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Introduction

Recent technological breakthroughs in high-resolution mass spectrometry (HRMS) have made it an increasingly popular technology in environmental analysis,1 biomonitoring2,3 and metabolomics.⁴ Often combined with suspect screening (SS) and non-targeted analysis (NTA) workflows, HRMS has shown great promise in the discovery of lesser-known chemical structures and in comprehensively characterizing the chemical composition of complex mixtures.^{2,3,5} One critical challenge associated with the application of HRMS in SS or NTA is the limited availability of analytical standards for many anthropogenic/synthetic chemicals and many endogenously produced metabolites.6,7

Modeling the relative response factor of small molecules in positive electrospray ionization*

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Technological advancements in liquid chromatography (LC) electrospray ionization (ESI) high-resolution mass spectrometry (HRMS) have made it an increasingly popular analytical technique in non-targeted analysis (NTA) of environmental and biological samples. One critical limitation of current methods in NTA is the lack of available analytical standards for many of the compounds detected in biological and environmental samples. Computational approaches can provide estimates of concentrations by modeling the relative response factor of a compound (RRF) expressed as the peak area of a given peak divided by its concentration. In this paper, we explore the application of molecular dynamics (MD) in the development of a computational workflow for predicting RRF. We obtained measurements of RRF for 48 compounds with LC - quadrupole time-of-flight (QTOF) MS and calculated their RRF. We used the CGenFF force field to generate the topologies and GROMACS to conduct the (MD) simulations. We calculated the Lennard-Jones and Coulomb interactions between the analytes and all other molecules in the ESI droplet, which were then sampled to construct a multilinear regression model for predicting RRF using Monte Carlo simulations. The best performing model showed a coefficient of determination (R^2) of 0.82 and a mean absolute error (MAE) of 0.13 log units. This performance is comparable to other predictive models including machine learning models. While there is a need for further evaluation of diverse chemical structures, our approach showed promise in predictions of RRF.

> When it comes to endogenous metabolites, the lack of commercially available analytical standards is often due to certain compounds being less well-characterized and thus not yet synthesized or purified. For anthropogenic chemicals, one of the reasons is that chemical manufacturers in the U.S. are not required to produce analytical standards for the chemicals that they manufacture and release to the environment.7 One exception to this rule is pesticides.⁷ It is important to note at this stage that the requirement for analytical standards does not extend to the transformation and breakdown products of these chemicals. So even in a hypothetical scenario where manufacturers would be required to produce analytical standards, that would cover only the parent compounds and not all the transformation products. The U.S. Environmental Protection Agency (EPA) has prioritized about 1.2 million chemicals of environmental importance and has created a database called EPA's CompTox Chemicals Dashboard (henceforth referred to as "the dashboard").8 Nuñez et al.6 estimated that out of the 1.2 million chemicals on EPA's Dashboard, less than 2% are available as analytical standards.

> There is thus a need to develop computational approaches to confirm and quantify detected compounds without analytical standards.6,9,10 While detection and tentative confirmation can be achieved with MS/MS libraries, quantification remains a more challenging task.^{11,12} Liquid Chromatography (LC) -

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Paper

Electrospray Ionization (ESI) HRMS is one of the most commonly used HRMS techniques in SS and NTA studies. One critical challenge in ESI is that abundances expressed as peak areas or peak heights are not easily translatable to concentrations. Two compounds of the same concentration can exhibit peak areas that differ by 3 orders of magnitude because of differences in ionization.12,13 Abundances may be used as a surrogate for concentrations in certain situations when comparing the same chemical across different samples, however, they cannot be used to compare two chemicals to each other.14 We should note at this point that while we focus on HRMS in our paper, low-resolution MS such as triplequadrupole instruments can also be used to study the ionization efficiency of chemicals. We focus primarily on HRMS and NTA because that is when one often meets with the lack of available analytical standards and when predictive models can help circumvent that problem.

While ESI is extensively used in mass spectrometry for the analysis of both small (*e.g.*, metabolites) and large molecules (*e.g.*, proteins), the precise mechanism has not been fully understood. Briefly, during ESI, the solution containing the analyte passes through a metal capillary that is charged at an electric potential of thousands of volts (kV). The solution forms a tip at the end of the capillary known as a Taylor cone that emits a spray of fine droplets. The droplets start in the μ m range and shrink in size as they undergo evaporation often accelerated with heating of the capillary. The density of the charged ions in the droplet is controlled by repulsive coulombic forces between positively charged ions. The upper limit of that density is described as the Rayleigh stability limit:^{15,16}

$$z_{\rm R} = \frac{8\pi}{e} \sqrt{\varepsilon_0 \gamma R^3} \tag{1}$$

where, *e* is the number of elementary charged particles, ε_0 is the vacuum permittivity, γ is the surface tension of the droplet and *R* is the droplet radius.

Conceptual models have been proposed to describe the process that molecules undergo to become ionized and transferred to the gas phase during ESI. We focus our discussion on the commonly used positive ESI under which positive ions are formed. Small molecules (<1000 Da) are thought to ionize and be transferred to the gas phase by the ion evaporation model (IEM).¹⁵ According to IEM, the analyte is protonated already while inside the droplet and eventually moves from the center towards the surface of the droplet. As the positively charged analyte meets the positively charged solvent molecules on the surface of the droplet, the ion is transferred to the gas phase through repulsive forces of positively charged ions and by the excess droplet charge.

