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An efficient data filtering strategy for easy metabolite detection from the direct analysis of a biological fluid using Fourier transform mass spectrometry

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Short title of up to 70 characters: Data filtering strategy for metabolite detection from DI-HRMS data
Abstract

RATIONALE

High throughput analyses require an overall analytical workflow including robust and high speed technical platform but also dedicated data processing tools able to extract the relevant information. This work aimed at evaluating post-acquisition data mining tools for selective extraction of metabolite species from direct introduction high resolution mass spectrometry data.

METHODS

Investigations were performed on spectral data in which seven metabolites of vinclozolin, a dicarboximide fungicide containing two chloride atoms, were previously manually identified. The spectral data obtained from direct introduction (DI) and high resolution mass spectrometry (HRMS) detection were post-processed by plotting the mass defect profiles and applying various data filtering methods based on accurate mass values.

RESULTS

Exploration of mass defect profiles highlighted, in a specific plotting region the presence of compounds containing common chemical elements and pairs of conjugated and non-conjugated metabolites resulting from classical metabolic pathways. Additionally, the judicious application of mass defect and/or isotope pattern filters removed many interfering ions from DI-HRMS data, greatly facilitating the detection of vinclozolin metabolites. Compared to previous results obtained by manual data treatment, three additional metabolites of vinclozolin were detected and putatively annotated.

CONCLUSIONS

Tracking simultaneously several specific species could be efficiently performed using data mining tools based on accurate mass values. The selectivity of the data extraction was improved when the isotope filter was used for halogenated compounds, facilitating metabolite ion detection even for low abundance species.

Keywords: mass defect, data mining, direct introduction mass spectrometry, Fourier transform mass spectrometry, metabolite identification
Introduction

Nowadays, human exposure to external factors, *i.e.* environment and diet or lifestyle, is one of the major issues in public health. The study of the xenometabolome is useful for evidencing such exposure and particularly to assess the simultaneous exposure to mixtures of several contaminants (*e.g.*, pesticides, drugs or persistent organic pollutants). A rapid approach including efficiency, selectivity and sensitivity is required to perform characterization of the xenometabolome, especially for large scale analysis in cohort studies. Targeted acquisition methods for detection and quantitative analysis of residues have been reported, which allow the simultaneous screening of several families of compounds.\(^1,2\) However, such approaches have some limitations such as the cost of trace analyses and the limitation of the search for new markers, which may reflect exposure to new contaminants or to a combined exposure to several xenobiotics. Therefore, global approaches combining metabolite fingerprinting with appropriate data processing tools seem to be the most suitable for the rapid screening of expected as well as unknown markers of exposure to toxicants.

Mass spectrometry with on-line liquid chromatography, gas chromatography or capillary electrophoresis (*i.e.* LC/MS, GC/MS or CE/MS) has been the most often used approach in metabolite studies, but the use of on-line separation techniques greatly limits the throughput of analyses. Direct introduction mass spectrometry (DIMS) enabling mass spectrum acquisition in a few minutes is a valuable high throughput approach.\(^3\) Such direct analyses require appropriated ionization techniques for performing rapid mass spectral acquisition.\(^4\) Most of them are atmospheric pressure sources. Typically, the ambient ionization techniques such as direct analysis in real-time (DART), desorption electrospray (DESI) and extractive electrospray ionization (EESI) can be applied for direct analysis of complex mixtures with limited or no sample preparation. Their applications in many different fields (*e.g.*, forensics, environmental and drug studies) have been reported.\(^5\) Nevertheless, the electrospray ionization (ESI) coupled to chromatography system is the method commonly used in metabolite studies but is increasingly used in the DIMS approach.\(^6-10\) ESI is sensitive to matrix effects since all compounds present in the matrix, including endogenous components as well as xenobiotics and their metabolites, undergo simultaneously the ionization process. Hence, ionization competition between different compounds occurs, leading to ion suppression phenomenon. In addition to the sample dilution, which reduces the matrix effects, the use of Fourier transform mass spectrometry (FTMS) instruments, providing high performances as mass resolving power and dynamic range, can reduce the drawbacks of...
The distinction of isobaric ions (i.e. same nominal mass but different elemental compositions) and the simultaneous detection of a large number of species in the same mass spectrum become possible with this instrumentation. However, the inability of DIMS to distinguish isomers hampers metabolite identification even using the most powerful instrument. The resulting mass spectral data remain complex and difficult to be interpreted because of the large number of ions detected simultaneously in a single experiment, potentially with close m/z values.

