncertainty was lower when samples had 1000 peaks, relative to samples with 100 peaks; in Figures 9-11 all A/B and C/D panels have 100 and 1000 peaks, respectively. MonotonieWe observed monotonic relationships and lower uncertainty with more peaks were found for both within-sample and between-sample error; in Figures 9-11 all A/C and B/D panels have within-sample and between-sample errors, respectively. For Shannon diversity, observed values were consistently lower than true values, but all observed vs. true relationships were linear (Fig. 9). For Bray-Curtis, inclusion of between-sample error resulted in an overestimation of values and non-linear (but monotonic) relationships between observed and true values (Fig. 10). For mean trait values, the observed

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values had uncertainty but there were we found no systematic quantitative biases, and the relationships between observed and true values were consistently linear (Fig. 11). Furthermore, the

The variation in observed values explained by true values (via a linear model) increases rapidly with the number of peaks, and sharply asymptotes beyond ~500-1000 peaks per sample (Fig. S2). We and sharply asymptotes beyond ~500-1000 peaks per sample (Fig. S2). Sample-to-sample changes in the value of ecological metrics can, therefore, be interpreted with increasing confidence as the number of peaks increases. Qualitative gradients are, therefore, more robust with more peaks. The absolute magnitude of some ecological metrics, however, are shifted away from their true magnitude even when there are large numbers of peaks (e.g., Fig. 10D). Quantitative comparisons from one dataset to another may, therefore, require further simulation-based evaluation. We further caution that the number of peaks needed to reach the asymptote, thereby minimizing error, is likely dataset dependent, and 500-1000 peaks should not be taken as a general rule- for real-world datasets. We encourage researchers to complete such simulations using the numbers of peaks present across their own real-world datasets to better understand their ability to make statistical and conceptual inferences.

#### 6 Conclusions and Recommendations

There is increasing interest in using ecological metrics with FTMS data to study organic matterNOM chemistry across a broad range of environments and settings. It is vital that this growing body of work be based on rigorous use of the data-to-develop trust in the associated conceptual and mechanistic inferences. To do so, This requires deep understanding of the metrics themselves, full, awareness of the data limitations of the OM data from mass spectrometers, and careful use of the metrics informed by the data limitations. We suggest that studies/publications that use using FTMS peak intensities need to include material that directly discusses the data limitations, what peak intensities do and do not represent (e.g., tree-like vs. bird-like data; Fig. 5), and how knowledge of those limitations was used to select specific metrics.

We have provided both strong theoretical reasoning and empirical observations showing that peak intensities do not directlynecessarily map to concentrations of the associated organic molecules within NOM-like complex mixtures of organic molecules. This is particularly true for between-peak comparisons of intensity, and statistical post-hoc normalizations of peak intensity data do not solve this problem. That is, there are no situations that we are aware of in whichchallenge. We caution against using between-peak differences in intensity indicate between-peak differences in concentration. We therefore assert that between peak differences in intensity within HRMSfrom FTMS data cannot be used to make direct inferences related to between-peak variationdifferences in abundance or concentration. This means that HRMS data are unlikely to provide informativehas implications for some ecological analyses based directly on variation in species abundances. In particular, estimation of 'species abundance distributions' isare likely to be invalidproblematic. Analyses that bin peaks into high and low abundance groups based on between-peak differences in concentration are, likewise, almost certainly invalid, also likely to be problematic. We did not directly evaluate these types of analyses, however, and we suggest that future work should expand upon the ecological metrics examined here via simulation.

