Accepted Manuscript

This document is the Accepted Manuscript version of a Published Work that appeared in final form in Analytical Chemistry, copyright © American Chemical Society after peer review and technical editing by the publisher. To access the final edited and published work see http://dx.doi.org/10.1021/acs.analchem.7b00743

> Saer Samanipour, Malcolm J. Reid, Kevin V. Thomas. 2017. Statistical variable selection: an alternative prioritization strategy during the nontarget analysis of LC-HR-MS data. Analytical Chemistry . 89(10): 5585-5591. ISSN 0003-2700.

It is recommended to use the published version for citation.

Statistical variable selection: An alternative prioritization strategy during the non-target analysis of LC-HR-MS data

Saer Samanipour,^{*,†} Malcolm J. Reid,[†] and Kevin V. Thomas^{†,‡}

†Norwegian Institute for Water Research (NIVA), 0349 Oslo, Norway
‡Queensland Alliance for Environmental Health Science (QAEHS), University of Queensland, 39 Kessels Road, Coopers Plains QLD 4108, Australia

E-mail: saer.samanipour@niva.no

Abstract

1

Liquid chromatography coupled to high resolution mass spectrometry (LC-HR-2 MS) has been one of the main analytical tools for the analysis of small polar organic 3 pollutants in the environment. LC-HR-MS typically produces a large amount of data 4 for a single chromatogram. The analyst is therefore required to perform prioritization 5 prior to non-target structural elucidation. In the present study we have combined 6 the F-ratio statistical variable selection and the apex detection algorithms in order 7 to perform prioritization in data sets produced via LC-HR-MS. The approach was 8 validated through the use of semi-synthetic data, which was a combination of real 9 environmental data and the artificially added signal of 31 alkanes in that sample. 10 We evaluated the performance of this method as a function of four false detection 11 probabilities namely: 0.01, 0.02, 0.05, and 0.1%. We generated 100 different semi-12 synthetic data sets for each F-ratio and evaluated that data set using this method. 13 This design of experiment created a population of 30,000 true positives and 32,000 14

true negatives for each F-ratio, which was considered sufficiently large enough in order 15 to fully validate this method for analysis of LC-HR-MS data. The effect of both the F-16 ratio and signal to noise ratio (S/N) on the performance of the suggested approach were 17 evaluated through normalized statistical tests. We also compared this method to the 18 pixel-by-pixel as well as peak list approaches. More than 92% of features present in the 19 final feature list via F-ratio method were also present in conventional peak list generated 20 by MZmine. However, this method was the only approach successful in classification of 21 samples, thus prioritization, when compared to the other evaluated approaches. The 22 application potential and limitations of the suggested method discussed. 23

24 Introduction

A large number of small polar organic pollutants are considered as chemicals of emerging 25 concern (CECs) due to their fate and behavior in the environment (as reviewed by Klečka 26 et al.¹ and La Farre et al.²). Liquid chromatography coupled to the high resolution mass 27 spectrometry (LC-HR-MS) has become the leading analytical instrumentation for analysis 28 of these pollutants in different environmental compartments.^{3–5} Measuring these pollutants 29 in the environment takes place through three different and/or complementary approaches, 30 namely target analysis, suspect analysis, and non-target analysis.^{4,6–8} For target analysis 31 the analyst has all the necessary information, including the retention time and the spec-32 tral information, for confident identification of a target analyte in complex environmental 33 samples.⁹ On the other hand for the suspect screening only limited information is available 34 while during non-target analysis the analyst does not have any prior information regarding 35 the identity of the analytes in the sample.^{7,9} Even though non-target analysis is the most 36 difficult and the least certain of the three mentioned approaches, this method is essential for 37 the discovery of new CECs in the environment.^{4,6,7} 38

39

40 Confident identification of pollutants based only on the data generated via non-target

analysis on LC-HR-MS, is a challenging task due to the volume and complexity of the data 41 .^{10,11} During the non-target analysis, each sample may produce thousands of features, where 42 each includes a measured exact mass, intensity, and the retention time.^{12,13} Therefore, the 43 analyst may have to prioritize among the features for structural elucidation. For LC-HR-44 MS data, there have been different approaches used for prioritization during the non-target 45 analysis. The simplest approach applies the absolute intesity and the detection frequency 46 as the main criteria for prioritization.^{9,12} However, high signal intensity and the detection 47 frequency does not guaranty environmental relevance. Another approach utilizes either tox-48 icity information (through effect-directed analysis¹⁴) or the elemental composition (through, 49 for example, filters for halogenated compounds¹²). However, the mentioned approaches are 50 complicated and may be biased towards a certain family of compounds, for example halo-51 genated ones. A less used approach, particularly in the field of environmental analysis, has 52 been the application of unsupervised and/or supervised statistical methods, such as princi-53 pal component analysis (PCA) and partial least square discrimination analysis (PLS-DA) 54 for prioritization of the relevant features.^{13,15} These statistical methods perform well when 55 used in metabolomics due to a more clear change in the sample composition. However, these 56 same methods may suffer when there is a high level of redundancy/similarity in the analyzed 57 samples.^{16,17} Recent studies have shown the superior performance of the supervised F-ratio 58 method combined with PCA for analysis of the data via gas chromatography coupled to 59 low resolution mass spectrometry (GC-MS) of complex samples.^{16–20} However, the F-ratio 60 method has never been used/optimized for the non-target analysis of the data recorded via 61 LC-HR-MS and particularly for complex environmental samples. 62

