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¹ Enumeration of Labile Hydrogens in Natural Organic Matter by Use ₂ of Hydrogen/Deuterium Exchange Fourier Transform Ion Cyclotron 3 Resonance Mass Spectrometry

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 - Supporting Information

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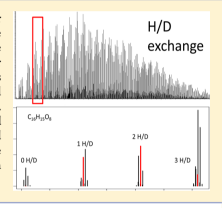
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ABSTRACT: A method to enumerate labile hydrogens in all constituents of molecular ensemble of natural organic matter (NOM) based on our previously developed simple hydrogen/deuterium (H/D) exchange (electrospray ionization (ESI) ion source (Kostyukevich et al. Anal. Chem. 2013, 85, 5330) and ultra-high-resolution Fourier transform ion cyclotron resonance mass spectrometry is presented. The method was applied for analysis of Suwannee River fulvic acid (SRFA), which is an International Humic Substances Society standard, as well as Siberian crude oil; and lignosulfonate. We found that SRFA and lignosulfonate molecules contain 2-5 labile hydrogens, and their number increases with the number of oxygens in the molecule. Also, we observed that compounds of Siberian crude oil ionizing in positive-ESI mode do not have labile hydrogens, while compounds ionizing in negative-ESI mode have one labile hydrogen that detaches during ESI ionization.



Tatural organic matter (NOM) presents a complex mixture of degraded natural organic compounds that plays an 26 important role in carbon cycling. However, molecular structures 27 of NOM remain largely unknown. This is because the extreme 28 complexity of NOM renders impossible their separation into 29 individual compounds by conventional techniques such as liquid 30 chromatography, electrophoresis, or isoelectric focusing. The 31 same is true for structural studies of NOM by powerful physical 32 methods that work perfectly for monomolecular samples (e.g., 33 NMR spectroscopy, soft X-ray spectroscopy, fluorescence 34 polarization) but are not capable of revealing the individual 35 structures of NOM despite many efforts.^{2,3}

Fourier transform ion cyclotron resonance mass spectrometry 37 (FTICR-MS) proved to be an important and useful tool for 38 investigation of NOM on the molecular level, making it possible 39 to determine accurate masses and elemental compositions of all 40 compounds, 4,5 perform fragmentation studies of isolated mole-41 cules, 6 and implement gas-phase reactions inside the FTICR cell 42 with previously separated compounds. 7,8

In mass spectrometric studies, one of the most important 44 methods of obtaining chemical and structural information about 45 a molecule (from metabolites to proteins)^{9,10} is based on the 46 exchange of labile hydrogens for deuteriums. Labile hydrogens 47 belonging to functional groups such as amide, carboxyl, or 48 hydroxyl can be easily replaced by deuterium in solution or in the

gas phase. At the same time, hydrogens attached to the carbon 49 backbone may be exchanged only under chemical ionization 50 conditions.1

As a consequence, the number of hydrogens exchanged by 52 deuterium in a molecule under soft experimental conditions 53 might serve as an equivalent to the number of the corresponding 54 functional groups containing labile hydrogens in this molecule. 55 Thus, simultaneous determination of labile hydrogens in all 56 compounds of NOM might yield chemical and structural infor- 57 mation about different molecules of a complex mixture.

Labile hydrogens in the large class of NOM belong to car- 59 boxylic and hydroxylic groups, and their determination by con- 60 ventional techniques based on exchange in liquid phase is impos- 61 sible due to a fast back-exchange in the ionization source. 9,10,12 62 Gas-phase exchange based on the infusion of deuterated gaseous 63 phase (such as D₂O vapor or ND₃) in the FTICR cell is difficult 64 to implement on the arbitrary FTICR and requires long-lasting 65 postexperimental cleaning.

Here we report a use of the simple in-source H/D exchange 67 method, previously developed in our group, combined with ultra- 68 high-resolution FTICR mass spectrometry¹² for simultaneous 69

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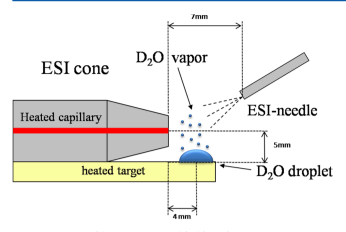


Figure 1. Design of the ESI source modified for H/D exchange experiments.

70 determination of labile hydrogens in all compounds composing 71 NOM. We investigated Suwannee River fulvic acid (SRFA),

which is an International Humic Substances Society standard, as 72 well as Siberian crude oil and lignosulfonate. Those substances 73 correspond to different degrees of NOM transformation in the 74 environment.