While certain there are challenges and limitations in the use of ecological analyses of HRMS metrics with FTMS data-are likely to be invalid, we foundshow that there is a tangible path forward. In particular, our simulation model revealed good performance of some common metrics. These cological metrics were originally designed to use relative abundances of biological species. Our simulation modeling indicated that at least some of  $\alpha$ -diversity,  $\beta$ -diversity, and functional trait metrics are likely to provide valid qualitative patterns. That is, values. We infer that conceptual and mechanistic inferences are likely to be valid when based on analyses such as comparing peak-intensity-based ecological metrics across experimental treatments or variation along environmental gradients. The performance of intensity-weighted mean trait values was particularly good in terms of both qualitative and quantitative aspects. We emphasize that we studied a small set of metrics (Shannon diversity, Bray Curtis, and

intensity weighted trait values) and our inferences only extend to these metrics. Fortunately, it is relatively straightforward to extend the simulation model to additional metrics (e.g., Hill numbers; Hill, 1973) and analyses (e.g., species abundance distributions; McGill et al., 2007) and we). We suggest that users of such datasets wanting to use additionalFTMS data do this before applying abundance-based ecological metrics/analyses test them using simulation models before applying\_to real-world datasets. This will provide objective guidance on how to use (and whether to ascertain if these avoid) specific metrics hold given the known biases in these analyses and metrics for specific FTMS datasets.

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To enable robust use of HRMSFTMS peak intensity data in future studies, we recommend use of and further development of the simulation model developed here. The simulation model is the only tool we are aware of that can provide objective guidance on what analyses are not valid and the levelevaluations of uncertainty and potential biases associated with valid analyses. It should using FTMS peak intensities to compute ecological metrics. The model should not be taken as a static or mature tool, however. The model should be expanded in a number of ways by including We encourage future work to expand it to include additional ecological metrics/analyses, situations with more than two-sample, sample situations-to-sample variation in peak richness, links between peak richness and peak intensity, other ways of modeling error, and measured levels of error between concentrations and peak intensities. This These evaluations are outside of the scope of this work, but will be straightforward to include in future versions of the simulation model. Such additions will allow each study to customize the model for their specific application. It should be possible to include the number of samples, the number of peaks in each sample, the peak intensity distributions, number of replicates, and the specific ecological analyses that will be applied. In turn, simulation model outcomes can provide objective guidance tailored to each study. One may think of the resulting guidance as akin to a power analysis whereby the simulation can indicate what can and cannot be inferred from a given dataset. For example, the model indicates that observed Bray-Curtis values have little to no correspondence to true values when Bray-Curtis is below ~0.2 (Fig. 10B, D). Bray-Curtis near and below ~0.2 are commonly observed in HRMSFTMS studies (e.g., Hawkes et al., 2016; Derrien et al., 2018; Bao et al., 2018), and this disconnect between observations and truth is maintained even with 1000 peaks per sample (Fig. 10D). In turn, HRMSFTMS studies that observe Bray-Curtis below ~0.2 may not be able to use those observations to make valid conceptual inferences. However, quantitative guidance must be developed for each study and we recommend that a version of the simulation model should be used by all future studies using peak intensities to conduct ecological analyses of HRMSFTMS data. It may be that in time we understand the general rules well enough to leave the simulation behind, but for now, failing to use it (or a similar tool) leaves analyses open to criticism and potentially spuriouswe suggest its use is warranted to ensure robust inferences.

In addition to further use and development of the simulation model, we recommend translation of other modeling approaches for use with HRMSFTMS data. Two potential approaches are based in machine learning and hierarchical modeling. Machine learning could be used within very tightly controlled systems to understand the magnitude and nature of non-quantitative biases that disconnect peak intensity from concentration. In this case, one could to model the instrument response for a diverse chemical space in typical environmental samples to learn how measured signal intensities may relate to startingtrue concentrations. Even if such a model does not yield highaccuracy results, it may nonetheless help understand error, /biases, and provide additional guidance for robust use of peak intensity data. Furthermore, such a model would be constrained to the system it was built around, and application outwith this system could be wrong. Potentially in concert with machine learning, hierarchical modeling could be translated from its application in ecological analyses (Iknayan et al., 2014) for use with HRMSFTMS. This approach has been used to model sources of error that lead to variation in detectability across biological species, such as variation in species visibility (e.g., Dorazio and Royle, 2005). In turn, data can essentially be corrected by accounting for the modeled sources of error (Roth et al., 2018), even revealing 'hidden diversity' (Richter et al., 2021). There are likely direct analogs to FTMS data in terms of variation among molecules in detectability due to variation in ionization and molecular interactions discussed in previous sections. Machine learning could be used to understand sources of error and, in turn, inform hierarchical models aimed at improving the mapping between peak