63

The mentioned statistical variable selection approaches can be applied to either a peak list^{11,13,15} or the whole chromatogram.^{16–24} Even though processing of the peak list is faster than the whole chromatogram due to its smaller size compared to the whole chromatogram, the raw chromatogram must go through preprocessing steps such as signal deconvolution, peak finding, peak picking, and peak integration in order to generate a final peak list useful to a prioritization method. All these preprocessing steps are prone to error when dealing with highly complex samples.^{22–25} The application of the statistical variable selection approaches to the whole chromatogram has been shown to result in reliable models and therefore, reliable prioritization.^{16–25}

73

The aim of this study is to adapt, comprehensively validate, and test the applicability 74 of the F-ratio method for the non-target analysis of LC-HR-MS chromatograms of complex 75 environmental samples. The F-ratio was applied to the whole chromatogram in order to 76 minimize the data manipulation and produce a reliable statistical model. We combined the 77 F-ratio method with the apex detection as well as adduct and isotope removal algorithms, 78 in order to adapt this method to be used for non-target analysis of LC-HR-MS data. We 79 comprehensively validated this method using a semi-synthetic data set, which consisted of 80 the background signal generated from the real environmental samples with the addition of 81 the signal of 31 alkanes randomly distributed as true positives and true negatives, and noise. 82 This data set was evaluated 400 times where the random selection of the alkanes and the 83 background signal caused generation of a completely different sample for each evaluation. 84 Finally, the chromatograms of 15 sludge extracts from three different locations in Norway 85 and three blanks were analyzed using the F-ratio method as well as conventional peak picking 86 algorithms. We also applied the F-ratio method to the peak list and compared this feature 87 list to the one produced via using the whole chromatogram. The feature lists via F-ratio 88 were compared to the peak lists generated by a conventional peak pick method, in order to 89 further evaluate and/or validate the applicability of this method for non-target analysis of 90 LC-HR-MS data. 91

⁹² Experimental Methods

⁹³ Environmental Sampling and Sample Preparation

⁹⁴ 15 sludge samples from three different wastewater treatment plants (WWTP) in Norway were ⁹⁵ collected (i.e. five replicates for each WWTP), during the spring of 2015. These WWTPs ⁹⁶ were located at Oslo, Hamar, and Gjøvik. The plants in Oslo and Hamar were equipped ⁹⁷ with a three stage treatment process, including physical, chemical, and biological treatment ⁹⁸ whereas the plant in Gjøvik has only the physical and chemical treatments. More details on ⁹⁹ the chemicals, the suppliers, and the sample preparation steps are provided in section S1 of ⁹¹⁰ Supporting Information.

¹⁰¹ Instrumental Conditions and Analysis

All the extracts were analyzed employing, Waters Acquity UPLC system (Waters Milford, 102 MA, USA). An Acquity UPLC HSS C18 column $(2.1 \times 150 \text{ mm}, \text{ particle size } 1.8 \text{ mm})$ (Wa-103 ters, Milford, MA, USA) was used for all the separations. A mixture of solvent A, 5 mM 104 ammonium formate at pH 3.0 and solvent B, acetonitrile with 0.1% formic acid at a constant 105 flow rate of $0.4 \text{ ml} \text{min}^{-1}$ was used for the chromatographic separations. The gradient varied 106 from 87% of solvent A to 5% of solvent A. More details regarding this method are provided 107 elsewhere.²⁶ Both the analytical column and the column guard were kept as 50 °C during 108 the separations. 109

110

¹¹¹ Xevo G2-S Q-TOF-MS (Waters Milford, MA, US) was used for analysis of all 18 samples, ¹¹² including the 15 sludge extracts and 3 blanks. The MS^1 , with a collision energy of 6 eV, and ¹¹³ the MS^2 , with a collision energy ramp between 15 to 45 eV were simultaneously recorded ¹¹⁴ during the whole chromatogram. We employed a mass range of between 90 Da and 700 Da ¹¹⁵ with a sampling frequency of 1.8 Hz. More information regarding the mass spectrometer ¹¹⁶ conditions is available elsewhere.²⁶

¹¹⁷ Data Processing and Workflow

All the chromatograms were recorded in profile mode employing MassLynx (Waters Milford, 118 MA, US). The chromatograms were then exported as netCDF files, using DataBridge package 119 (Waters Milford, MA, US) incorporated in the MassLynx software. The MS¹ channel (i.e. 120 low collision energy channel 6 eV) chromatograms were used for the data analysis. All the 121 exported chromatograms were then imported into Matlab.²⁷ These files were processed using 122 the following sequnece of steps in the stated order including the binning, retention alignment, 123 data matrix generation, F-ratio calculation, null-distribution validation, zero mask applica-124 tion, chromatogram folding, apex detection, and finally adducts and isotope removal. All the 125 steps taken in this workflow are explained in detail below and section S2 of the Supporting 126 Information. 127