As well, to analyze such complex mass spectrum, data processing procedures based on mass defect can be efficiently used to facilitate detection and selection of chemically-relevant ions. The term ‘mass defect’, first introduced by E. Kendrick, refers to the difference between the accurately measured m/z value and its nominal value. The Kendrick method has been used to process high resolution mass spectrometry (HRMS) data. It is considered as a data classification method, facilitating the distinction of homologous compounds having different numbers of same base units. Kendrick plots have been successfully applied in the field of petroleomics, and few examples have also been reported for organic matter and metabolomic analyses. Werner et al. reported the Kendrick plot of 7,000 compounds extracted from the KEGG (Kyoto Encyclopedia of Genes and Genomes) database, resulting in the classification of compounds in homologous series as illustrated for the carboxylic fatty acids series. Alternatively, this method helps in the calculation of the elemental compositions of high m/z ions based on the known elemental compositions of lower m/z ions from the same series of compounds. Yet, due to the large chemical diversity of small molecules constituting the metabolome, the Kendrick approach seems less appropriate for detection of metabolites in metabolomics than for detection of apolar compounds in petroleomics.

Afterwards, Zhang et al. used the mass defect based approach to process complex HRMS data in drug metabolism studies. This approach called “mass defect filter” (MDF) is a data filtering method using a defined mass defect window centered at the mass defect value of the precursor compound ions. This allows to discard most of interfering ions. Such a data mining tool facilitates detection and subsequent annotation of specific exogenous compounds as well as their metabolites present in complex biological matrices. It has been successfully applied for the selective detection of drug metabolites in various biological materials. It has also been employed in the screening of contaminants in waste water, in the field of lipidomics and in natural product studies. A recent study from Geng et al. has
demonstrated the usefulness of MDF for the specific detection of traditional Chinese medicine components and their corresponding in vivo metabolites from biological materials. MDF is a real data filtering approach compared to the classical post-processing methods, which generate extracted or reconstructed chromatograms of predicted ions (EIC) as a visualization approach for detection of only expected ions.\textsuperscript{20} Hence, it represents an efficient data mining tool for simplifying large mass spectral datasets. Above all, MDF enables to detect both expected and unexpected metabolite species.

Until now, mass defect filtering has mainly been applied for post-processing complex data generated from liquid or gas chromatography coupled to high resolution mass spectrometry. Very few applications of MDF to mass spectral data acquired by DI-HRMS technique have been reported. Erve et al. tested the performances of DI-HRMS acquisition and data post-processing techniques for evaluating ZipTip cleaning procedures applied to biological matrices.\textsuperscript{8,9} In our previous work, the detection and characterization of seven metabolites of vinclozolin \[3-(3,5-dichlorophenyl)-5-methyl-vinyl-1, 3-oxazolidine-2,4-dione\], a dicarboximide fungicide, were manually performed on mass spectral data obtained by direct analysis (DI-HRMS) of urine of rats treated with this compound.\textsuperscript{10} Additionally, an isotope filtering procedure was tested, facilitating the detection of the seven vinclozolin metabolites ions.

In this work, investigations were done on a data filtering strategy developed for DI-HRMS data. First, the exploration of the mass defect profile was done to understand the distribution of different species based on their mass defect values. The search for organic compounds bearing common chemical elements and for metabolites generated from classical metabolic pathways was performed. Then, different data filtering templates (i.e. mass defect and isotope filters) were applied to evaluate their efficiency in selecting urinary metabolites of vinclozolin from raw DI-HRMS data. Three new low abundance metabolites were then detected by this strategy. Subsequent LC/MS and MS/MS (tandem mass spectrometry) experiments were performed to consolidate the putative metabolite annotation.

**Materials and Methods**

*Animal experimentation*

Urine of rats treated with vinclozolin was obtained as already described.\textsuperscript{28} Three male rats were individually housed in metabolic cages. They were fed by gavage with [\textsuperscript{14}C]-vinclozolin
and unlabeled vinclozolin dissolved in dimethyl sulfoxide to obtain a final activity of 0.481 MBq [\textsuperscript{14}C]-vinclozolin/kg and a 1 mg/mL final concentration of vinclozolin. Urine was collected for 24 h and stored at -20 °C until analysis.

**Analytical method**

Mass spectrometric analyses were performed in negative ESI mode, using a hybrid ion trap LTQ Orbitrap Fourier transform mass spectrometer (LTQ Orbitrap XL Thermo Fisher Scientific, Bremen, Germany).

**DIMS** data were obtained from direct analysis of rat urine sample using high resolution mass spectrometry with a mass resolving power (full width at the half maximum height, **FWHM**) set at 60 000 for m/z 400. Both flow injection analysis (**FIA**) and LC/MS approach were performed as described in previous work.\textsuperscript{10}

**MS/MS** experiments were performed on both the monoisotopic and (M+2) isotope peaks of vinclozolin metabolites using **CID** (collision-induced dissociation) conditions in the linear ion trap (LTQ) device. The following parameters were used: isolation width of precursor ions of 2 u, activation time of 30 ms, and normalized collision energy of about 20 % (arbitrary units). All **CID** experiments were carried out under high resolution detection with a mass resolving power of 7 500 (**FWHM**) for m/z 400.

**Data processing**

The proprietary .raw data file format was first converted into .cdf format using the file converter tool of Xcalibur 2.0.7 software (Thermo Fisher Scientific, Courtaboeuf, France). The mass spectral data was further processed using a script written in R 3.1.2,\textsuperscript{29} a statistical programming language (the script is freely available upon request). It was used to generate the mass defect profiles by plotting the mass defect value of each peak versus the measured m/z value. Different data filter templates were also created.