intensity and concentration. If successful, this would increase the quality of information provided by peak intensities in both existing and future datasets, thereby enabling more robust conceptual and mechanistic inferences.

In summary, HRMSFTMS has many strengths and weaknesses just like any analytical platform. Other types of compositional data also contain biases and uncertainties, such as the lack of true quantitation in sequence-based microbiome data (Gloor et al., 2017). Careful use of FTMS peak intensity data informed by objective, model-based guidance can overcome some of its weaknesses. Despite peak intensities notWe encourage further development of the model presented here and inclusion of additional methods developed to address issues that arise in similar data types (e.g., Gloor et al., 2017; Hardwick et al., 2018; Vicira-Silva et al., 2019). While these are important directions, we emphasize that despite peak intensities not necessarily reflecting concentrations, ecological metrics overall appear to perform well. This is likely due to the law of large numbers as HRMSFTMS, especially FTICR MS, datasets often contain 1000 or more peaks per sample. Our simulation results indicate that large numbers of identified peaks allow ecological metrics to essentially track towards their true valuevalues. We are encouraged by this outcome and look forward to further applications of ecological metrics, concepts, and theory to organic matterNOM chemistry.

628 7 Code Availability: R code for running the simulation models is available on GitHub: 629 https://github.com/stegen/Peak Intensity Sims. Python code used to process the empirical da

https://github.com/stegen/Peak\_Intensity\_Sims. Python code used to process the empirical data and to generate the associated figures will be available upon publication.

- 8 Data Availability: Raw and processed data will be made publicly available upon manuscript acceptance.
- 9 Author Contributions: WK contributed to conceptualization, experimental data curation, formal analysis, methodology, software, visualization, writing-original draft, writing-review/editing; AMP contributed to conceptualization, methodology, visualization, writing-original draft, writing-review/editing; CHC and SMC contributed to investigation and writing-review/editing; JE contributed to sample preparation and writing-review/editing; MMT contributed to conceptualization, methodology, writing-review/editing; JH contributed to conceptualization and writing-review/editing; RKC contributed to project administration, conceptualization, experimental data curation, methodology, writing-review/editing; JCS contributed to conceptualization, simulation data curation, formal analysis, funding acquisition, investigation, methodology, software, visualization, writing-original draft, writing-review/editing.
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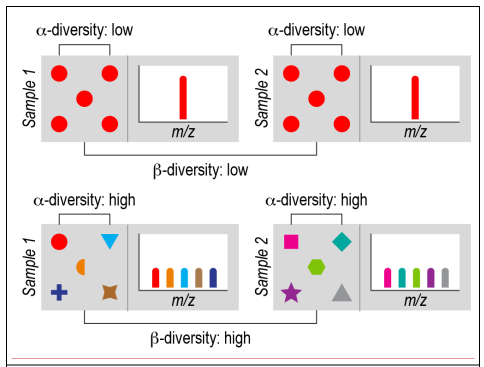


Figure 1. Ecological concepts of α-diversity and β-diversity. Each gray box represents a sample of an ecological community or collection of organic molecules (i.e., an OM assemblage). Symbols represent individual organisms or molecules. Different biological or molecular species are represented by a combination of shape and color. (Top) Each sample has one biological species (red circles) or one chemical species (red bar), and the species are the same within and between the samples. This reflects minimal  $\alpha$ -diversity because there is a single species. This also reflects minimal  $\beta$ -diversity because there is no difference in which species are present in each sample. (Bottom) Each sample has five species (biological or chemical) represented by different colors and symbols. There are no shared species between samples. This reflects maximum  $\alpha$ -diversity because every individual is a different species within each sample, and maximum  $\beta$ -diversity because there are no species shared between samples. In real ecological and OM samples,  $\alpha$ -diversity and  $\alpha$ -diversity fall between these extremes.