Larger molecules such as globular proteins (natively folded proteins) are thought to ionize and be transferred to the gas phase by the Charged Residue Model (CRM).^{15,17} According to CRM, solvent droplets containing a single protein molecule gradually evaporate to dryness and as the solvent molecules evaporate, the charge is transferred to the analyte. The droplets remain close to the Rayleigh stability limit while evaporating, which indicates that the droplet loses some of the electric charge as it shrinks in size. This is supported by experimental

observations where the size of solvent droplets positively correlated with the droplet charge following an exponential curve. Contrary to small molecules, the ejection of globular proteins is not kinetically favorable. The repulsive forces of the excess surface charge are not sufficiently strong for the molecule to be ejected and transferred to the gas phase. CRM is supported by experimental evidence where ionization of globular proteins produces ions with a charge of $[M + z_R H]^{z_R+}$, where z_R is the charge at the Rayleigh limit of water droplets of the same size as the globular protein.¹⁵

While these two models are considered distinct, both of these two processes could apply to small or medium-sized molecules, especially in the case of heated ESI where the evaporation of the water droplets is assisted through heating of the capillary. This is also supported by molecular dynamics (MD) simulations studies where native (unmodified) carbohydrates ionize through CRM, while their permethylated derivatives ionize through IEM.¹⁸

Ionization efficiency is a property that has proven to be difficult to predict from a small number of chemical descriptors or physicochemical properties. Numerous investigations have examined the correlation between ionization efficiencies and diverse physicochemical properties, including but not limited to their pK_a , log P, surface area, charge delocalization, and gasphase proton affinity.19-24 These findings have led to the development of various predictive models for ionization efficiencies,^{22,25-28} utilizing both the physicochemical properties of analytes and solvent characteristics as fundamental inputs. These models commonly rely on parameters associated with the analyte's hydrophobicity (e.g., log P, WAPS, WANS, C/H ratio) and ionizability (such as pK_a and the degree of ionization).²⁹ Data-driven approaches involving machine learning, such as random forest models, have shown great promise in providing predictions with reasonable uncertainties.11,12,30-32 It should be noted, however, that these approaches require large datasets in the order of hundreds of chemicals with diverse structures and properties, and the predictions are tied to a specific method and instrumentation.

Theory-driven approaches that are based on quantum chemistry and computational chemistry principles could provide an alternative when large datasets are not available or are difficult to obtain. Molecular dynamic simulations (MD) have been previously applied in a number of studies to understand the mechanism of ionization in salt ions, peptides, and proteins³³⁻³⁷ and to a lesser extent in small molecules^{18,38} However, to the best of our knowledge, there appears to be little on the predictions of their ionization efficiency using MD simulations. Our study aims to fill this gap by employing molecular dynamics to model the behavior of chemicals in the ionization chamber and evaluate the potential of such theory-driven approaches to make predictions and assess their uncertainties.

Materials and methods

Workflow diagram

The individual steps of the experimental and computational aspects of the study are presented in Fig. 2.

Experimental section

Chemicals and solutions. The analytes were provided by the US EPA for the purposes of this study and were developed as part of EPA's Non-Targeted Analysis Collaborative Trial (ENTACT). The preparation of the chemical mixtures is described in detail in the study of Ulrich et al.39 For the purposes of this study, we used mixtures 504, 506, and 508. The mixtures were diluted in a series of dilutions first with methanol (99.9% Millipore Sigma) and then with HPLC water (99.9% Millipore Sigma) from 20 mM to 100, 50 and 25 μ M with a final water content of >99%. For the purposes of this study, we selected chemicals that ionize in positive electrospray ionization mode (ESI+) and whose calibration curves showed an R^2 of 0.8 or higher. The chemical structures and chemical identifiers of the chemicals involved in the study (n = 48) are shown in ESI spreadsheet 1.[†] The complete chemical lists from mixtures 504, 506, and 508 are also shown in ESI spreadsheet 1.[†] We should note at this point that characterizing the mixtures and maximizing the number of detected and identified compounds is beyond the scope of the study. As our study requires extensive computations that take days to complete, we have to limit our efforts to a small subset of compounds that satisfy the criteria of detection (<5 ppm mass difference from the monoisotopic mass) and linearity ($R^2 \ge 0.8$).

Instrumental analysis. The mixtures were analyzed with an Agilent 1290 ultra-high performance liquid chromatography (UPLC) coupled to an Agilent quadrupole time-of-flight (QTOF) mass spectrometer. The UPLC was equipped with an Agilent Eclipse Plus C18 column (2.1 \times 100 mm, 1.8 μ M) for the chromatographic separation of the analytes. The mobile phase consisted of the two following solutions. Solution A: HPLC water (Sigma-Aldrich, \geq 99.5%) with 0.1% methanol (Sigma-Aldrich, 99.9%) and 5 mM ammonium acetate (Sigma-Aldrich, \geq 98%). Solution B: 90% methanol with 10% HPLC water and 5 mM ammonium acetate. The two solutions were mixed under the following gradient program: 0 min 10% B and 90% A, 0-15 min increase to 100% B, 16-20 min equilibration at 100% B. The solvent gradient over time is also shown in Fig. S1.[†] All mixtures were injected twice at an injection volume of 5 µL. Two no-injection blanks and one HPLC water blank were also analyzed in the beginning of the sequence.

The instrument was operated in both positive electrospray ionization mode (ESI+) and full scan mass spectra (MS1) were acquired in the range of 100–1000 Da with a resolving power of 40 000 and a mass accuracy of <5 ppm. The instrument was calibrated before the analysis and the mass difference was corrected with reference standards using masses 121.050873 and 922.009798 for positive ionization mode.