First, compounds containing specific chemical elements were searched based on the detection of \textsuperscript{12}C/\textsuperscript{13}C, \textsuperscript{14}N/\textsuperscript{15}N, \textsuperscript{16}O/\textsuperscript{18}O, \textsuperscript{32}S/\textsuperscript{34}S and \textsuperscript{35}Cl/\textsuperscript{37}Cl isotope ion pairs by selecting peaks displaying a pre-defined m/z difference value: \( \Delta = 1.0034 \pm 0.0002 \) u for \textsuperscript{12}C/\textsuperscript{13}C isotopes, \( \Delta = 0.9970 \pm 0.0002 \) u for \textsuperscript{14}N/\textsuperscript{15}N isotopes, \( \Delta = 2.0042 \pm 0.0002 \) u for \textsuperscript{16}O/\textsuperscript{18}O isotopes, \( \Delta = 1.9958 \pm 0.0002 \) u for \textsuperscript{32}S/\textsuperscript{34}S isotopes and \( \Delta = 1.9970 \pm 0.0002 \) u for \textsuperscript{35}Cl/\textsuperscript{37}Cl isotopes.

Similarly, specific searches for peaks of the potential ion pairs corresponding to non-conjugated and conjugated metabolites were performed. The following mass differences were
used: $\Delta = 176.0321 \pm 0.0002$ u for glucuronide conjugates, $\Delta = 79.9568 \pm 0.0002$ u for sulfate conjugates, $\Delta = 255.9889 \pm 0.0002$ u for glucurono-sulfate conjugates, $\Delta = 305.0682 \pm 0.0002$ u for glutathione conjugates, $\Delta = 119.0041 \pm 0.0002$ u for cysteine conjugates and $\Delta = 161.0147 \pm 0.0002$ u for N-acetyl-cysteine conjugates.

Various data filtering methods were then employed to process the mass defect profiles.

A MDF using a mass defect window (i.e. either $\pm 0.050$ u or $\pm 0.060$ u) centered at -0.0113 u, the theoretical mass defect value of the deprotonated vinclozolin (the theoretical accurate m/z value of [M-H]$^-$ ions is 283.9887) was applied to eliminate species with mass defect values outside the window used.

Alternatively, isotope pattern filter (IPF) templates were applied based on the accurate mass difference of 1.9970 u between $^{35}$Cl and $^{37}$Cl using a $\pm 0.0003$ u window for tracking M and (M+2) diagnostic isotope ion pairs characteristic of species containing chlorine atom(s). A complementary criterion was also added to specifically select dichlorinated species by taking into account the relative isotope abundance of the (M+2) peak compared to the monoisotopic one using the ratio interval ranging from 58 % to 70 %, i.e. 64 ± 6 %, which corresponds the theoretical value for dichlorinated species.

The resulting processed data were converted into reconstituted mass spectrum format in order to facilitate the comparison with the original data.

**Results and Discussion**

(Figure 1 near here)

The mass spectrum of diluted urine of rats treated with vinclozolin is presented in Figure 1a. This spectrum was obtained in previous work using DI-HRMS approach without neither prior sample clean-up nor chromatographic separation. Seven metabolites of vinclozolin were detected: M1, M4, M5 and M6 as non-conjugated metabolites, the sulfate conjugate of M5 and glucuronide conjugates of M4 and of M5 (Table 1). However, such manual search is time consuming and cannot be easily applied for large size and complex spectral data, especially when there are low abundance metabolites and presence of interfering ions with m/z values close to that of metabolites.

Here, an automated data processing tool was built for performing a fast detection of vinclozolin metabolites in urine sample from DI-HRMS data. From this dataset, the seven
previously annotated metabolites of vinclozolin (displayed as colored dots in the reported figures) were scrutinized first to test the efficiency of the data filtering methods.

**Mass defect profiles**

The mass spectral data were processed in the mass defect scale, which consists in the projection of projecting all detected signals in a two dimensional representation as proposed by Zhang et al., providing the so-called mass defect profile (Figure 1b and Figure 2). Each dot represents a peak detected at an accurate m/z value and a specific mass defect value. From the rat urine analyzed in this work, a region containing a high dot density is located at m/z values lower than 600 u and mass defect values ranging from -0.10 to 0.20 u, as shown on the dot density plot in the Figure 1b.