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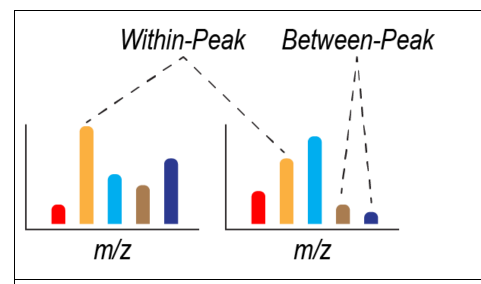


Figure 2. Summary of within-peak and between-peak comparisons of peak intensity. Two idealized mass spectra (i.e., from two samples) are shown with each peak defined by a mass-to-charge ratio (m/z) and represented by a different color. The intensity of each peak in each sample is represented by the height of each colored bar. Within-peak comparisons of intensity are based on comparing intensities at the same m/z across two or more samples. Between-peak comparisons of intensity are based on comparing intensities at two or more m/z values. Between-peak comparisons can be done within a sample (as shown) or between samples (not shown).

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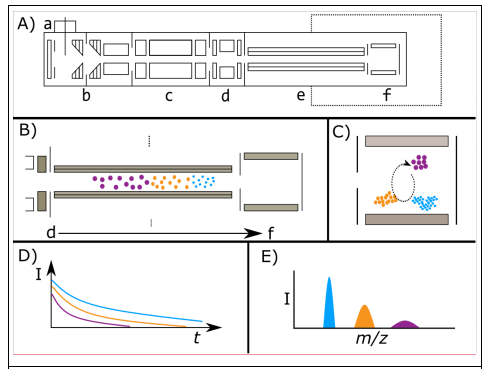
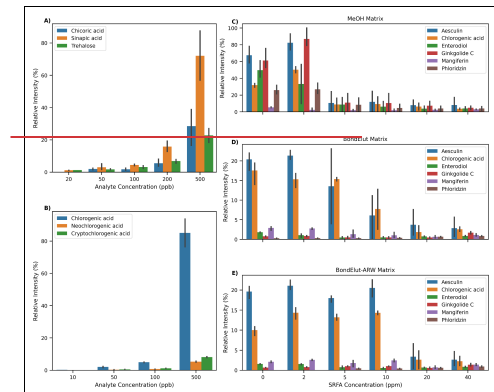


Figure 3. Illustrative example of a generic FTICR mass spectrometer (panel A), showing common and key biases between FTICR signal intensity and m/z of ions (B-E). Panel A shows the major elements of a generic FTICR mass spectrometer (based loosely on a Bruker solariX FTICR MS geometry). Panel A elements include; a - atmospheric pressure ionization source (i.e. ESI source), b - source ion optics (i.e. dual ion funnels), c - mass selecting quadrupole, d - collision cell, e - transfer multipoles to ICR cell, f - ICR cell. Dashed line indicates the magnetic field. Note: diagram is deliberately simplified and not to scale. Panel B) demonstrates the time-of-flight bias along the transfer multipoles (e) in the 'flight tube', from the collision cell (d) to the ICR cell (f). Lower m/z ions travel faster, as indicated by the smaller icons reaching the ICR cell first. Ions are shaded to aid visualization. Panel C) visualizes the effect of a variable excitation radii for ions of different masses, as may happen with a CHIRP excitation pulse. Lower m/z ions are closer to the detection electrodes (shaded in gray) and therefore will induce a larger image current. Note also the ion populations have been adjusted from B) to indicate biases from the time-of-flight effect. Panel D) shows the time-domain recorded signal intensity against time, with the ions having an initial intensity roughly proportional to the number of ions in that cloud. However, as time progresses the less abundant ion clouds lose coherence and destabilize more rapidly, resulting in an attenuation of their signal. Note that the real signal would follow a damped sinusoidal function; here an absolute value approximation is shown for simplicity. Panel E) shows the mass spectrum post-Fourier transform, demonstrating that the impact is not only on intensity (peak height), but also resolution (peak width). In all cases, effects are deliberately exaggerated and not-to-scale to aid interpretation.