76

METHODS

Sample Preparation. SRFA was dissolved in methanol 77 (normal or deuterated) and diluted to a concentration of 1 g/L. 78 Siberian oil was first dissolved in toluene and then diluted in 79 methanol (MeOH or MeOD). Final concentration of oil was 80 1 g/L. Solvent composition was 30% toluene and 70% methanol. 81 Lignosulfonate was dissolved in water (normal or deuterated) 82 and diluted to a concentration of 1 g/L. Solvent composition was 83 50% water and 50% methanol.

Fourier Transform Ion Cyclotron Resonance Mass 85 Spectrometry. All experiments were performed on LTQ FT 86 Ultra (Thermo Electron Corp., Bremen, Germany) equipped 87 with a 7 T superconducting magnet. The obtained resolving 88

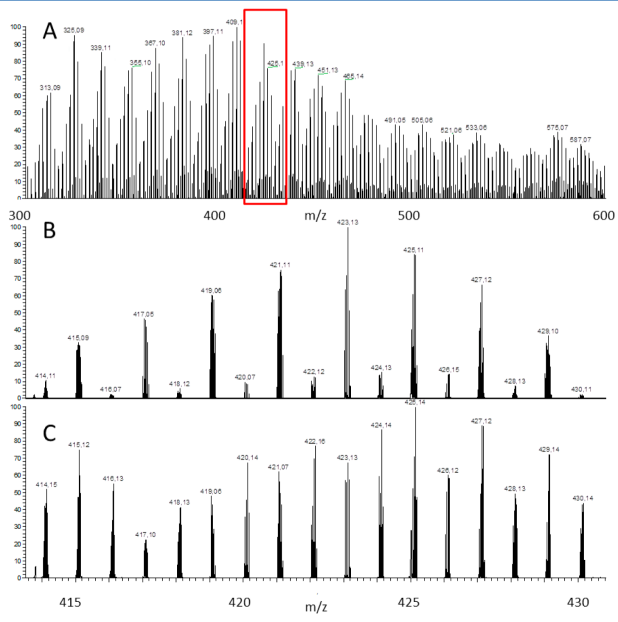


Figure 2. (A) FTICR mass spectrum of SRFA. (B) Magnified portion of FTICR mass spectrum of SRFA. (C) Magnified portion of FTICR mass spectrum of SRFA after H/D exchange.

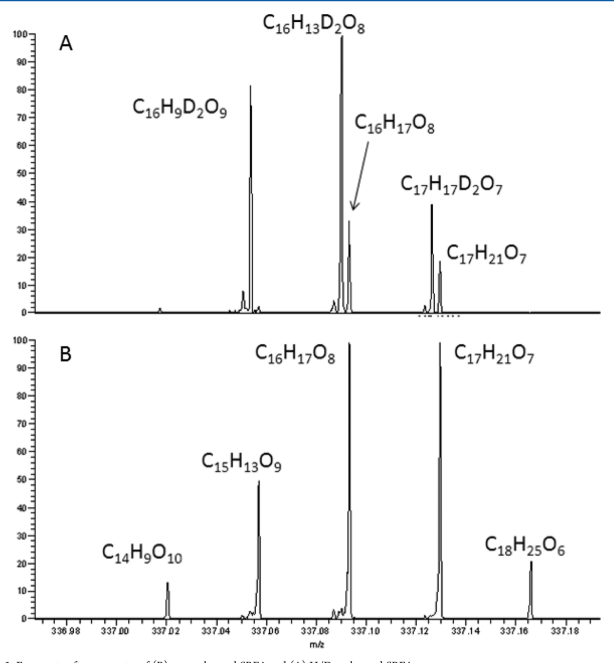


Figure 3. Fragments of mass spectra of (B) nonexchanged SRFA and (A) H/D-exchanged SRFA.

89 power was 400 000 at m/z = 400. Each spectrum was a sum of 400 90 consecutive scans. Ions were generated by IonMax Electrospray 91 ion source (Thermo Electron Corp., Bremen, Germany) in 92 positive and negative electrospray ionization (ESI) mode.