Data collection and file processing. All the collected data files were processed with MS-DIAL, an open-source software for mass spectrometry that was developed by the University of California, Davis, and by RIKEN (Japan). The detected features were aligned across samples and were matched to the monoisotopic masses of the chemicals contained in the mixtures within a 10 ppm mass difference (ESI spreadsheet†). The peak areas of the analytes were calculated by taking the average of the duplicate injections and they were corrected by subtracting the average area measured in the blanks. The MS-DIAL settings and parameters used to process the data files are presented in the ESI spreadsheet.[†]

Calculation of RRF. Ionization efficiency describes the extent to which molecules of an analyte in the liquid phase can transition to the gas phase as ions during the process of electrospray ionization. The ionization efficiency of an analyte can be mathematically described by the relative response factor of the analyte as follows:

$$RRF = \frac{A}{C}$$
(2)

where, RRF is the relative response factor, A is the abundance (peak area or peak height) and C is the concentration of the analyte.

Molecular dynamics simulations

Input generation. We generated mol2 format files for all chemicals in the dataset using UCSF Chimera and SMILES as inputs for the mol2 files. The protonation of each molecule was determined by generating pKa diagrams for each chemical using Chemaxon and Chemicalize⁴⁰ and identifying the dominant species at the pH that would be relevant to our experiments (pH = 5).⁴¹ All the pKa diagrams and the protonation states are uploaded as .png images on GitHub under https://github.com/dimitriabrahamsson/electro-spray.

Topology generation. Topologies were generated with CGenFF force field (CHARMM General Force Field) using the CHARMM-GUI⁴² online platform (https://www.charmm-gui.org/) and the mol2 files from the previous step. The protonation of the analytes was examined once more to ensure that it was correct and that no changes were made while importing the mol2 files in GHARMM-GUI. The generated files from CHARMM-GUI were then converted to GROMACS format using the charmm2gromacs-pvm.py script (uploaded on GitHub under https://github.com/dimitriabrahamsson/electro-spray).

System preparation. All preparation steps were conducted using GROMACS (version 2023.2). GROMACS uses periodic boundary conditions (PBC) where the atoms of the simulation system are put into a space-filling box, which is surrounded by translated copies of itself. Thus, the system does not have finite borders during the simulation, but it allows for the removal of PBC for post-simulation calculations.43 Two different systems were considered for the MD simulations. System 1 aimed at approximating the composition of a nm electrospray droplet which included the analyte, the H⁺ produced during hydrolysis, and the water and methanol molecules as shown in the first step of IEM in Fig. 1. With the second system (System 2), we aimed to approximate the composition of the droplet surface and the analyte located at the surface before evaporation as shown in the second step of IEM in Fig. 1. One critical challenge when describing both systems is describing the H⁺ ions in solution. In water, the H⁺ ions, also referred to as H_(aq)⁺, produced from the hydrolysis of water molecules react with other water molecules to form hydronium, $H_3O_{(aq)}^+$, also known as oxonium, following the reactions below:

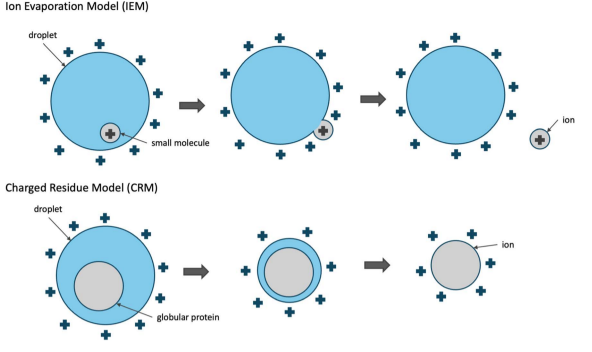


Fig. 1 Conceptual models describing the mechanism of electrospray ionization. Small molecules are thought to be ionized through the ion evaporation model (IEM),¹⁵ while larger molecules, such as globular proteins are thought to ionize through the charged residue model (CRM).¹⁵

$$H_2O_{(l)} \rightleftharpoons H_{(aq)}^{+} + OH_{(aq)}^{-}$$
(3)

$$H_2O_{(l)} + H_{(aq)}^{+} \rightleftharpoons H_3O_{(aq)}^{+}$$
(4)

System 1. The hydronium ion was described using a TIP3P that included additional hydrogen (3H in total) and was modified to include the specific parameters for H_3O^+ from the study of Wolf and Groenhof.⁴⁴ The distance between O and H (r_{OH}) was set at 1.02 Å, the angle for H–O–H (θ_{HOH}) was set at 112°, and the charges for O (q_O) and H (q_H) were set at –0.59 and 0.53*e*. The droplet was represented by a three-dimensional cube and the size was set at 64 nm³ (4 × 4 × 4 nm). The number of H_3O^+ molecules was approximated based on the experimental observations of Smith *et al.*⁴⁵ who determined the charge (*e*) of water and methanol droplets in ESI+ as a function of the droplet diameter. The calculations are described in detail in Text S1 in ESI.[†]

The numbers of water (TIP3P) and methanol molecules were determined based on the gradient mixing (Fig. S1[†]) of the two solvents during LC and based on the retention time of each chemical (ESI spreadsheet[†]). This means that for every chemical the number of water and methanol molecules was different depending on when it was eluted from the LC column. Previous MD studies⁴⁶ on ESI droplets have also suggested that the amount of methanol in the droplet plays a critical role in the ionization efficiency of the analytes. As the volume of methanol increases, the evaporative rate increases, as does the ionization efficiency, for many molecules.⁴⁶