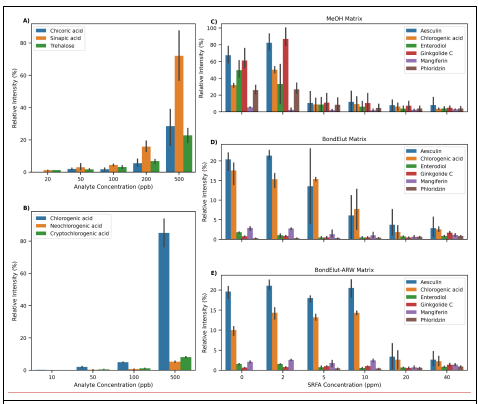


Figure 4 - A) Barplot visualization of the relationship between signal intensity (relative intensity) and concentration of analyte for three chemically distinct molecules analyzed contemporaneously but independently in pure methanol solvent. Relative intensity indicates data were scaled to the largest signal in any replicate from the associated series of spectra. Replicates are combined to show their mean and 95% confidence interval. B) As with A), but for three structural isomers of chlorogenic acid. C-E) Compounds spiked into three different solvent matrices (methanol, BondElut methanol, and BondElut artificial river water (ARW)) at a fixed concentration (100ppb100 ppb), but with addition of SRFA at varying concentrations from 0 to 10ppm40 ppm. In all cases, [M-H]- ion only is shown, but other ions (i.e. [M+CI]-) were detected. 95% confidence intervals represent the results of triplicate measurements. Intensities have been scaled per plot for A and B, and are on the same scale for C-E).

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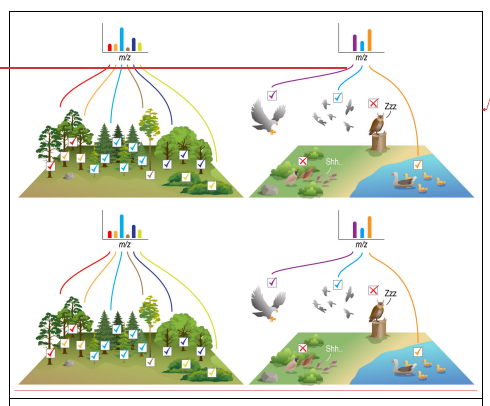
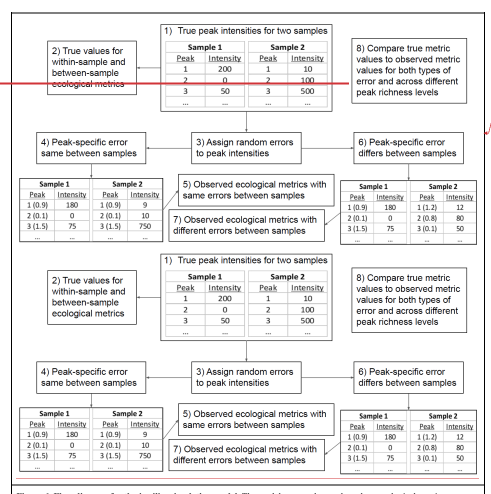