H/D Exchange Reaction. The design of the modified ESI 94 source for H/D exchange experiments is presented in Figure 1. A 95 copper plate was installed just beneath the ESI cone and a droplet 96 of D_2O was placed on it. The desolvating capillary was 97 maintained at a temperature of 250 °C, and the needle voltage 98 was 2400 V. Due to evaporation of the droplet, an atmosphere of 99 D_2O vapor in the region between the ESI needle and the MS 100 entrance capillary is created. D_2O molecules can penetrate ESI 101 droplets or participate in ion—molecular reactions inside the 102 desolvating capillary or in the vacuum capillary skimmer region. 103 As a result of these processes, H/D exchange takes place. The 104 relative intensity of the peak corresponding to n exchanges with 105 depth of exchange P equals

$$h(n) = C_N^{\ n} P^n (1 - P)^{N-n} \tag{1}$$

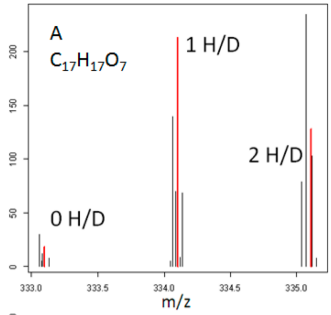
Here N is the total number of labile hydrogens. Mass difference 107 between peaks in the series formed by incomplete H/D exchange 108 reaction of the same parent ion equals 1.006 277 Da. 109

■ RESULTS AND DISCUSSION

Suwannee River Fulvic Acid. The mass spectrum of SRFA 111 is presented in Figure 2A. It can be seen that the spectrum (just as 112 of any other NOM) is very complicated and contains thousands 113 of peaks. Elemental compositions of SRFA molecules that are 114 ionized in negative-ESI mode can be presented as $C_cH_hO_o$. 115 A magnified portion of the SRFA mass spectrum is presented in 116 Figure 2B. Major peaks correspond to monoisotopic com- 117 pounds, and minor peaks correspond to ^{13}C isotopologue peaks. 118

The observed changes in mass spectra of the nonexchanged 119 sample and of deuterated SRFA are represented in Figure 2 B,C 120

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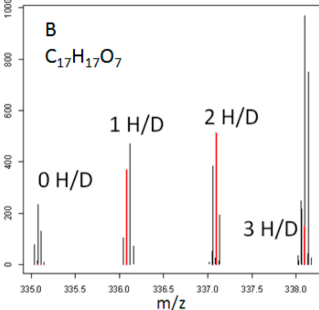


Figure 4. Distribution of H/D exchange peaks for SRFA compounds: (A) $C_{17}H_{17}O_7$ and (B) $C_{16}H_{16}O_8$.

 121 and in Figure 3. It can be seen that, by use of ultra-high-resolution 122 FTICR-MS, it is possible to unambiguously determine peaks 123 corresponding to the H/D exchange reaction of a certain parent 124 ion and quantitate those peaks.

125 H/D exchange of fast exchangeable hydrogens is always 126 incomplete because of the interaction with traces of atmosphere 127 water. As a consequence, the mass spectrum of H/D-exchanged 128 SRFA becomes much more complicated as compared to the 129 original one. We observed a factor of 3—4 increase in the number 130 of peaks that can be reliably identified.

Filtration and extraction of peaks corresponding to H/D sechange reaction of the same parent ion is a challenging problem due to the shifting of peaks and disappearance of small peaks in the mass spectrum. To enumerate labile hydrogens in SRFA, we used the following procedure. First we observed that the parent (nondeuterated ion) is not always present in the H/D exchange mass spectrum, but the peak corresponding to one

H/D exchange can be identified clearly. So for known elements in 138 SRFA, we identified peaks corresponding to one H/D exchange, 139 and for those peaks we extracted peaks with mass differences close 140 to the mass difference of the H/D exchange reaction. Then those 141 series were manually analyzed (see Supporting Information) to 142 identify the number of labile hydrogens in each parent ion.

The peaks corresponding to the H/D exchange reaction of 144 arbitrarily chosen compounds of SRFA are shown in Figure 4. It 145 can be seen that isotopic distribution obeys distribution. Relative 146 intensities of peaks corresponding to the H/D exchange reaction 147 of several compounds are presented in Table 1. We calculated the 148 depth of H/D exchange reaction by use of the following formula: 149

