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Development in aquatic humic chemistry

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Abstract – There are always specific achievements which affect scientific work. This also applies to the field of aquatic humic chemistry. Some of the milestones were: (1) the standardization of isolation procedures for humic substances and by this the availability of reference materials; (2) the application of powerful tools of NMR spectroscopy and mass spectroscopy in combination with pyrolysis for the identification of molecular building blocks; (3) the application of gel chromatography with multidetection, including vibrational and electronic spectroscopy and on-line measurement for direct characterisation of organic matter in aquatic samples; (4) the application of static and time-resolved fluorescence to study excited states and complex formation; (5) the investigation of reactions with well defined partners; and (6) the development of computer models for energy minimized molecular structures.

Résumé – Développements en chimie des substances humiques aquatiques. Le travail scientifique dépend toujours de savoirs spécifiques déjà acquis. Ceci s’applique notamment au domaine de la chimie des substances aquatiques. Certaines étapes importantes ont été : (1) la standardisation des procédés d’isolement des substances humiques et avec elle la disponibilité de substances de référence ; (2) l’application des outils puissants de la spectroscopie RMN et de la spectrométrie de masse combinés à la pyrolyse pour identifier les blocs moléculaires constitutifs ; (3) l’application de la chromatographie sur gel avec détection multiple y compris la spectroscopie vibrationnelle et électronique et la mesure en ligne pour la caractérisation directe de la matière organique dans les échantillons aquatiques ; (4) l’application de la fluorescence statique et résolue en temps pour étudier les états excités et la formation de complexes ; (5) l’étude des réactions avec des partenaires bien définis et (6) le développement des modèles informatiques pour les structures moléculaires dont le niveau d’énergie est minimum.

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1. Introduction

Humic matter is as old as life on earth. As a sink of organisms and a source for new living cells it plays a fundamental role in the environment. Already at the end of the 18th century, Achard [3] investigated peat and extracted soil with base. Berzelius [8] isolated humic substances from a Swedish spring. Only since about the beginning of the 20th century has the scientific interest for these substances emerged in a broad scale [18, 33, 47, 75]. It was the field of soil and agriculture which stimulated investigations of the structure, properties and function of the brownish heterogeneously compositied organic material [21, 25, 38, 61, 69, 70, 79].

Even though it was clear from the beginning that the water soluble part of the material was the most interesting one for reactions and transport phenomena, it was not until the second half of the century that systematic work began on aquatic humic matter [36, 51, 68]. Since then, it has become widely accepted that humic substances are ubiquitous in water [73], and a great interest in their chemical structure and role in aquatic systems has developed [22], including technical applications [71]. In 1983, the International Humic Substances Society (IHSS) was founded and members have collected data and results on refractory matter in soil and water [5, 13, 17, 29, 65]. The term “refractory” commonly reflects a relatively low biodegradability, but it has not been defined properly. The number of publications on aquatic humic substances has increased significantly in recent years. It is the aim of this paper to give an overview on important milestones in aquatic humic chemistry, and on the consequences for scientific progress in this field.

2. Definitions

According to the complex structure of humic substances (HS) they can not be defined precisely. Based on fractionation procedures and on specific research views, operational definitions are often used. Some common terms are given in Table I.

<table>
<thead>
<tr>
<th>Abbreviation</th>
<th>Meaning</th>
</tr>
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<tbody>
<tr>
<td>AHS</td>
<td>Aquatic Humic Substances</td>
</tr>
<tr>
<td>DOM</td>
<td>Dissolved Organic Matter</td>
</tr>
<tr>
<td>FA</td>
<td>Fulvic Acids (acid and base soluble)</td>
</tr>
<tr>
<td>HA</td>
<td>Humic Acids (base soluble)</td>
</tr>
<tr>
<td>HS</td>
<td>Humic Substances</td>
</tr>
<tr>
<td>NHS</td>
<td>Non Humic Substances</td>
</tr>
<tr>
<td>NOM</td>
<td>Natural Organic Matter</td>
</tr>
<tr>
<td>ROM</td>
<td>Recalcitrant Organic Matter</td>
</tr>
<tr>
<td>ROS</td>
<td>Refractory Organic Substances</td>
</tr>
<tr>
<td>SOM</td>
<td>Soil Organic Matter</td>
</tr>
<tr>
<td>DOC</td>
<td>Dissolved Organic Carbon (&lt; 0.45 µm)</td>
</tr>
<tr>
<td>TOC</td>
<td>Total Organic Carbon</td>
</tr>
<tr>
<td>POC</td>
<td>Particulate Organic Carbon</td>
</tr>
</tbody>
</table>

It is necessary to note that some of the terms are synonymous. Since most terms are operationally defined, they need a detailed description of the experimental procedure of their isolation, preparation and origin.