Figure 5. Graphical summary of how <a href="HRMSFTMS">HRMSFTMS</a> peak intensity data are often treated (left), which is distinct from the reality of those data (right). When surveying the number of individuals of each species within a tree community, there is good confidence that the measured abundances are close to real abundances. This is because there is relatively little variation across species in the ability to detect individuals. <a href="HRMSFTMS">HRMSFTMS</a> peak intensity data are often used as though they are like tree-community data. However, <a href="HRMSFTMS">HRMSFTMS</a> data are more like bird-community data. That is, the ability to detect different species varies due to intrinsic factors (e.g., activity patterns, how loud and often birds call, etc.) and extrinsic factors (e.g., habitat structural complexity, predator-induced behavioral changes, etc.). Similarly, the intrinsic physics of a given molecule will impact its ability to ionize and thus its observed peak intensity, and in environmental samples there are thousands of molecular species that impact the ionization 'behavior' of each other. <a href="HRMSFTMS">HRMSFTMS</a> data being more bird-like than tree-like needs to be accounted for when performing ecological analyses using <a href="HRMSFTMS">HRMSFTMS</a> data.



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Figure 6. Flow diagram for the *in silico* simulation model. The model was used to evaluate how ecological metrics are impacted by variation in ionization across organic molecules (i.e., peaks). The true peak intensities are what is expected if intensity is linearly to concentration, and all peaks fall along the same linear function. Variation in ionization adds error around this idealized linear relationship. The error is modeled in two ways: the error applied to a given peak is either the same between samples (i.e., there are no variable matrix effects on ionization) or varies randomly between samples (i.e., there are variable matrix effects on ionization). In the lower tables the proportional error applied to each peak is provided parenthetically. The tables are for demonstration and show only three peaks per sample. The number of peaks per sample was set to either 100 or 1000.



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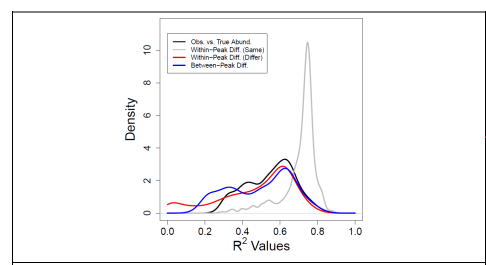
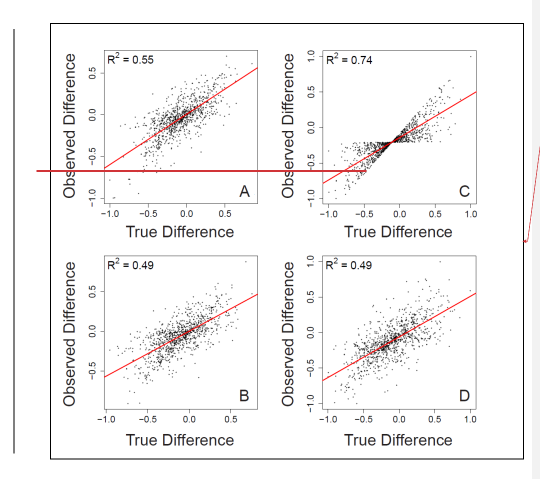


Figure 7. Variation in observed intensity explained by true abundance. Kernel density functions are shown for different relationships and types of error. Density functions were fit using R² values collated from across simulation iterations. Higher R² values indicate a stronger link (i.e., lower uncertainty) between observed intensities and true abundances. Black is for the relationship shown in Figure S1. Blue is for between-peak within-sample differences (example relationships shown in Figures 8A,CB). Gray is for within-peak between-sample differences when the same peak-level error was used for both synthetic samples within a given simulation iteration (example relationship shown in Figure SBSC). Red is for within-peak between-sample differences when different peak-level error was used across the synthetic samples within a given simulation iteration (example relationship shown in Figure 8D). While there are central tendencies in all four distributions, there is also significant variation in the degree to which observed intensities reflect true abundances.