$$P = \frac{1}{N} \frac{\sum_{i=0}^{N} i h_i}{\sum_{i=0}^{N} h_i}$$
 (2) ₁₅₀

Table 1. Number of Labile Hydrogens Observed in Different Compounds of SRFA

brutto formula a	$\mathrm{D0}^{b}$	D1	D2	D3	D4	d^c	P^d
$C_{16}H_{15}O_{7}$	2	27	10	0	0	3	0.6
$C_{16}H_{17}O_{7}$	5	81	37	0	0	6	0.63
$C_{16}H_{19}O_7$	17	180	75	0	0	3	0.6
$C_{17}H_{17}O_7$	3	46	29	0	0	3	0.66
$C_{17}H_{19}O_7$	10	113	50	0	0	4	0.61
$C_{18}H_{19}O_7$	6	41	16	0	0	5	0.58
$C_{15}H_{13}O_{8}$	0	26	78	36	0	4	0.69
$C_{14}H_{15}O_9$	0	83	256	71	0	1	0.65
$C_{16}H_{13}O_{8}$	0	22	65	18	0	3	0.66
$C_{15}H_{15}O_9$	3.3	155	232	82	0	3	0.61
$C_{16}H_{13}O_{9}$	0	16	67	44	0	6	0.74
$C_{14}H_{11}O_9$	0	13	60	93	21	15	0.66
$C_{14}H_{11}O_{10}$	0	10	53	73	50	9	0.72

^aElemental composition. ^bIntensity of peak corresponding to H/D exchange (D1, one exchange; D2, two exchanges, etc.). ^cMaximum mass difference between neighbor peaks and mass difference corresponding to H/D exchange (10⁻⁵ amu). ^dDepth of exchange.

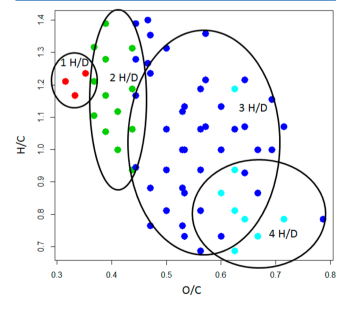


Figure 5. Van Krevelen diagram for reliably investigated SRFA compounds: (red) one H/D exchange, (green) two H/D exchanges, (dark blue) three H/D exchanges, (light blue) four H/D exchanges. The number of labile hydrogens in the molecule is bigger by 1 because ions were generated in negative mode and one hydrogen detached during ionization.

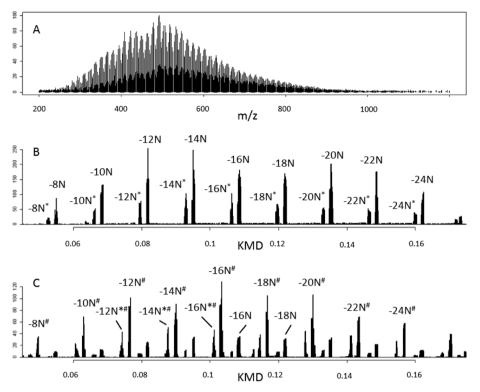


Figure 6. (A) Positive ESI FT ICR mass spectrum of Siberian crude oil. (B, C) Weighted Kendrick mass defect histograms for (B) normal and (C) H/D exchange conditions. (*) ${}^{12}C_{c-1}{}^{13}C_1$ species; (#) compounds with one H/D exchange that belong to the same homologous series.

151 Here h_i is the intensity of the peak corresponding to the ith H/D 152 exchange and N is the total number of labile hydrogens.

In Figure 5 is presented the Van Krevelen diagram for reliably investigated parent ions. We can see the trend that the number of labile hydrogens increases with the number of oxygens in the molecule.

The latter trend and the fact that only hydrogens attached to 157 carboxyl and hydroxyl groups are labile under soft H/D 158 experimental conditions (in ESI source) 12 allow us to surmise 159 that we can obtain information about types of oxygen in the 160 molecule by comparing the number of labile hydrogens with the 161 total number of oxygens. For example, the compound $C_{17}H_{17}O_{7}$ 162 has three labile hydrogens. Even if all these hydrogens were from 163 carboxyl groups (-COOH), there is still one oxygen left and it 164 must stem from other functional groups, such as carbonyl or 165 ether group or others.

Crude Oil. Crude oil is another example of very complex natural organic matter (Figure 6A). Unlike dissolved organic matter, crude oil is composed of long homologous series that make it possible to implement a compact visual analysis method based on Kendrick mass defect approach. The elemental composition of a crude oil molecule can be represented as $C_c H_{2c+Z} N_n O_o$, and then a homologous series formed by $C_c H_{2c}$ can be referred to as $ZN_n O_o$. It is convenient to introduce a Kendrick mass cale by setting the mass of $C_c H_{2c}$ fragment (14.015 65) to equal 14:

$$Kendrick mass = \frac{(IUPAC mass)(14)}{14.01565}$$
(3)

176 Compounds with the same constitution of heteroatoms will 177 have identical Kendrick mass defect (KMD):

175

178

$$KMD = nominal Kendrick mass - exact Kendrick mass$$
(4)

179 If one plots KMD versus mass, then compounds belonging to the 180 same homologous series forms a straight horizontal line on the

plot. This approach works perfectly for general analysis of crude 181 oil, but for H/D exchange experiments it is more convenient to 182 introduce a weighted Kendrick mass defect histogram. To create 183 a weighted Kendrick mass defect histogram, one must first 184 calculate Kendrick mass defect for all compounds and then plot a 185 histogram where the number of entries of each KMD is pro- 186 portional to the intensity of the corresponding peak.