System 2. As mentioned earlier, System 2 aimed at approximating the behavior of the analyte on the surface of the droplet

where most of the charged ions are expected to be located and where the $[M + H]^+$ ions are likely formed. While the presence of hydronium is well established, there are important differences between how the proton (H^+) is bound to H_2O in H_3O^+ and how it is bound in $[M + H]^+$. In the case of H_3O^+ , the proton is bound to the oxygen atom in a covalent bond. Oxygen in H₂O has two lone pairs of electrons. When the extra proton attaches, it forms a dative (or coordinate) covalent bond with one of these lone pairs, where both electrons in the bond come from the oxygen atom. In this covalent bond, oxygen shares one of its lone pairs with the extra hydrogen, creating a stable bond within the hydronium ion.⁴⁷ In the case of $[M + H]^+$, however, H⁺ does not form a covalent bond with the analyte (M). Instead, it is stabilized through ionic interactions. The proton attaches itself to a site on the molecule where it can stabilize the positive charge, typically near a region with lone pairs (such as nitrogen or oxygen atoms) or near a π -system (in the case of aromatic molecules).15,48 In order to account for this discrepancy, in System 2, we describe the H⁺ ions as freely floating ions that are not covalently bound to water molecules or to the analyte. In this case, H⁺ was described in the same way as other ions like Na⁺ and Cl⁻ are described in GROMACS using the CGenFF force field. In this description, the mass of H⁺ was set at 1.0080 g mol^{-1} and the charge (q) was set at +1. As a point of reference, Na⁺ ions in CGenFF are described as single atoms with a mass of 22.98977 g mol⁻¹ and a charge of +1. A 4 \times 4 \times 4 nm solvent box was created with approximately 600 water molecules (TIP3P) and 600H⁺ ions. The number of 600H⁺ was determined based on pilot simulations so that H⁺ would remain evenly distributed inside the box throughout the simulation to ensure continuous interactions with the analyte. A smaller number of

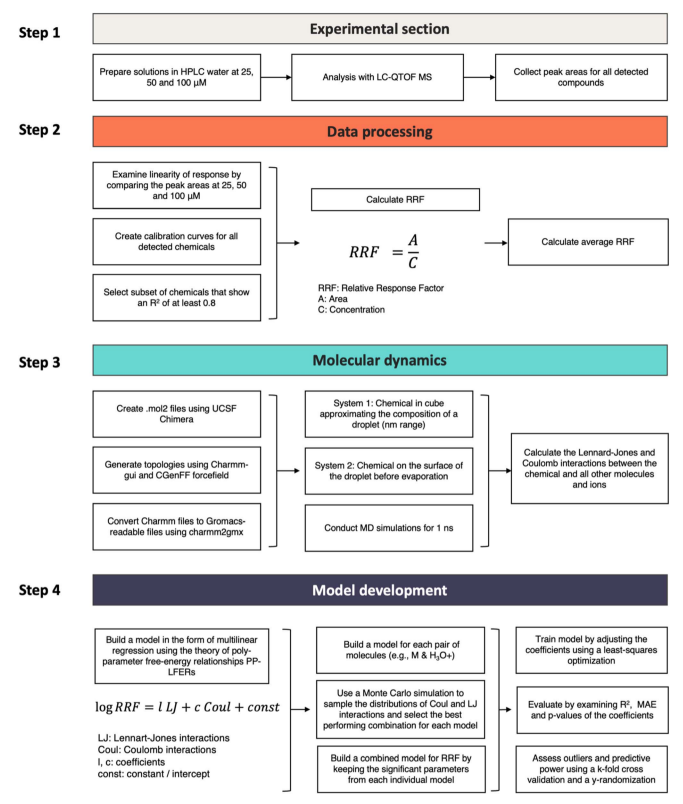


Fig. 2 Workflow diagram for the processing steps in the experimental and computational parts of the study.

ions resulted in the ions starting evenly distributed but during the simulation moving to the outer parts of the box and not sufficiently interacting with the chemical which often remained towards the center of the box (Fig. S3–S5 and Text S2†). **Simulation setup.** The simulations were conducted using GROMACS version 2023.2. The simulation protocol started with the steepest descend minimization with 50 000 steps as the maximum number of minimization steps to perform and

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<1000 kJ mol⁻¹ nm⁻¹ as the threshold at which the minimization process can stop. The minimization and subsequent simulation steps were run using Verlet as the cut-off scheme for neighbor searching and Fast Smooth Particle-Mesh Ewald electrostatics (FSPME or PME in GROMACS) for modeling the electrostatic interactions. The short-range electrostatic cut-off points for Coulomb and van der Waals interactions were set to 1.2 nm which is recommended for CGenFF.49 The temperature was set at 300 K and it was controlled with a Berendsen thermostat in NVT and a Parrinelo-Rahman barostat in NPT. System equilibration was conducted in two stages, the NVT stage where volume and temperature were kept constant, and the NPT stage where pressure and temperature were kept constant. The simulation step was set at 0.5 fs using a leapfrog integrator and the simulation length was 200 ps. The production step following equilibration was conducted using the same simulation step and integrator as previously but in this case, the simulation length was 1000 ps (1 ns). All the mdp files for the minimization, equilibration, and production steps with all the details are available on GitHub (https://github.com/ dimitriabrahamsson/electro-spray).