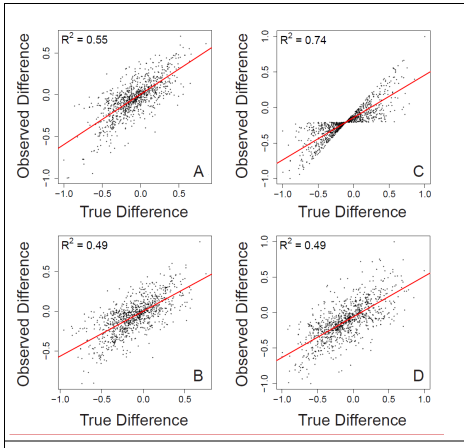
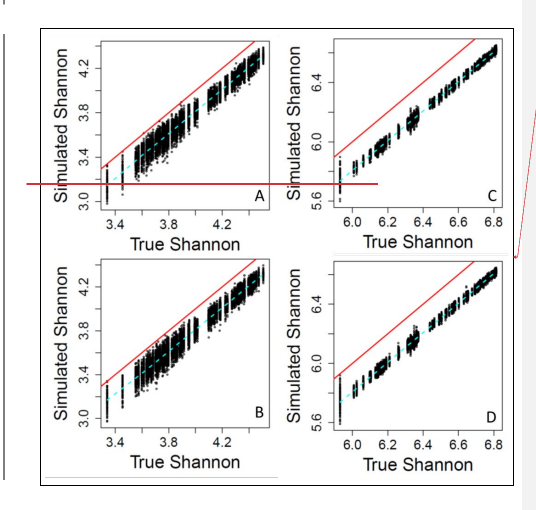


Figure 8. Observed differences in peak intensity as a function of true differences in peak intensity across both within-peak and between-peak comparisons and across both kinds of error. (A) Between-peak differences with the same error applied to a given peak between samples. (B) Within-peak differences with the same error applied to a given peak between samples. (C(B)) Between-peak differences with different errors applied to a given peak between samples. (C) Within-peak differences with the same error applied to a given peak between samples. (D) Within-peak differences with different errors applied to a given peak between samples. On all panels the red line represents the linear regression model, and the associated R<sup>2</sup> value is provided.



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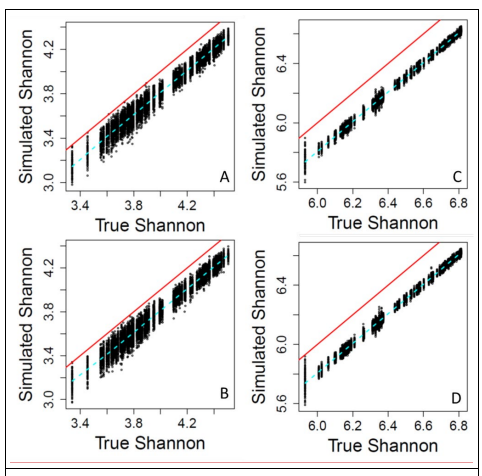
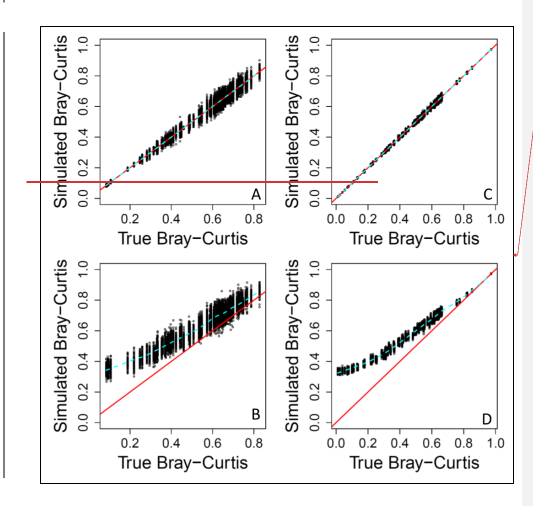


Figure 9. Shannon  $\alpha$ -diversity that includes simulated error regressed against true Shannon, across different scenarios. (A) The same error applied to a given peak between samples, and 100 peaks per sample. (B) Different errors applied to a given peak between samples, and 100 peaks per sample. (C) The same error applied to a given peak between samples, and 1000 peaks per sample. (D) Different errors applied to a given peak between samples, and 1000 peaks per sample. On all panels the red line represents the one-to-one line and the dashed line is a spline fit to the data. All data are from the simulation model.