128

We binned the exported chromatograms using a bin thickness of 10 mDa, which was 129 based on the observed mass accuracy of \pm 5 mDa in our data set (section S2.1). The mass 130 accuracy was defined based on the shift observed in the measured mass of the calibrant 131 injected every 20 s into the source. The binned chromatograms were then retention aligned 132 with a home-developed algorithm inspired by the piecewise method previously developed and 133 validated by Synovec group.^{25,28} We added an additional mass spectral correlation control 134 in order to increase the accuracy of the retention alignment. More details regarding both 135 binning and retention alignment processes are provided in Supporting Information section 136 S2. The retention aligned chromatograms were then unfolded to create a long vector of in-137 tensities for every single measured m/z value. These vectors were then stacked on top of each 138 other in order to produce a large matrix which was used for the statistical prioritization. 139 Every row in this matrix was a sample while every column was an independent variable. 140 The F-ratio was calculated for each variable,¹⁶ or column of the matrix, based on *a priori* 141 knowledge of the sample classification (section S2.3). An F-ratio threshold was calculated 142 using the probability distribution generated via null-distribution analysis.¹⁹ This procedure 143

aims to minimize the number of false positive detections as well as the method validation 144 during the analysis (see section S2.4). The variables that had an F-ratio smaller than the 145 defined threshold, based on the null-distribution, were set to zero in the data matrix. This 146 process was referred to as the zero mask application. Each zero mask applied chromatogram 147 then was folded back into a matrix where a row was one scan and a column was the signal 148 for a m/z value. We performed apex detection in the folded chromatograms (see section 149 S2.5 of Supporting Information). The apex detection groups the non-zero and statistically 150 meaningful variables which can be represented as a feature in the chromatogram. For ex-151 ample all the non-zero variables in a chromatographic peak can be grouped and represented 152 via only one pair of retention time and m/z value, thus a feature. Therefore, the apex de-153 tection generates a list of unique retention time and m/z value pairs for each sample. This 154 differs from conventional peak picking algorithms in that apex detection does not perform 155 signal modeling and/or integration therefore minimizes the signal manipulation. Finally, 156 the adducts and the isotopes were removed from this list in order to create the final unique 157 feature list for each chromatogram. This workflow provides the necessary initial information 158 for discovery-based non-target analysis of complex samples analyzed via LC-HR-MS. 159

160

We also performed F-ratio analysis on the peak list produced by conventional peak picking algorithm, MZmine 2²⁹(explained in detail below). The peak list was retention-aligned using a home-developed method using a mass window of 2 mDa and a retention window of 2 S. The retention aligned peak tables were used for F-ratio and null-distribution calculations. The peaks in the peak list with an F-ratio larger than the threshold were kept in order to produce the feature list. The feature list, finally, was processed for adduct and isotope removal in order to generate the final feature list.

168

169 Data pretreatment

During the validation process of the F-ratio method (i.e. analysis of the semi-synthetic data), 170 we did not employ any data pre-treatment methods such as mean-centering, standardiza-171 tion, and normalization. This choice enabled us to comprehensively evaluate the effect of 172 introduced noise on the performance of the F-ratio method. For the environmental sample 173 analysis, we tested different data pre-treatment methods such as mean-centering, standard-174 ization, and normalization before processing the data set with F-ratio method. However, 175 these pre-treatments did not affect the final unique feature list for the analyzed data set. 176 Therefore, we decided to work with the raw data and avoid performing any type of pre-177 treatment. 178

179 Computations

All the mentioned data processing steps were performed via Matlab, employing a Windows
7 Professional version (Microsoft Inc, USA) workstation computer with 12 CPUs and 128
GB of memory.

¹⁸³ MZmine Peak Picking

The conventional peak list for each chromatogram was generated using MZmine 2.²⁹ The 184 peak picking was performed by mass detection followed by GridMass 2D peak detection. A 185 five scan window was selected for the smoothing of the chromatogram in the time dimension 186 and a 10 mD window was used in the mass dimension. A minimum signal of 300 counts was 187 required for a peak to be considered as a meaningful peak. These parameters were optimized 188 based on the observed mass accuracy and the peak widths in both time and mass domains. 189 These parameter settings resulted in feature numbers varying between 7,500 for blanks and 190 12,500 for the samples from Oslo WWTP. 19

¹⁹² Principal Component Analysis (PCA)

¹⁹³ We employed principal component analysis (PCA)³⁰ for classification/separation of the sam-¹⁹⁴ ple groups. We performed PCA on the peak list generated via MZmine, variable selected peak ¹⁹⁵ list (i.e. the F-ratio applied to the peak list generated via MZmine) with F-ratio method, ¹⁹⁶ the whole chromatogram (i.e. pixel-by-pixel), and the variable selected chromatogram em-¹⁹⁷ ploying F-ratio method. The PCA was performed on the mean centered data utilizing the ¹⁹⁸ singular value decomposition algorithm.³⁰