(Figure 2 near here)

In order to better understand the distribution of all dots across the mass defect profile, species containing chemical elements commonly found in organic compounds such as C, N, O, S and Cl were automatically searched and represented by colored dots in the Figure 2a. This involved the search of isotope ion pairs having specific accurate mass differences between $^{12}$C and $^{13}$C, $^{14}$N and $^{15}$N, $^{16}$O and $^{18}$O, $^{32}$S and $^{34}$S and $^{35}$Cl and $^{37}$Cl (with an error of 0.0004 u). Most of these species were located inside a specific plotting region described above, indicating the presence of most of species, which are characteristic of the sample, as reported by Zhang et al. for several biological matrices (i.e. plasma, urine, bile, and feces). However, the authors mentioned that dot distribution of common drugs overlapped dot cluster in the mass defect profile of urine samples whereas better separation was observed for other biological matrices. Here, the seven expected metabolites of vinclozolin were also detected in the region of high dot density. The signals located outside the dense dot clusters may be attributed to endogenous and contaminant compounds as demonstrated by the occurrence of isotope ion pairs (e.g., $^{12}$C/$^{13}$C in the Figure 2a). However, other dots characterized by anomalous mass defect values were also detected, suggesting the presence of artifact peaks, which could potentially result from Fourier transform experiments. It should be underlined that mass defect profiles generated from LC/MS data displayed large dot densities forming vertical lines at the low-mass threshold (i.e. about m/z 150), which should correspond to artifact peaks generated from chemical noise. Such vertical lines were not observed from our DI-HRMS data.
Alternatively, potential conjugate metabolite species can be selectively detected through the search of known biotransformation reactions as proposed in Figure 2b and Figure 2c. These metabolites can be selected by searching accurate mass differences corresponding to the mass shifts resulting from classical metabolic pathways. For example, a $\Delta = 176.0321 \pm 0.0002$ u m/z difference was used for tracking glucuronic acid conjugates and their corresponding non-conjugated compounds in the Figure 2b. The colored dots corresponding to glucuronide, sulfate, glucurono-sulfate, glutathione, cysteine and N-acetyl-cysteine conjugates and their non-conjugated forms are mostly located in the same region of high dot density, which should contain the most relevant chemical information of the biological matrices (Figure 2b and Figure 2c). Here, conjugated vinclozolin metabolites (i.e. conjugated M4 and M5) as well as their non-conjugated analogues (i.e. M4 and M5) were found among the detected species. Thus, this approach enabling the simultaneous detection of conjugated and non-conjugated metabolite pairs appears to be an efficient way to reveal the (combined) exposure to a (or several) specific xenobiotic(s).

Nevertheless, data filtering methods such as MDF and other data mining tools can be applied to process more efficiently such profiles.

**Data filtering methods**

As reported in the literature,20 most of metabolites produced from phase I or phase II biotransformations display mass defect shifts within a $\pm 50$ mu window relatively to the mass defect value of the parent ion (except for glutathione conjugates which display a larger mass defect shift, i.e. 68 mu). Here, a MDF window ranging from -0.060 u to 0.040 u was applied, which is centered to the mass defect of the deprotonated vinclozolin.

The mass defect profile from DI-HRMS data contains more than 3400 dots (i.e. m/z values) as shown in the Figure 1. The extraction procedure using the MDF template removed many interfering ions leading to a strong reduction of the data size. The processed mass defect profile is composed of 667 variables in which ions of six metabolites of vinclozolin are detected, except the M5 glucuronide (data not shown). The mass defect of ions of this latter metabolite (i.e. 47 mu for the m/z 468.0467 monoisotopic peak) is out of the prior MDF window used. The selectivity of the MDF strongly depends on the closeness of the mass defect values of metabolites to that of the parent molecule and that of the studied xenobiotics relatively to that of endogenous compounds of biological matrices. Therefore, the choice of MDF templates strongly depends on the specific class of metabolites to be selected, e.g.
metabolites formed from minor changes (oxidation, methylation, demethylation…) have small mass defect shifts relatively to that of the parent molecule. In the case of vinclozolin, a restrained mass defect window cannot be applied due to the large isotope patterns of this molecule and, most importantly, because of the already shown occurrence of sulfate and glucuronic acid conjugates, which display large mass defect shifts compared to the parent molecule mass defect. The mass defect value of monoisotopic ions of the M5 glucuronide (*i.e.* 47 μm) is 58 μm higher than that of deprotonated vinclozolin (*i.e.* -11 μm). Hence, a larger MDF window (*i.e.* ± 0.060 u) ranging from -0.070 u to 0.050 u was used to retain ions of the seven known vinclozolin metabolites. The resulting mass defect profile was composed of about 796 variables, in which ions of the seven metabolites of vinclozolin were extracted (*Figure 3a* and 3b). Nevertheless, hundreds of peaks likely corresponding to interferences (*i.e.* endogenous compounds and contaminant species) overlapped the xenobiotic metabolite dots in this mass defect profile.