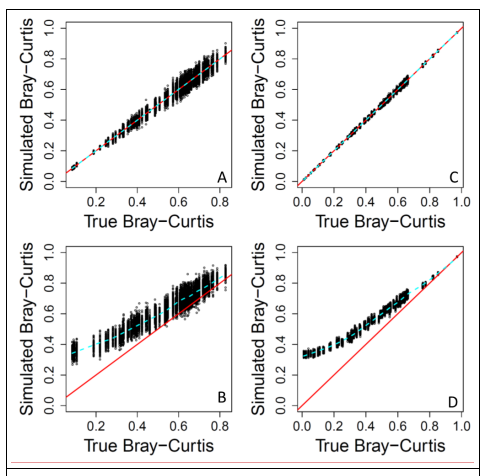
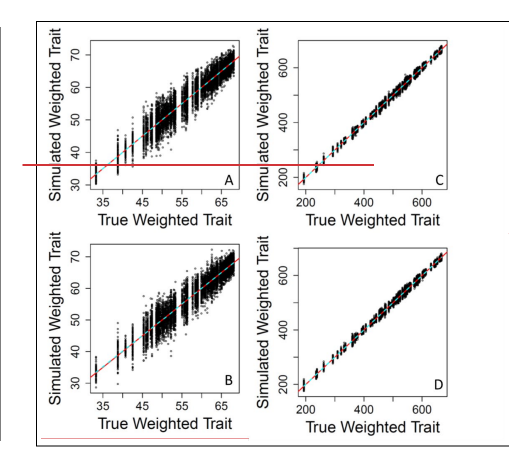


Figure 10. Bray-Curtis dissimilarity as a measure of  $\beta$ -diversity that includes simulated error regressed against true Bray-Curtis, across different scenarios. (A) The same error applied to a given peak between samples, and 100 peaks per sample. (B) Different errors applied to a given peak between samples, and 100 peaks per sample. (C) The same error applied to a given peak between samples, and 1000 peaks per sample. (D) Different errors applied to a given peak between samples, and 1000 peaks per sample. On all panels the red line represents the one-to-one line and the dashed line is a spline fit to the data. All data are from the simulation model.







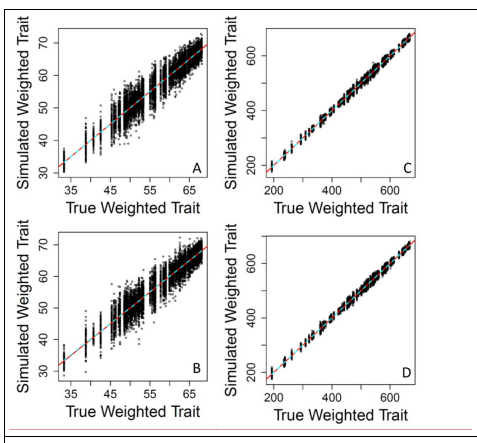


Figure 11. Mean peak-intensity-weighted trait values that include simulated error regressed against true mean peak-intensity-weighted trait values, across different scenarios. (A) The same error applied to a given peak between samples, and 100 peaks per sample. (B) Different errors applied to a given peak between samples, and 100 peaks per sample do a given peak between samples, and 1000 peaks per sample. (D) Different errors applied to a given peak between samples, and 1000 peaks per sample. On all panels the red line represents the one-to-one line and the dashed line is a spline fit to the data. All data are from the simulation model.