¹⁹⁹ Results and Discussion

We validated the F-ratio method for data generated via LC-HR-MS, employing both semi-200 synthetic data and the real environmental data. The use of semi-synthetic data enabled us 201 to perform a large number of evaluations (i.e. total number of detection cases $62,000\times 4$) 202 knowing exactly the added signal, noise, and relative intensity of the added signal which, 203 translated into comprehensive validation of the proposed method. This would not have 204 been possible using spiked samples due to the limitation on the number of standards and 205 injections as well as the potential interference between the sample and the standard mixture. 206 This study is the first implication of this method for the data generated via LC-HR-MS as 207 well as adaptation of this method in order for it to be included in non-target identification 208 workflows. This method enables the direct prioritization of the unique features, which are 209 the main cause of the separation of different sample groups. Therefore, the identification 210 efforts can be focused on the prioritized unique features. 211

²¹² Validation via Semi-synthetic Data

We employed a semi-synthetic data set, which consisted of a combination of real environmental data and synthetic signal, for comprehensive validation of the F-ratio method. The signal of 31 alkanes (i.e. the neutral monoisotopic masses, Table S1) was added at different

concentrations to a background signal, which came from the real environmental data. During 216 each analysis, these 31 alkanes were divided in two randomly selected groups where the first 217 group of 15 alkanes was added to the background signal at concentration levels that were 218 statistically meaningful. Therefore, for these 15 alkanes the resulting F-ratios were larger 219 than the threshold. For the second group, 16 alkanes were added to the background at a sta-220 tistically constant concentration. Four different F-ratios of 208, 30, 28, and 13 having false 221 positive detection probabilities of 0.01, 0.02, 0.05, and 0.1 %, respectively, were evaluated. 222 Each F-ratio value was evaluated 100 times with different: background signal, combination 223 of alkanes, concentration levels, and retention times of true positives (i.e. 15 alkanes) and 224 true negatives, thus a total of 400 evaluations. The generation of the these semi-synthetic 225 data is described in detail in Supporting Information, section S3. Alkanes were selected for 226 our analysis because these compounds are not ionized by ESI source therefore we were sure 227 that these compounds were not present in the real background signal. This design of exper-228 iment created a set of 15 true positives, 16 true negatives, and a different background signal 229 during each evaluation, which enabled us to comprehensively examine the capabilities and 230 limitations of the F-ratio method. The number of evaluation (i.e. 100 for each F-ratio) was 231 selected based on our preliminary assessment, that showed that 100 analysis for each F-ratio 232 would generate a large enough population of true positives (TP) 30,000 and true negatives 233 (TN) 32,000 for that F-ratio, in order to fully validate this method. To compare the effect 234 of different F-ratio probability value on the final results, we employed normalized statistical 235 parameters such as rate of false positive, rate of false negative, sensitivity, specificity, and 236 accuracy.³¹ 237

238

Increased F-ratios resulted in a smaller number of false positives and a larger number of false negatives. The number of false positive detection ranged from 2,518 cases for the F-ratio of 208 having a probability of 0.01% to 9,525 cases for the F-ratio of 13 with a probability of 0.1%, Table 1 and Figure S7. The largest number of false negative detections of 2204 was

observed for an F-ratio of 208 whereas the smallest number of false negative detections of 243 1,404 was caused by an F-ratio of 13, Table 1. These trends were due to the fact that the 244 selection of a large F-ratio value (i.e. more strict selection criterion) lowers the probability 245 of false positive detection while increasing the probability of false negative detections. The 246 observed changes in F-ratio method performance were better projected through normalized 247 statistical parameters such as rate of false positive detection, rate of false negative detection, 248 sensitivity, specificity, and accuracy, Table 1. For example, the drop in the specificity and 249 accuracy observed for F-ratio of 13 (probability of 0.1%) showed the inadequacy of this F-250 ratio for the analyzed data set (Figure S7). This drop also indicated that this F-ratio may 251 cause a large number of false positive detections when analyzing this data set. Therefore, 252 the analyst is required to find an optimized F-ratio value in order to minimize the number 253 of potential false positive detection while limiting the number of false negatives. Among the 254 four F-ratios evaluated, the value of 28 (probability of 0.05%) showed to be the optimized 255 one, considering that this value provided the largest accuracy level, second largest sensitivity 256 level while maintaining a high level of specificity (Figure S7). 257