(*Figure 3 near here*)

Obviously, the use of a generic MDF template was not sufficiently to remove many interfering signals and to facilitate the detection of xenobiotic ions in urine sample, as suggested by Zhang et al.19 The use of multiple MDF templates in parallel can also lead to a more comprehensive detection of various classes of metabolites. Ruan and Zhu investigated the bioactivation pathways of ticlopidine in rat liver microsomes using stable isotope-labeled GSH adducts, LC/HRMS data acquisition and multiple MDF templates among which, the filtering of doubly charged ions was applied to selectively detect GSH adducts.31 Note that the structure of the studied parent compound should be carefully considered to ensure that unexpected metabolites are not lost during the filtering process.20

Alternatively, MDF can be combined with other data mining tools to selectively extract chemically relevant data.32-34 Here, a further data mining tool was performed by using an isotope pattern filtering (IPF). Few applications of isotope filters have been reported.10,33-37 For example, IPFs allowed selective detection of organochlorides in environmental samples from very large data sets produced by GC×GC–HRMS on a TOF instrument.37 However, to the best of our knowledge, the use of IPF to process spectral data obtained from DIMS experiments has never been reported until our previous work.10

Here, a combination of MDF (using a ± 0.060 u mass defect window) and IPF templates was tested for facilitating detection of the expected vinclozolin metabolites and potentially
additional unknown metabolites which were not evidenced in our previous work. Indeed, vinclozolin and its metabolites possess highly selective isotopic patterns. Hence, in addition to MDF, the template based on the accurate mass difference between $^{35}$Cl and $^{37}$Cl isotopes was applied, enabling the extraction of 100 dots (Figure 3c). Another IPF template taking into account an additional pre-defined relative abundance of the (M+2) isotopic peak compared to the monoisotopic peak was also performed in a similar way as Zhu et al. More precisely, MDF was combined with an IPF template including both an accurate mass difference window ranging from 1.9967 to 1.9973 u and a relative abundance of isotope pairs falling in the 58 %-70 % range for searching specifically the dichlorinated species. The processed mass defect profile is reported in the Figure 3d. A quite broad window (64 % ± 6 %) of relative peak intensity was used here because of an uncertainty in the relative isotope abundance (RIA) measurements even using FT/MS instrumentation. Indeed, a systematic error of about 20 % has been reported for RIA determinations from complex mixture samples analyzed in LTQ-Orbitrap instrument due to ion suppression phenomena and existence of a signal intensity threshold. In addition, isotopes of common chemical elements constituting the molecule such as C (at least two atoms), O, Cl and S atoms, contribute to the signal of the (M+2) isotope peak, leading to an increase of the relative abundance of this peak (theoretical value of 63.9 % for dichlorinated species), especially when the mass resolving power used is not enough for discriminating those (M+2) isotopic peaks.

Anyway, the mass defect profile resulting from the application of both MDF and IPF templates led to a drastic simplification of the dataset containing only 22 variables, among them, the seven known metabolite ions were clearly predominant (Figure 3d), showing the high selectivity of such data mining tools for processing DI-HRMS data. In addition, four other patterns characteristic of dichlorinated species were selected. Their ion pairs (i.e. monoisotopic and (M+2) isotopic peaks) were detected at i) m/z 159.9729 and 161.9699, ii) m/z 289.9991 and 291.9964, iii) m/z 297.9347 and 299.9318 and iv) m/z 387.9666 and 389.9637. The examination of LC/MS experiments showed the co-elution of each ion pair (as displayed by the extracted ion chromatograms based on the accurate m/z values of oxidized form of M5 metabolite in the Figure S1 in supporting information), confirming their chemical relation and then their putative belonging to the same dichloride isotopic pattern. The m/z 159.9729 ions could be putatively annotated as the deprotonated 3,5-dichloro-aniline species (with 1.6 ppm of mass error) coming from the M3 metabolite of vinclozolin. The m/z 289.9991 ions could correspond to deprotonated species of oxidized form of M5 metabolite (i.e. M5 with one double bond or a ring formation) and could be annotated as
C_{11}H_{10}O_{4}NCl_{2} with -0.3 ppm mass error. The latter m/z 387.9666 ions could be putatively annotated as C_{11}H_{12}O_{8}NCl_{2}S (-0.2 ppm error), which may be ion formulae of ionized sulfate conjugate of hydroxylated M5 metabolite. These three metabolites were not characterized as urinary metabolites of vinclozolin in previous studies.\textsuperscript{10,28} Unfortunately, the m/z 297.9347 value did not give any plausible elemental composition by considering the vinclozolin moiety. (Table 1 near here)

(Figure 4 near here)

To test the selectivity of IPF in our case, this latter data filter template (including the criteria of accurate mass difference and relative abundance for chloride isotopes) was directly applied on the unprocessed data (Table 1 and reconstituted mass spectrum from processed data in the Figure 4). A significant reduction of the data size from more than 3400 to only 24 variables was obtained, among them all ions (22 variables) selected from the previous processing using the combination of MDF and IPF filters were still present. One additional dichloride isotopic pattern was selected: the m/z 469.0502 and m/z 471.0471 ions should correspond to the (M+1) and (M+3) isotopic peaks of the glucuronide acid conjugate of M5. This ion pair was not selected when using MDF because the mass defect value of the m/z 469.0502 ions (slightly higher than 50 mu) is outside the used MDF window (i.e. ranging from -0.070 u to 0.050 u). This result demonstrated the high efficiency of IPF to extract selectively vinclozolin metabolites.