Calculation of interactions and model development. For both System 1 and System 2, we calculated the short-range Lennard-Jones and short-range Coulomb interactions between the analyte and each group of molecules in the system. For System 1, the sets were (i) analyte and water, (ii) analyte and methanol, and (iii) analyte and H_3O^+ ions. For System 2, the sets were (i) analyte and water and (ii) analyte and H⁺ ions, however, we only considered the set of analyte and H⁺ ions since the interactions with water were already described in System 1. As these are short-range interactions, it is important to point out that this includes only the molecules around the analyte that are within the short-range distance, which was set at 1.2 nm. The interactions were calculated using the gmx energy command in GROMACS. The generated files contained the interaction energies (kJ mol⁻¹) over time (ps) in the form of a time series. An example of the Coulomb and Lennard-Jones interactions for caffeine is shown in Fig. 3. The top figures show the interactions over time and the bottom figures show the distribution of the observed interaction energies using the kernel density estimate.

Our model is based on the idea that the RRF of a given compound in ESI+ can be described as a function of the

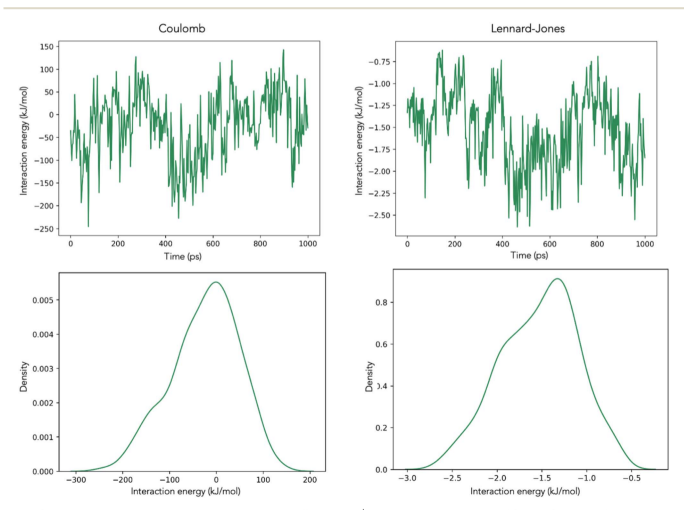


Fig. 3 Coulomb and Lennard-Jones interactions between caffeine and H^+ ions during the simulation of System 2. The top figures show the interactions over time and the bottom figures show the distribution of the observed interaction energies.

Coulomb and Lennard-Jones interactions between the compound and all other molecules in the solution. RRF was expressed as a function of the Coulomb and Lennard-Jones using a multilinear regression model.

The model was as follows:

$$\log RRF = lLJ + cCoul + const$$
(5)

where, LJ is the Lennard-Jones interactions, and Coul is the Coulomb interactions.

One critical challenge when incorporating these interactions into a model is finding which metrics are meaningful for the purposes of the model. We applied a Monte Carlo simulation approach to randomly sample the Coulomb and Lennard-Jones distributions (3 percentile points per distribution plus the standard deviation) 100 times for each set of molecules (*i.e.*, System 1: (i) analyte and water, (ii) analyte and methanol, and (iii) analyte and H_3O^+ ions; System 2: analyte and H^+ ions) and selected the best-performing model for each set.

Expanding the *l*LJ and *c*Coul terms of the equation we get:

$$lLJ = l_1LJ_{p1} + l_2LJ_{p2} + l_3LJ_{p3} + l_4LJ_{std}$$
(6)

$$c\text{Coul} = c_1\text{Coul}_{p1} + c_2\text{Coul}_{p2} + c_3\text{Coul}_{p3} + c_4\text{Coul}_{\text{std}}$$
(7)

where, *p*1, *p*2, and *p*3 are the 3 percentile points from the distribution (*e.g.*, 20, 50, and 70) and std is the standard deviation of the distribution.

The coefficients and the intercept of the model were determined through a least-squares minimization using the statsmodels package (version 0.14.0) in Python (version 3.10.11). The GitHub is available on (https://github.com/ script dimitriabrahamsson/electro-spray). The model was evaluated by examining the R^2 , the mean absolute error (MAE), and the *p*-values of the coefficients. After selecting the best-performing model for each set of molecules from both systems, we built a composite model with the parameters whose *p*-values were lower than 0.1. We purposely chose a higher cutoff point at this stage in order to be more inclusive, however, in the final model, only the p-values below 0.05 were considered significant. The final model was evaluated based on its R^2 , the mean absolute error (MAE), and the *p*-values of the coefficients. We also tested whether the addition of other physicochemical properties, *i.e.* vapor pressure (P_V) , water solubility (S_W) , the equilibrium partitioning ratio between octanol and water (K_{OW}) , air and water (K_{AW}) , methanol and water (K_{MW}) , methanol and air (K_{MA}) , and the innate charge of the molecule improved the predictive accuracy of the model. $P_{\rm V}$ and $S_{\rm W}$ were calculated with OPERA 2.6 (ref. 50) available on the dashboard.⁸ K_{OW} , K_{AW} , K_{MW} , and $K_{\rm MA}$ were calculated with UFZ-LSER.⁵¹ The innate charge of the molecule was determined by examining the structure and its protonation state at pH 5 and noting 0 if it was neutral, +1 (or more) if it had a positive innate charge, and -1 (or less) if it had a negative innate charge. The properties were tested iteratively by adding each property to the model and recording its performance. Only one property was tested at a time and only the properties whose coefficient showed a *p*-value of less than

0.05 were considered significant and were included in the model. A parameter with a mere increase in R^2 without a significant *p*-value would not be included in the model.