258

Further evaluation of our data set, showed that for F-ratios ≥ 28 more than 70% of the 259 false positive cases were coming from the background signal rather than the true negatives 260 (i.e. added signal of alkanes at constant concentrations). The observed trend was caused 261 by the high level of variability artificially introduced into the background signal during the 262 background generation. For F-ratio of 13, around 50% of the false positives were true neg-263 atives. In this case even though the signal of true negatives did not have a large level of 264 variability between sample groups, once added to the background, the variability in that 265 signal increased due to the inherent large variance in the background signal. Therefore, 266 these true negatives produced a large enough F-ratio, which met the F-ratio threshold and 267 were selected as positive detections. When looking at the false negative cases, for all four 268 F-ratios, the main causes of false negative detection were the large variability in the back-260

ground signal and the S/N threshold setting during the apex detection. Also in this case, 270 the random variability introduced into the true positives signal was not large enough to 271 overcome the variability present in the background signal. We tested these hypothesis by 272 increasing the initial concentration of the added signal of the true positives from 5% to 15%273 and also increasing the concentration factor from 2-8 to 2-20 (see section S3 of SI for more 274 details). With an F-ratio of 28, increasing these parameters reduced drastically the number 275 of false positive detection from 2,864 to 253 cases as well as the number of false negatives 276 from 1,570 to 35 cases after 100 simulations. These results indicated that the combination 277 of low level concentration of added alkanes, their low between sample group variability, and 278 finally the large level of variability introduced into the background signal have an important 279 effect on the performance of this algorithm. 280

281

Mean centering and standardization (i.e. division by the square root of standard devia-282 tion of each variable) with an F-ratio of 28, added signal of 5%, and the concentration factor 283 of 2-8 reduced the number of both false positive detection and false negative detection from 284 2,864 to 350 cases and from 1,570 to 97 cases, respectively after 100 simulations. These 285 pre-treatments' approaches both decrease the noise levels in the data set while emphasizing 286 the underlying trend.³⁰ This implies that these data pre-treatments reduced the effect of 287 artificially introduced noise in the data set while emphasizing the between group variability, 288 thus a decrease in the number of false positive and false negative detection. The type of data 289 pre-treatments employed prior to the F-ratio analysis is data set and objective dependent.³⁰ 290 Therefore, the analyst is required to optimize these data pre-treatments approaches in ad-291 vance in order to be able to produce reliable results. Further investigation on the effect of 292 these parameters on the F-ratio method are needed and will be subject of our future studies. 293

Table 1: The number of false positive detections, number of false negative detections, rate of false positive, rate of false negative, sensitivity, specificity, and accuracy parameters calculated for four different F-ratio values based on 100 evaluations for each F-ratio probability value.

		(1	U	1	/
Parameter	208 (0.01)	30(0.02)	28 (0.05)	13(0.10)	
False positive detection ^{a} (FP)	2,518	$3,\!172$	2,864	9,525	
False negative detection ^{a} (FN)	2,204	2,220	$1,\!570$	$1,\!404$	
Rate of false positive detection ^{b} (%)	7.3	9.0	8.0	23.0	
Rate of false negative detection ^{c} (%)	6.8	6.9	5.0	4.5	
Sensitivity ^{d} (%)	93.2	93.1	95.0	95.5	
Specificity ^{e} (%)	92.7	91.0	91.8	77.1	
$Accuracy^f$ (%)	92.9	92.0	93.3	85.0	

| F-ratio values (probability of false positive detection %)

^aThis parameter represents the number false positive detection out of total number of detections of 62,000, including 30,000 true positives (TP) and 32,000 true negatives (TN); ^bThe rate of false positive³¹ was calculated as FP/(FP+TN); ^cThe rate of false negative³¹ was calculated as: FN/(TP+FN); ^dThe sensitivity³¹ values were calculated using: TP/(TP+FN); ^eThe specificity³¹ values were calculated with: TN/(TN+FP); ^fThe accuracy³¹ values were calculated employing: (TP+TN)/(TP+FP+FN+TN).

²⁹⁴ The effect of S/N on F-ratio algorithm

The S/N is an important parameter, which affects the performance of the F-ratio algorithm 295 particularly during the apex detection. The apex detection step aims to reduce the level 296 of redundancy in the data set by grouping variables, that can be represented by a unique 297 one (see section S2.5 for more detailed information). We evaluated the effect of S/N on the 298 results of the algorithm with an F-ratio of 28. This evaluation was performed by varying 299 the S/N from 1 to 10 (i.e. 1, 3, and 10) and performing 20 analysis for each S/N value. The 300 F-ratio of 28 was selected based on the fact that it appeared to be the optimized F-ratio for 301 the evaluated data set. 302

303

We observed a slight decrease in the number of false positive detection as a function of increase in the S/N while the increase in the S/N had a positive effect on the number of detected false negatives, Table 2. However, the changes in the S/N did not appear to cause a large variation in the normalized statistical parameters such as rate of false positive, rate ³⁰⁸ of false negative, sensitivity, specificity, and accuracy.³¹ This suggested that the S/N ratio

³⁰⁹ has a less relevant effect on the performance of this method compared to the F-ratio value.

310 However, these results may be case dependent, therefore optimization of this parameter

³¹¹ based on the data set should be considered by the analyst.

Table 2: The number of false positive detections, number of false negative detections, rate of false positive, rate of false negative, sensitivity, specificity, and accuracy parameters calculated for four different S/N values, having an F-ratio of 28, based on 20 simulations for each S/N.