**Complementary LC/MS and MS/MS experiments**

LC/MS and MS/MS experiments were further performed to consolidate the putative annotation of the vinclozolin metabolites (Table 2). No MS/MS experiments could be performed for low abundance ions of M1, M3 and M6 metabolites. Because of the stability of the dichloroaniline structure, no decomposition from M3 metabolite ions could occur. From LC/MS analysis, extracted ion chromatograms were generated by using the accurate m/z values of the vinclozolin metabolites with mass tolerance of 5 ppm (Figure S1 in supporting information). Several chromatographic peaks were detected at the same m/z value, indicating the presence of isomer ions for almost vinclozolin metabolites. The existence of isomers of vinclozolin metabolites (i.e. diastereoisomers as well as regio-isomers) has already been reported in the previous metabolic work using radio-labelled parent compound.\textsuperscript{28} It should be noted that in addition to those structural isomers, isobaric species may also be detected (within the mass tolerance of 5 ppm used for EIC traces). Different ionic states of
vinclozolin metabolites could eventually exist since no organic or acid additives were added to the chromatographic solvent to improve the separation. This probably explains the detection of multiple chromatographic peaks for most of vinclozolin metabolites (Figure S1 in supporting information). To eventually resolve the isomer structure, LC/MS/MS experiments were performed. The resulted fragmentation patterns did not allow to elucidate the exact structure of isomers since almost identical MS/MS spectra were obtained for the different chromatographic peaks detected at the same accurate m/z value. Only the characterization of the conjugation type is provided by the formation of diagnostic product ions and/or the loss of neutral. For example, sulfate conjugate species (i.e. M5 and hydroxylated M5 sulfate) mainly decomposed by loss of SO₃ into the formation of non-conjugated species (i.e. m/z 292.0145 and m/z 308.0108 product ions from deprotonated M5 sulfate and hydroxylated M5 sulfate, respectively; Table 2). From glucuronide conjugates (i.e. M4 and M5 glucuronides), m/z 235.0456 product ions were mainly formed under CID conditions. These ions should correspond to deprotonated hydroxy-ethyl glucuronide (C₈H₁₁O₈ formulae), but do not give indication about the exact position of the conjugation. Nevertheless, the formation of these latter product ions suggest that conjugation may occur on one of the two hydroxyl groups located on the positions 3 and 4 of the butanamide of M4 or M5 metabolites.

The comparison of the MS/MS spectra of M5 and the possible oxidized form of M5 metabolites showed the main product ions detected at m/z values shifted by 2 u. The detection of m/z 229.9798 fragment ions from m/z 290 parent ions (i.e. oxidized form of M5) instead of m/z 231.9935 from m/z 292 parent ions (i.e. M5) suggested that one additional insaturation or cyclization should occur within positions 1 and 2 of the butanamide moiety. A putative structure involving a ring formation was proposed in the Table 2. Anyway, metabolite identification cannot be performed without availability of authentic reference compounds, according to the Metabolomics Standards Initiative (MSI).40 (Table 2 near here)

Conclusion

Our study constitutes one of the few examples of applications of dedicated data mining tools based on the mass defect for processing HRMS data generated from direct analysis of biological fluids without requiring any prior sample clean-up or chromatographic separation. Such a data processing strategy showed a good selectivity for the complex HRMS data
extraction and greatly improved the detection of xenobiotics and that of their metabolites in biological matrices.

In our model, the ability to resolve stable isotopes based on their accurate masses and relative intensities demonstrated the high efficiency of the approach used for the selective detection of urinary metabolites of vinclozolin. Interestingly, three additional metabolites, which have not been characterized in the previous metabolic work using $^{14}$C-radiolabelled parent compound,$^{28}$ were detected and putatively annotated.

Although our model is a favorable case since vinclozolin displays a highly specific isotopic pattern, such an approach can be applied for any compound displaying characteristic isotopic pattern. Alternatively, stable isotopes can also be synthetically incorporated into biological materials for screening specific compounds and their metabolites. Indeed, as shown in proteomic studies, the “mass defect tags” method, as employed in bio-molecular analysis,$^{41}$ when using reactive molecules having discernable mass defect values, constitutes a very attractive way for tracking desired labeled species.$^{33,34,42}$

Finally, such a high throughput metabolomic approach combining direct MS analysis of biofluids and complementary data mining tools appears to be very promising for phenotyping large cohorts and tracking exposure to various xenobiotics.

Anyway, it must be underlined that MDF and other data mining tools constitute a useful step in the global strategy for screening and characterizing the structure of metabolites which further requires complementary experiments and computational tools for metabolite identity assignment, e.g., empirical formulae determination and/or the interpretation of their product ion spectra. The molecular sites prone to biotransformation can be subsequently elucidated using the structural information from the $\text{MS}^n$ experiments.