The predictive power of the model was further evaluated with a 10-fold cross-validation and a y-randomization. During the 10fold cross-validation, the dataset was first divided into 10 equally sized sub-datasets. Then, during each fold one dataset was set as the testing set and the remaining sub-datasets were compiled into a training set. The model was trained on the training set and tested on the testing set. The process was repeated 10 times (10-fold). It is important to note at this point that when applying a k-fold cross-validation and when dividing the dataset into training and testing there is always a possibility of encountering compounds in the testing set that are outside the applicability domain of the trained model. In order to account for this discrepancy, if a prediction was 2 log units higher than the highest value in the dataset or 2 log units lower than the lowest value in the dataset it was considered outside the applicability domain and it was excluded from the evaluation. The compounds that were excluded from any particular fold of the cross-validation exercise were still included in the discussion section of the paper. The purpose of the k-fold crossvalidation is to evaluate the predictive power of the model outside its training set and to identify outlier compounds in the dataset. These compounds are considered outliers in the sense that they represent physicochemical properties that are dissimilar to the ones in the training set and in order for the model to make accurate predictions, they have to be included in the training set.

For the *y*-randomization, the *y* variable, in this case the RRF was randomly shuffled, and the model was developed as previously by dividing the dataset into training and testing sets. The process was repeated 5 times, and the predictions were averaged across the 5 iterations. The purpose of the *y*-randomization is to evaluate the extent to which the model predictions are different from random predictions. This helps to determine whether the model is making meaningful predictions and whether it has been overparametrized. The lower the R^2 of the *y*-randomization and the more different it is from the R^2 of the cross-validation, the higher the likelihood that the model is making meaningful predictions that are distinct from random predictions.

One of the challenges we encountered is that the generated CGenFF topologies often included high penalties (>50) for a charge, a bond, an angle, a dihedral, or an improper group. While high penalties do not necessarily mean large errors, they do denote a low similarity with the build-by-analogy structure in CGenFF and it is recommended to apply caution when using such structures because they may require further validation. This may be an important issue in the case of protein dynamics and ligand binding, however, in our case, it is unclear how these penalties or uncertainties may influence our calculations. To address this issue, we tested the robustness of the model by incrementally removing compounds with high penalties starting from the ones with the highest penalties to the ones with the lowest penalties. This resulted in 10 different models with a different cutoff point as the maximum acceptable penalty

ranging from 500 to 50. We then examined the changes in the R^2 of the model as the number and type of chemicals in the dataset changed. In order to avoid introducing errors in the first steps of the model development, we developed the first iteration of the model with chemicals that had a penalty of less than 300.

Results and discussion

Experimental measurements

The observed log RRF values for the chemicals in our dataset ranged from 1.73 to 3.17 with Cinchophen showing the lowest value and Thiabendazole showing the highest value (ESI spreadsheet[†]). As RRF is the ratio of abundance to concentration, higher RRF values indicate higher ionization efficiency (higher abundance at lower concentrations). This observation is in agreement with data from our previous study12 where Cinchophen showed a lower RRF compared to Thiabendazole. It should be noted that the two studies use the same mixtures (in part) but different methods and different instruments (same type - Agilent LC-QTOF-MS - but different instrument). Despite the differences in methods and instrumentation, the differences in the RRF values of the two chemicals are preserved. This observation is supportive of the ionization efficiency (IE) scale approach developed by Oss et al.25 where a set of RRF values can be represented as a scale of relative ionization efficiencies and that scale should in principle be transferable across different methods.

Model development

From the models developed for System 1, the best-performing models for predicting log RRF showed an R^2 of 0.42 when using the analyte-water interactions, 0.37 when using the analyte-methanol interactions, and 0.39 when using the analyte- H_3O^+ interactions (ESI spreadsheet[†]). From the models developed for System 2, the best-performing model for predicting log RRF showed an R^2 of 0.71 (ESI spreadsheet[†]). This observation indicates that the final stage of ionization $[M + H]^+$ is better described by the interactions of the analyte with the H⁺ ions on the surface of the droplet (Fig. 1 - step 2 of IEM) than by the interactions of the analyte with the other molecules while in the center of the droplet (Fig. 1 - step 1 of IEM). This is not to say that there is no predictive value in the interactions of the analyte with the solvent molecules. Previous studies have demonstrated the impacts of different solvents on the ionization efficiency of small molecules^{52,53} and this is in agreement with our calculations from System 1. This observation just indicates that the interactions of the analyte on the droplet are potentially more determining the ionization efficiency of the analyte. The final composite model consisted of the following parameters. System 1: p2, p3 and the standard deviation for Lennard-Jones interactions between the analyte and water; System 2: p1, p2, p3 and the standard deviations for Lennard-Jones and Coulomb interactions between the analyte and H⁺ ions. For system 1, p1 was not included because its *p*-value (p = 0.117) was higher than the 0.05 cutoff point. The derived coefficients for the abovementioned parameters showed p-values below 0.05 (Table S1[†]).

Out of all the physicochemical properties that we tested, the only one that showed a statistically significant contribution was the water solubility of the analyte (S_W). The final model showed an R^2 of 0.82 and an MAE of 0.13.