	S/N values		
Parameter		3	10
False positive detection ^{a} (FP)	654	629	583
False negative detection ^{a} (FN)		151	166
Rate of false positive detection ^{b} (%)		9.0	8.0
Rate of false negative detection ^{c} (%)		2.5	2.7
Sensitivity ^{e} (%)		97.5	97.3
Specificity ^{f} (%)	90.7	91.1	91.7
Accuracy ^{g} (%)	94.1	94.1	94.3

^aThis parameter represents the number false positive detection out of total number of detections of 12,400, including 6,000 true positives (TP) and 6,400 true negatives (TN); ^bThe rate of false positive³¹ was calculated as FP/(FP+TN); ^cThe rate of false negative³¹ was calculated as: FN/(TP+FN); ^eThe sensitivity³¹ values were calculated using: TP/(TP+FN); ^fThe specificity³¹ values were calculated with: TN/(TN+FP); ^gThe accuracy³¹ values were calculated employing: (TP+TN)/(TP+FP+FN+TN).

312 Comparison between the unique feature list and the conventional

313 peak list

Once the F-ratio method was validated via semi-synthetic data, we processed the chromatograms of the 15 sludge samples plus 3 method blanks using this algorithm. The same data set was also processed via MZmine, employing previously optimized parameters. The F-ratio method produced a list of unique features for each sample whereas MZmine created a conventional peak list for the same samples. We compared the unique feature lists produced via F-ratio method to the conventional peak lists by MZmine as well as the unique feature lists produced via application of F-ratio method to both the whole chromatogram and the
peak list by MZmine. These comparisons enabled us to further evaluate/validate the F-ratio
method for analysis of the data generated via LC-HR-MS.

323

More than 92% of the unique features via F-ratio method were also present in the con-324 ventional peak list via MZmine. For example, for one of the Oslo samples after the adducts 325 and isotopes removal 109 out of total 112 (i.e. 97%) unique features were also present in 326 the peak list of the same sample generated by MZmine. The number of features, via F-327 ratio method, before adducts and isotope removal ranged from 403 features for one of the 328 blank samples to 127 for the Oslo sample whereas after the adducts and isotope removal 329 the unique features numbers ranged between 302 for the blank sample and 112 for the Oslo 330 sample. For the conventional peak list, we observed around 7500 peaks for the blank whereas 331 this number was around 12500 for the sludge samples. When comparing the unique feature 332 list to the conventional peak list, the number of discrepancy cases varied between 3 cases 333 for Oslo sample and 23 cases for the blank samples. A discrepancy case is defined as a 334 unique feature detected via F-ratio that is not present in the conventional peak list. All 335 the discrepancy cases were classified in two categories, for ease of explanation. The first 336 category and the most dominant one, particularly in the blank samples belonged to unique 337 features, which appeared to be noise rather than analytical signal. Considering the large 338 number of variables evaluated occurrence of a certain number of false positives was likely. 339 The second category was mainly caused by the fact that MZmine performs peak modeling 340 during the peak picking and uses the modeled apex for estimation of both m/z value and 341 the retention time. Using this approach, this algorithm may group shoulders of a peak with 342 the main peak. The F-ratio method however treats the shoulders as independent variables 343 and thus potential unique features. This category of discrepancy cases may also be caused 344 by the resolution of our instrument of 35,000. All considered, the F-ratio combined with the 345 apex detection algorithm method showed to have a large number (i.e. $\leq 92\%$) of common 346

unique features with the conventional peak picking approach which is an indication of its
robustness. Furthermore, these results imply that this method can be implemented in the
non-target workflows for structural elucidation.

350

The F-ratio applied to the whole chromatograms of the environmental samples resulted 351 in 250 unique feature in average while producing 3 unique features in average when applied 352 to the peak list generated via MZmine. A large number of the observed discrepancy cases 353 were due to the signal deconvolution, which was caused by the complexity of the analyzed 354 samples. The unsuccessful signal deconvolution was directly translated into the large within 355 group variability in the area of the integrated peaks, thus their lack of detection. The second 356 group of discrepancies was due to the peak modeling algorithm in MZmine, which failed to 357 detect the shoulder of a peak in the m/z domain as a separate peak, therefore their absence 358 from the unique feature list. It should be noted that the mentioned sources of failure in 359 the F-ratio applied to the peak list may be case dependent and may vary from dataset to 360 dataset. Further investigation of the potential sources of discrepancy between the F-ratio 361 variable selection applied to the whole chromatogram vs the peak list are needed. 362