**Acknowledgments**

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References


Table and figure captions

Table 1 Ions selected from the application of MDF combined with IPF or from the application of IPF alone to data from DI-HRMS analysis of urine sample of rats treated with vinclozolin. The MDF processing used a ± 0.060 u mass defect window centered at -0.0113 u; the IPF includes the accurate mass difference and relative intensity between M and (M+2) isotopic peaks, i.e. corresponding to dichlorinated species

Table 2 Main product ions putatively annotated from MS/MS experiments.

Figure 1 (a) Mass spectrum from diluted urine sample of rat treated by vinclozolin using FIA-MS (negative ESI ion mode) and high resolution mass spectrometry (LTQ-Orbitrap) and (b) the corresponding mass defect plot showing dot density

Figure 2 Mass defect plots with (a) colored and bold dots corresponding to peaks displaying accurate mass differences of the $^{12}\text{C}/^{13}\text{C}$, $^{14}\text{N}/^{15}\text{N}$, $^{16}\text{O}/^{18}\text{O}$, $^{32}\text{S}/^{34}\text{S}$ and $^{35}\text{Cl}/^{37}\text{Cl}$ isotope ion pairs. In (b) is reported the same mass defect profile with colored dots displaying aglycone species and their conjugated metabolite ions (i.e. glucuronide, sulfate and sulfo-glucuronide conjugates) whereas possible thioether type metabolites were shown in (c) (i.e. glutathione, cysteine and N-acetyl-cysteine conjugates) (see experimental section)

Figure 3 Processed mass defect profiles obtained after application of (a) MDF (using a ± 0.060 u mass defect window centered at -0.0113 u) and (b, c and d) Y-zoomed mass defect profiles after using (b) MDF and (c and d) combined MDF with IPF filter (see the experimental section). The profiles (c) and (d) exhibited ions having mass difference of 1.9970 ± 0.0003 u whereas the profile (d) showed only ion pairs having a relative isotope abundance of 64 ± 6 % for the (M+2) isotopic peak compared to the M monoisotopic peak (i.e. corresponding to dichlorinated species)

Figure 4 Reconstituted mass spectrum from data processed with IPF template only
<table>
<thead>
<tr>
<th>Measured m/z</th>
<th>Mass defect (mu)</th>
<th>(M+2)/M ratio $^{(a)}$</th>
<th>Ion chemical formulae</th>
<th>Mass error (ppm)</th>
<th>Putative annotation</th>
</tr>
</thead>
<tbody>
<tr>
<td>159.9729$^{(d)}$</td>
<td>-27</td>
<td>65</td>
<td>C$_6$H$_4$NCl$_2$</td>
<td>1.6</td>
<td>3,5-dichloroaniline (M3)</td>
</tr>
<tr>
<td>161.9699$^{(d)}$</td>
<td>-30</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>289.9991$^{(d)}$</td>
<td>-1</td>
<td>62</td>
<td>C$<em>{11}$H$</em>{10}$O$_2$NCl$_2$</td>
<td>-0.3</td>
<td>Oxidized M5 (or M5 with one double bond)</td>
</tr>
<tr>
<td>291.9964$^{(d)}$</td>
<td>-4</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>292.0147</td>
<td>15</td>
<td>63</td>
<td>C$<em>{11}$H$</em>{12}$O$_2$NCl$_2$</td>
<td>-0.6</td>
<td>N-(3,5-dichlorophenyl)-2,3,4-trihydroxy-2-methylbutanamide (M5)$^{(b,c)}$</td>
</tr>
<tr>
<td>294.0118</td>
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<td></td>
<td></td>
</tr>
<tr>
<td>297.9347$^{(d)}$</td>
<td>-65</td>
<td>63</td>
<td>Unknown</td>
<td>-</td>
<td></td>
</tr>
<tr>
<td>299.9318$^{(d)}$</td>
<td>-68</td>
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<tr>
<td>301.9991</td>
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<td>61</td>
<td>C$<em>{12}$H$</em>{10}$O$_2$NCl$_2$</td>
<td>-0.3</td>
<td>2-(3,5-dichlorophenyl)-carbamoyl-oxo-2-methyl-3-butenoic acid (M1)$^{(b,c)}$</td>
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<td>303.9961</td>
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<tr>
<td>317.9940</td>
<td>-6</td>
<td>62</td>
<td>C$<em>{12}$H$</em>{10}$O$_2$NCl$_2$</td>
<td>-0.4</td>
<td>3-(3,5-dichlorophenyl)-5-(1,2-dihydroxyethyl)-5-methyl-1,3-oxazolidine-2,4-dione (M4)$^{(b,c)}$</td>
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<td>C$<em>{12}$H$</em>{12}$O$_2$NCl$_2$</td>
<td>-0.6</td>
<td>2-(3,5-dichlorophenyl)-carbamoyl-oxo-2-methyl-3,4-dihydroxybutenoic acid, (M6)$^{(b,c)}$</td>
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<td>371.9714</td>
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<td>66</td>
<td>C$<em>{12}$H$</em>{12}$O$_2$NCl$_2$S</td>
<td>-0.8</td>
<td>Sulfate conjugate of M5$^{(b)}$</td>
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<tr>
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<td>387.9666$^{(d)}$</td>
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<td>65</td>
<td>C$<em>{12}$H$</em>{12}$O$_2$NCl$_2$S</td>
<td>-0.2</td>
<td>Sulfate conjugate of hydroxylated M5</td>
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<td>389.9637$^{(d)}$</td>
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<td>468.0467</td>
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<td>C$<em>{17}$H$</em>{20}$O$_{10}$NCl$_2$</td>
<td>-0.7</td>
<td>Glucuronic acid conjugate of M5$^{(b,c)}$</td>
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<td>469.0502$^{(d)}$</td>
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<td>470.0435</td>
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<td>471.0471$^{(d)}$</td>
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<td>494.0260</td>
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<td>69</td>
<td>C$<em>{18}$H$</em>{18}$O$_{11}$NCl$_2$</td>
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<td>Glucuronic acid conjugate of M4$^{(b,c)}$</td>
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<td>496.0228</td>
<td>23</td>
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</table>