The coefficients and intercept of the developed model were determined to be as follows:

$$\log RRF = lLJ + cCoul - 0.14S_W + 2.51$$
(8)

where,

$$ILJ = 0.63LJ_{p1}^{H} - 5.80LJ_{p2}^{H} + 4.92LJ_{p3}^{H} + 3.49LJ_{std}^{H} + 0.21LJ_{p2}^{W} - 0.20LJ_{p3}^{W} + 0.35LJ_{std}^{W}$$
(9)

$$c\text{Coul} = -0.01\text{Coul}_{p1}^{\text{H}} + 0.03\text{Coul}_{p2}^{\text{H}} - 0.03\text{Coul}_{p3}^{\text{H}} - 0.04\text{Coul}_{\text{std}}^{\text{H}}$$
(10)

where, LJ^H are the Lennard-Jones interactions between the analyte and H^+ ions from System 2. Coul^H are the Coulomb interactions between the analyte and H^+ ions from System 2. LJ^W are the Lennard-Jones interactions between the analyte and water molecules from System 1. The values for *p*1, *p*2, and *p*3 in System 2 were 0.5, 34 and 50. The values for *p*2, and *p*3 in System 1 were 44 and 89.

The *p*-values of the coefficients and the intercept were all below 0.05 (Table S1[†]). The standard errors for the derived coefficients are presented in Tables S1 and S2.[†] The R^2 and MAE of the model were comparable to those in the study of Oss *et al.*²⁵ where they observed an R^2 of 0.67 and a standard residual error of 0.86 log units. While the two studies are very different in the computational approaches, they both share datasets of similar size (48 *vs.* 62) and they both use multilinear regression models as their final predictive models thus allowing for meaningful comparisons.

We examined whether the differences between the experimental and modeled RRF (absolute errors) could be explained due to the different retention times (RT) of the chemicals and by extension due to the different ratios of water to methanol, but we did not observe a significant association between the two. Neither did we observe a significant association between RRF and RT (Fig. S6†).

Our modeling calculations showed that the interactions of the analyte with the water molecules in System 1 were similar but slightly more predictive than the interactions of the analyte with the H_3O^+ ions (R^2 : 0.42 vs. 0.39). Given the great collinearity of these two variables, including both of them in the model renders the coefficients for H_3O^+ insignificant (p > 0.05). This observation suggests that, at least in terms of interactions with the analyte, the H atoms in the H_2O molecules are not distinguishable from the H atoms in the H_3O^+ ions.

As mentioned earlier in the methods, we examined the effect that compounds with high penalties may have on the predictive power of the model. Including all compounds with penalties over 300 resulted in a small decrease in R^2 (0.82 vs. 0.74) and a small increase in the MAE (0.13 vs. 0.15) of the model (Fig. 4). After incrementally removing compounds with high penalties from the dataset, we observed that the R^2 of the model appeared

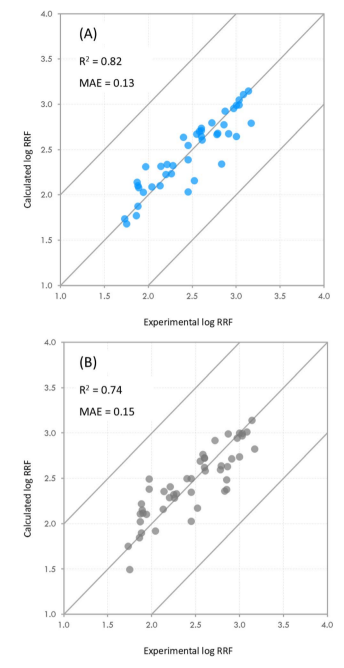


Fig. 4 Experimental and calculated values of log transformed RRF using the developed model. The diagonal lines show the 1-to-1 agreement line, and the $\pm 1 \log$ unit deviation line. Plot (A) shows the results of the model after removing compounds with a penalty over 300 (dataset n = 42). Six chemicals were excluded from the dataset in this iteration. Plot (B) shows the results of the model including all the chemicals in the dataset (dataset n = 48). R^2 is the coefficient of determination and MAE is the mean absolute error between the predictions and the experimental values.

to be consistent with an increase around cutoff points of 250 and 300 (Fig. S7†), which confirmed our initial cut-off point of 300. We should note at this point that while in this particular case, the effect of including compounds with high penalties appears to be minimal, we do not know how that may manifest in other datasets with different compositions and with compounds with higher penalties.

The 10-fold cross-validation showed an R^2 of 0.52 and an MAE of 0.25 for compounds that were not included in the training set (Fig. S8A[†]). This shows that the model can make reasonable predictions for chemicals that were not included in the training set. Two chemicals were shown to be outside of the applicability domain of the model (based on the definition in the Methods section). These two chemicals were Furalaxyl and Dicrotophos (Fig. S9[†]). During the cross-validation, Furalaxyl showed an absolute error of 10.1 log units, and Dicrotophos had an absolute error of 5.58 log units. Both chemicals had penalties lower than 300 so it does not seem that their penalties would be a likely explanation (ESI spreadsheet[†]). Most likely, these two chemicals are structurally and physicochemically distinct from the other chemicals in the dataset. This is supported by the observation that when these two compounds are included in the dataset their absolute errors are 0.001 log units for Furalaxyl and 0.03 log units for Dicrotophos (Fig. S10[†]).

The *y*-randomization showed that when the model is trained on random data the expected R^2 is 0.03 (Fig. S8B†). This is substantially lower than both the R^2 of the model with all the chemicals (0.74) and the R^2 of the 10-fold cross-validation (0.52). This observation suggests that the model is making meaningful predictions that are distinct from random predictions.