363

The F-ratio method appeared to be able to successfully separate the sample groups 364 while both peak list and pixel-by-pixel methods failed in carrying out this task, Figure 1. 365 Multivariate statistical methods such as principal component analysis (PCA) when dealing 366 with large, complex datasets with a large level of noise and redundancy may fail to classify 367 the samples in logical groups. Consequently, univariate methods such as F-ratio are used 368 prior to these tests in order to reduce the redundancy in the data set. Therefore, a clear 369 and logical separation of the samples in the score plots is a crucial indication of a successful 370 prioritization/variable selection. We performed PCA on zero mask applied chromatograms 371 following the variable selection, the retention aligned peak list via MZmine, and the whole 372 chromatogram (i.e. pixel-by-pixel analysis). In the case of the sludge samples the inherent 373

complexity of the background signal was translated into inability of both peak list based 374 and pixel-by-pixel based methods to separate these sample groups from each other properly. 375 The F-ratio method, on the other hand, was able to perform separation of the sample groups 376 because this method retains the variables that are causing the clustering of samples within a 377 particular group. We also performed the F-ratio variable selection on the peak list generated 378 via MZmine. In this case also the PCA was not able to separate the sample groups from 379 each other, Figure 1. Therefore, it was not possible to perform a prioritization based on 380 the peak list using the F-ratio method. Despite the mentioned complexity, the F-ratio 381 method was able to separate the sample groups from each other, thus performing successful 382 prioritization. These results also indicate the applicability of F-ratio method within the 383 structure elucidation workflows during non-target analysis of complex samples analyzed via 384 LC-HR-MS. 385



Figure 1: Figure depicting the PCA score plots of (a) peak list based classification, (b) peak list based after F-ratio variable selection (c) pixel-by-pixel analysis, and (d) the F-ratio method during the prioritization for non-target analysis of the 15 sludge samples plus 3 blanks.

³⁸⁶ Potential and Limitations

The F-ratio method combined with the apex detection showed to be a robust and reliable 387 approach for prioritization of the unique features that are relevant to the sample classifica-388 tion. This method was effective at prioritization even for cases where the other conventional 389 methods may fail due to the complexity of the analyzed data set, Figures 2 and 1. This 390 method minimizes the data manipulations such as peak picking and/or modeling and at the 391 same time results in a list of unique features which can be used for structure elucidation. 392 The F-ratio method reduces the redundancy in the data set and detects the relevant vari-393 ables in the data set enabling the analyst to focus on the identification of only the unique 394 and relevant features. This method also has the advantage of being less dependent on the 395 absolute intensity of the each chemical signal in the sample compared to the conventional 396 prioritization methods. In other words, as long as a chemical signal causes large enough 397 variability between the sample groups, independently from its absolute intensity, it will be 398 detected as a unique relevant feature (Figure 2). Additionally, this method can be used 399 for a battery of discovery-based non-targeted applications as long as there are replicates 400 present. Furthermore, by changing the initial hypothesis, one can interrogate the data set 401 in a completely different way. For example in case of the sludge samples in this study, by 402 assuming that all the sludge samples belonged to one group and the blanks to another group, 403 we could have selected the unique features which are in common in all the sludge samples 404 and simultaneously subtracted the blanks from our samples. 405

406

There are also some limitations to application of F-ratio method for non-target analysis of LC-HR-MS data. This method is computationally expensive due to the large data sets produced when employing LC-HR-MS. For example, each sludge sample chromatogram in this study produced around 180 million variables (Figure 2), which requires a large computational power in order to be done in a timely manner. Moreover, this method has to be complemented with target and suspect analysis using the conventional methods for ubiquitous chemicals or pollutants where measurable concentrations are more uniform. These
pollutants would not be detected as unique features with statistically significant differences
between sample groups. Also the use of data pre-treatment should be evaluated by the analyst on a case study base.

417

Considering capabilities and the limitations of the F-ratio method, this approach has a great potential to be applied to the LC-HR-MS non-target discovery-based analysis. The application of this method as well as its combination with the structural elucidation workflows are going to be subject of our future studies.



Figure 2: Figure depicting an overview of the F-ratio method vs the conventional methods as well as the venn diagrams of the comparison between the unique feature list and the conventional peak list generated via MZmine.

422 Associated Content

423 Acknowledgement

⁴²⁴ The authors are thankful to Prof. Bert van Bavel, Dr. Merete Grung and Dr. Jose A.
⁴²⁵ Baz-Lomba for their editorial input. We are also grateful to the Research Council of Norway
⁴²⁶ for the financial support of this project (RESOLVE, 243720).

427 Supporting Information

The Supporting Information including details regarding the sample preparation, analysis, sateps taken during the data processing, and semi-synthetic data generation is available free of charge on the ACS Publications website.

431 Author Information

- 432 Corresponding Author:
- 433 Saer Samanipour
- 434 E-mail: saer.samanipour@niva.no
- 435 Phone: +47 98 222 087
- 436 Address: Norwegian Institute for Water Research (NIVA)
- 437 0349 Oslo, Norway

438 References

- (1) Klečka, G.; Persoon, C.; Currie, R. Reviews of Environmental Contamination and Tox *icology Volume 207*; Springer, 2010; pp 1–93.
- (2) La Farre, M.; Pérez, S.; Kantiani, L.; Barceló, D. TrAC Trends Anal. Chem. 2008, 27,
 991–1007.