(a) Experimental values to be compared to the theoretical value of dichlorinated compound of 64%; Metabolite identity assignment from (b) Bursztyka et al.$^{28}$ and (c) from Rathahao-Paris et al.$^{10}$. (d) bold m/z values corresponding to dichloride isotopic patterns not detected in our previous studies$^{10,28}$ (c) (M+1) or (M+3) isotopic peak of metabolite M5.
<table>
<thead>
<tr>
<th>m/z value of parent ion and elemental composition of putative metabolite</th>
<th>Measured m/z values of product ions (ion relative abundance %)</th>
<th>Elemental composition of product ions (error in ppm)</th>
<th>Proposed structure and fragmentation of [M-H]⁻ species</th>
</tr>
</thead>
<tbody>
<tr>
<td>290 C₁₁H₁₀O₂NCl₂ Oxidized M5</td>
<td>229.9778 (100) 159.9730 (1)</td>
<td>C₉H₆O₂Cl₂ (0.8) C₆H₄NCl₂ (1.5)</td>
<td><img src="image" alt="Proposed structure and fragmentation of [M-H]⁻ species" /></td>
</tr>
<tr>
<td>292 C₁₁H₁₂O₂NCl₂ M5</td>
<td>231.9935 (100) 159.9729 (28) 131.0352 (21)</td>
<td>C₉H₆O₂Cl₂ (-0.9) C₆H₄O (1.7)</td>
<td><img src="image" alt="Proposed structure and fragmentation of [M-H]⁻ species" /></td>
</tr>
<tr>
<td>318 C₁₁H₁₉O₂NCl₂ M₄⁻</td>
<td>274.0037 (100) 257.9726 (2) 246.0091 (43) 215.9987 (2)</td>
<td>C₉H₆O₂Cl₂ (-1.2) C₆H₄O (2.1)</td>
<td><img src="image" alt="Proposed structure and fragmentation of [M-H]⁻ species" /></td>
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<tr>
<td>372 C₁₁H₁₉O₂NCl₂S Sulfate conjugate of M5</td>
<td>*311.9502 (1) 292.0145 (100) 231.9935 (2) 182.9969 (1) 159.9728 (1)</td>
<td>C₁₀H₆O₂Cl₂S (-1.3) C₆H₄O (0.2) C₆H₄NCl₂ (1.1)</td>
<td><img src="image" alt="Proposed structure and fragmentation of [M-H]⁻ species" /></td>
</tr>
<tr>
<td>388 C₁₁H₁₉O₃NCl₂S Sulfate conjugate of hydroxylated M5</td>
<td>308.0108 (100)</td>
<td>C₁₀H₆O₃NCl₂ (3.2)</td>
<td>No indication about sulfate and hydroxy group positions</td>
</tr>
<tr>
<td>468 C₁₇H₂₀O₁₀NCl₂ Glucuronic acid conjugate of M5</td>
<td>450.0358 (2) 292.0145 (6) 235.0456 (100) 217.0352 (1) 193.0354 (5) 175.0249 (2) 159.9728 (1)</td>
<td>C₁₇H₁₈O₉NCl₂ (-1.3) C₁₂H₁₀NCl₂ (-1.2) C₆H₄O (0.2) C₆H₄NCl₂ (1.1)</td>
<td><img src="image" alt="Proposed structure and fragmentation of [M-H]⁻ species" /></td>
</tr>
<tr>
<td>494 C₁₉H₁₈O₄NCl₂ Glucuronic acid conjugate of M₄</td>
<td>307.0668 (7) 235.0458 (100) 217.0353 (1) 175.0250 (1)</td>
<td>C₁₉H₁₆O₁₀ (-1) C₁₄H₁₀O₈ (-0.7) C₁₂H₈O₄ (-0.2)</td>
<td><img src="image" alt="Proposed structure and fragmentation of [M-H]⁻ species" /></td>
</tr>
</tbody>
</table>
Figure 1
Figure 2