In trying to understand the contributions of the different interactions to RRF we examined the different terms of eqn (8) for two chemicals that showed near 0 differences between experimental and calculated values of RRF. The two chemicals were (1) Cinchophen with a log RRF of 1.73 (calculated log RRF = 1.74), and (2) Loratadine with a log RRF of 3.14 (calculated $\log RRF = 3.15$). Both chemicals' $\log RRF$ is determined to a larger extent by the Lennard-Jones and Coulomb interactions and to a smaller extent by their water solubility. For both chemicals, the Lennard-Jones interactions appear to have a positive contribution to RRF while the Coulomb interactions appear to have a negative contribution (Fig. 5). This is consistent for all chemicals in the dataset. When comparing Cinchophen and Loratadine, it appears that the lower RRF of Cinchophen is due to smaller *l*LJ and *c*Coul terms (Fig. 5). The Lennard-Jones potential approximates the van der Waals interactions and the Coulomb potential represents the ability to engage in hydrogen bonding. Previous studies have suggested that increased non-polar character, which would be represented by the Lennard-Jones potential is associated with higher RRF, while increased polar character which would be represented by the Coulomb potential is associated with a decrease in RRF.^{25,26,30,54,55} Our observations appear to be in agreement with the findings from previous studies.

Water solubility appears to play a small (Fig. 5) yet significant role (Tables S1 and S2[†]) in the model. For all compounds in the dataset, the sS_W term has a positive contribution to RRF. Based on this observation, one would expect that compounds with higher water solubility would have a higher RRF. However, given that the term sS_W is several orders of magnitude smaller than the *l*LJ and *c*Coul terms, the influence of sS_W on RRF is minimal in comparison. In our developed model, sS_W acts as

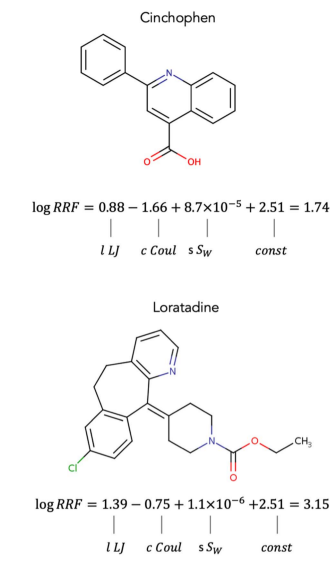


Fig. 5 Contributions of Lennard-Jones interactions, Coulomb interactions, and water solubility to the calculated log RRF for two compounds that showed near 0 errors between the experimental values and the predictions of log RRF. Cinchophen: experimental log RRF = 1.73 and calculated log RRF = 1.74. Loratadine: experimental log RRF = 1.73 and calculated log RRF = 1.74.

a corrective factor rather than a determining factor. Furthermore, water solubility is known to decrease with increasing molecular weight,⁵⁶ which is also what we observed in our dataset. The contribution of the sS_W for Cinchophen is slightly larger than that of Loratadine which is in agreement with their molecular weights (Cinchophen: 249.26 g mol⁻¹, Loratadine: 382.89 g mol⁻¹).

Limitations and future considerations

One limitation that needs to be acknowledged is that while our model showed good accuracy (R^2 : 0.82) the computational cost of our approach is much higher than other approaches that rely on simpler descriptor generators like PaDEL⁵⁷ or Mordred.⁵⁸ Running one simulation on one system for one chemical takes approximately 13 min using an ASUS GeForce GTX 1080 TI 11 GB Turbo GPU. Conducting simulations for 48 chemicals, 2 systems, and 3 replicates each comes up to 62.4 h. This may limit the ability of the model to be used as an online application as it would require access to GPUs. The workflow is, however, applicable in PCs with NVIDIA GPUs.

Another limitation that should be acknowledged is that, in this study, we examined only one type of force field (CGenFF). Future applications could examine whether using other types of force fields like gaff2 from AMBER and GROMOS from GRO-MACS can produce better predictions than CGenFF.

Finally, on the experimental side, it should be acknowledged that for the purposes of this study, we tested only two solvents for our LC gradient, HPLC water and methanol. As the ionization efficiency of chemicals is known to vary by different solvents,⁵³ the effect of that variability on the modeling calculations is something that needs to be investigated further.

Conclusion

Our study presents a novel approach for modeling the ionization efficiency of organic molecules. Our approach can be used in combination with existing approaches for concentration estimates of chemical compounds in environmental and biological samples. While there is a variety of modeling approaches for RRF, our view is that these approaches are complementary rather than competing. When trying to estimate concentrations of chemicals in environmental or biological samples, combining the results of multiple different approaches can help establish multiple layers of evidence that can be used in support of a prediction when analytical standards are unavailable.

Data availability

All code, ESI spreadsheets,[†] underlying datasets, and chemical structures used in this study are publicly available and can be accessed on GitHub under the following repository: https://github.com/dimitriabrahamsson/electro-spray. All Python scripts are accompanied by instructions on how to run them in order to replicate the findings of the study.

Author contributions

D. A. co-wrote the manuscript, designed the study, conducted the lab experiments, ran part of the simulations, and worked on the modeling section. L.-A. K. co-wrote the manuscript, ran simulations, and assisted with the modeling section. T. J. assisted with writing the manuscript and proofreading. J. Y. helped with proofreading, providing expertise on the analytical side, and assisting with revisions. X. J. helped with proofreading, providing expertise on the analytical side, and assisting with revisions. D. L. co-wrote the manuscript, supervised the experimental part of the study, provided advice and expertise on designing and executing the experiments, and interpreting the findings.

Conflicts of interest

The authors declare that they have no competing interests to report.

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