- (3) Giger, W. Anal. Bioanal. Chem. **2009**, 393, 37. 443
- (4) Krauss, M.; Singer, H.; Hollender, J. Anal. Bioanal. Chem. 2010, 397, 943–951. 444
- (5) Richardson, S. D. Anal. Chem. 2009, 81, 4645–4677. 445

447

- (6) Schymanski, E. L.; Singer, H. P.; Longrée, P.; Loos, M.; Ruff, M.; Stravs, M. A.; 446 Ripollés Vidal, C.; Hollender, J. Environ. Sci. Technol. 2014, 48, 1811–1818.
- (7) Gago-Ferrero, P.; Schymanski, E. L.; Bletsou, A. A.; Aalizadeh, R.; Hollender, J.; 448 Thomaidis, N. S. Environ. Sci. Technol. 2015, 49, 12333–12341. 449
- (8) Samanipour, S.; Langford, K.; Reid, M. J.; Thomas, K. V. J. Chromatogra. A 2016, 450 1463, 153–161. 451
- (9) others, et al. Anal. Bioanal. Chem. 2015, 407, 6237–6255. 452
- (10) Gorrochategui, E.; Jaumot, J.; Lacorte, S.; Tauler, R. Trends Anal. Chem. 2016, 82, 453 425 - 442.454
- (11) Yi, L.; Dong, N.; Yun, Y.; Deng, B.; Ren, D.; Liu, S.; Liang, Y. Anal. Chem. acta 455 **2016**, *914*, 17–34. 456
- (12) Chiaia-Hernandez, A. C.; Schymanski, E. L.; Kumar, P.; Singer, H. P.; Hollender, J. 457 Anal. Bioanal. Chem. 2014, 406, 7323-7335. 458
- (13) Schollee, J. E.; Schymanski, E. L.; Avak, S. E.; Loos, M.; Hollender, J. Anal. Chem. 459 **2015**, *87*, 12121–12129. 460
- (14) Thomas, K. V.; Langford, K.; Petersen, K.; Smith, A. J.; Tollefsen, K. E. Environ. Sci. 461 Technol. 2009, 43, 8066-8071. 462
- (15) Kalogiouri, N. P.; Alygizakis, N. A.; Aalizadeh, R.; Thomaidis, N. S. Anal. and Bioanal. 463 Chem. 2016, 408, 7955–7970. 464

- ⁴⁶⁵ (16) Pierce, K. M.; Hoggard, J. C.; Hope, J. L.; Rainey, P. M.; Hoofnagle, A. N.; Jack, R. M.;
 ⁴⁶⁶ Wright, B. W.; Synovec, R. E. Anal. Chem. **2006**, 78, 5068–5075.
- 467 (17) Beckstrom, A. C.; Humston, E. M.; Snyder, L. R.; Synovec, R. E.; Juul, S. E. J.
 468 Chromatogra. A 2011, 1218, 1899–1906.
- (18) Christensen, J. H.; Tomasi, G. J. Chromatogr. A 2007, 1169, 1–22.
- (19) Parsons, B. A.; Marney, L. C.; Siegler, W. C.; Hoggard, J. C.; Wright, B. W.; Synovec, R. E. Analytical chemistry 2015, 87, 3812–3819.
- 472 (20) Parsons, B. A.; Pinkerton, D. K.; Wright, B. W.; Synovec, R. E. J. Chromatogr. A
 473 2016, 1440, 179–190.
- ⁴⁷⁴ (21) Sinkov, N. A.; Sandercock, P. M. L.; Harynuk, J. J. Forensic Sci. Int. **2014**, 235, 24–31.
- 475 (22) Sinkov, N. A.; Harynuk, J. J. Talanta **2011**, 83, 1079–1087.
- 476 (23) Sinkov, N. A.; Harynuk, J. J. Talanta 2013, 103, 252–259.
- 477 (24) Adutwum, L.; Harynuk, J. Anal. Chem. 2014, 86, 7726–7733.
- 478 (25) Watson, N. E.; VanWingerden, M. M.; Pierce, K. M.; Wright, B. W.; Synovec, R. E.
 479 J. Chromatogr. A 2006, 1129, 111–118.
- 480 (26) Baz-Lomba, J. A.; Reid, M. J.; Thomas, K. V. Anal. Chem. acta 2016, 914, 81–90.
- 481 (27) MATLAB version 9.1 Natick, Massachusetts: The MathWorks Inc.,
- 482 (28) Nadeau, J. S.; Wright, B. W.; Synovec, R. E. *Talanta* **2010**, *81*, 120–128.
- 483 (29) Katajamaa, M.; Miettinen, J.; Orešič, M. *Bioinformatics* **2006**, *22*, 634–636.
- (30) Brereton, R. G. Applied chemometrics for scientists; John Wiley & Sons, 2007.
- 485 (31) Burke, D. S.; Brundage, J. F.; Redfield, R. R.; Damato, J. J.; Schable, C. A.; Put-
- 486 man, P.; Visintine, R.; Kim, H. I. N. Engl. J. Med. **1988**, 319, 961–964.



for TOC Only