In Figure 6B,C are presented weighted Kendrick mass defect 188 histograms for crude oil under normal and H/D exchange 189 conditions. Ions were generated in positive-ESI mode. We see 190 that major homologous series are compactly represented, and for 191 each series we observe a series corresponding to the isotope 192 exchange $^{12}\mathrm{C} \rightarrow ^{13}\mathrm{C}$. Under H/D exchange conditions we see 193 that all series exchanged one hydrogen for deuterium. In 194 positive-ESI mode, the molecule is ionized by attaching a 195 proton, and the fact that we observe only one H/D exchange 196 for all molecules indicates that all molecules in the fraction 197 ionizing in positive-ESI mode do not have labile hydrogens. 198 Only the ionizing proton exchanges.

We did not observe any differences in negative-ESI mass 200 spectra of crude oil under normal and H/D exchange conditions. 201 It proves that all molecules in this fraction have exactly one labile 202 hydrogen and this hydrogen detaches during ionization. 203

Lignosulfonate. Previously investigated compounds (SRFA 204 and crude oil) are products of the postmortem transformation of 205 living organic matter. Lignosulfonate is a derivative of lignin, one 206 of the most abundant organic polymers on Earth, ¹⁷ constituting a 207 quarter to a third of the dry mass of wood. Lignosulfonate is also a 208 complex mixture (see Figure 7A) containing many individual 209 molecules. The elemental composition of the lignosulfonate 210 molecule can be represented as $C_c H_h O_o S_s$. By the H/D exchange 211 approach, it is possible to enumerate labile hydrogens in 212 individual compounds. For example, we can see that compound 213 $C_{10}H_{13}O_8S_2$ contains exactly three labile hydrogens.

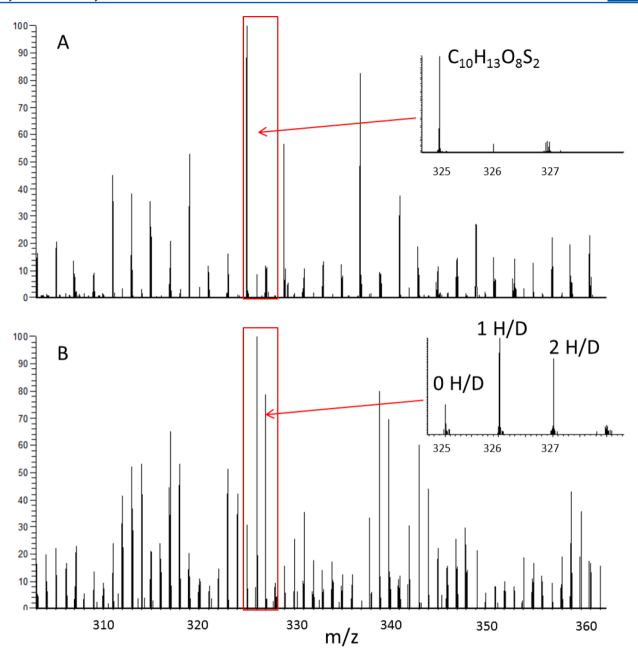


Figure 7. (A) FTICR mass spectra of lignosulfonate under (A) normal and (B) H/D exchange conditions (negative-ESI mode).

215 CONCLUSION

216 We demonstrate here the possibility to enumerate labile 217 hydrogens simultaneously in all compounds of natural organic 218 matter by using a simple experimental technique. Fulvic acids, 219 crude oil, and lignosulfonate represents different degrees of 220 transformation of living organic matter in nature. H/D exchange 221 FTICR mass spectrometry is capable of bringing new insights 222 into the investigation of carbon cycling in the Earth. Recent 223 advances in FTICR, such as increased resolving power and 224 dynamic range, ^{18,19} will be of great help for the further development of this technique.

226 ASSOCIATED CONTENT

Supporting Information

228 Manual analysis of mass spectral peaks as described in the text. 229 This material is available free of charge via the Internet at http:// 230 pubs.acs.org.

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The manuscript was written through contributions of all authors. 235

All authors have given approval to the final version of the 236

manuscript. 237

Notes 238

The authors declare no competing financial interest. 239

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