3. Isolation methods

The concentrations of organic carbon (OC) in aquatic systems range between 40 mg·L⁻¹ in brown water and soil seepage water and 0.5 mg·L⁻¹ typical for ground water and sea water [73]. These concentrations are too low for a direct application of most of the advanced analytical methods. In addition, matrix effects from high salt content or metal ions can lead to severe disturbances in structural investigations. Therefore isolation and concentration procedures are needed. The adsorption at low pH-values (pH ≤ 2) on synthetic resins like XAD 2 or XAD 8 followed by desorption at high pH-values (pH > 12) [48, 56] turned out to be a useful method. The IHSS recommended standard procedures for preparative isolation of HS [72] and supplied reference materials [46]. The principle of the isolation method has been applied in many variations, leading to new operationally defined products and fractions [1, 6, 45]. A scheme for the fractionation into so called hydrophobic and hydrophilic materials is shown in Figure 1.
The standardized methods for the isolation of HS and the availability of reference samples have stimulated the investigation of the structure and character of the material and have led to the intercalibration of analytical methods and by this to meaningful results. Even though there is no doubt about the merits of this approach, some pitfalls have to be kept in mind. The treatment of the water samples including high and low pH-values may result in structural changes. Especially when oxygen is not excluded, oxidation reactions are very likely to occur. To our knowledge, however, there are no publications on a reliable comparison of extraction procedures under oxic and anoxic conditions. There are other fundamental questions: how good the obtained fractions represent the natural situation, and whether the eliminated other water constituents influence the typical character of HS or not. A way out of that dilemma seem to be proper mass balances, e.g. on the basis of OC, and the development of sensitive methods which are suited for a direct application to the unchanged original aquatic samples. Other isolation and concentration methods like reverse osmosis or nanofiltration have also been introduced [1, 67].

4. Vibrational and electronic spectroscopy

The yellowish-brownish colour of HS is one of their typical properties [51]. The spectral absorption in the UV-range is even more pronounced.

Figure 1 shows a characteristic spectrum of a brown water. The absorbance decreases with increasing wavelength in a roughly exponential way and can be measured directly without any pretreatment of the water.

Even though the UV-vis spectra of HS are not too informative, they reflect the valence electrons and unsaturated character of the structural building blocks, especially if differential spectroscopy is used [39]. Related to the OC, the UV absorbance (e.g. specific UV absorbance SUVA at λ = 254 nm) can give a good estimation of the reactivity of the material against chlorine. This has become a valuable tool to predict the amount of by-product formation in the disinfection of drinking water.

Another specific property of HS is their fluorescence. This was shown by several researchers [12, 19, 66]. According to the Jablonski diagram, which is a simplified illustration of the processes taking place during absorption of radiation and relaxation, one way of obtaining information on the excited electronic states of a molecule is the recording of fluorescence spectra. Molecular differences can be most clearly seen in the synchronous mode [10]. Since complex formation of HS with paramagnetic metal ions and pollutants like polycyclic aromatic hydrocarbons (PAH) lead to a quenching of the original HS fluorescence, this can be used to learn about molecular interactions and electron donor-acceptor complexes [7, 23, 58]. Due to the sensitivity of fluorescence spectroscopy these investigations again can be done directly in aquatic samples.
recently time-resolved fluorescence spectroscopy was applied to aquatic HS. The principle of the method is given in Figure 3. It can be shown that there are at least three excited states which decay in different time windows of the nanosecond range (Fig. 4) [41].

As expected, chemical reactions like complex formation and oxidation change the decay rates and the distribution of the components. With an increased number of investigated samples this will lead to a better understanding of the reactivity of HS even though their molecular structure still remains unknown.

5. Solid sample investigation

Freeze drying of aquatic samples or aqueous fractions of isolates leads to solid HS samples. The freeze drying procedure is one of the most gentle methods for preparation, and opened the application of a number of powerful analytical tools like analytical pyrolysis (Py) coupled with mass spectrometry (MS) or gas chromatography/MS and solid state nuclear magnetic resonance (NMR). The application of those methods has led to information on building blocks and structural sub-units. However, the question remains open as to what extent the molecular structure revealed for the solid state is representative of the dissolved one.

5.1. Analytical pyrolysis

Oven pyrolysis in combination with field ionization mass spectrometry (FIMS) has turned out to be a useful tool for supplying ideas about possible building stones of HS in the range of up to a few hundred mass units. Especially, the high resolution mode for mass units gives reasonable suggestions for molecular structures including hetero-atoms of the pyrolysis product. A flexible temperature program and the temperature correlated product pattern open a wide field for macromolecular structure suggestions [62, 63]. A general scheme of PY-FIMS is given in Figure 5. Pitfalls come from a still poor understanding of the pyrolysis process and from uncertainties about the yield of the sample-specific products. A lack of oxygen in the products found

Figure 3. Principle of time-resolved fluorescence measurement of dissolved humic matter (flash lamp L; monochromator M; sample S; detector D).

Figure 4. Multi-component analysis (MCA) for HS regions with different fluorescence decay rates.

\[
\frac{I_F^0}{I_F} = 1 + K_{SV} \cdot [Q]
\]

\[
\frac{I_F^0}{I_F} = \text{DOC (mg/L)}
\]

\[
\text{pH 4}
\]

\[
\text{pH 11}
\]
compared with the amount shown by elemental analysis of the gross sample and an overestimation of aromatic structures are known problems.

Pyrolysis in combination with GC/MS is another method for the identification of molecular fragments. The volatilisation of parts of the macromolecules is normally done by Curie point pyrolysis which works in the millisecond-range. The products obtained (see Tab. II) are fractionated by gas chromatography and identified by MS [2, 9, 59].

5.2. NMR-spectroscopy

Nuclear magnetic resonance methods belong to the nondegradative analytical procedures and have been applied successfully for the characterization of complex samples and macromolecules [77]. The necessary masses and concentrations of samples have decreased significantly over the last few years, but are still far from original aquatic samples. Besides $^1$H and $^{13}$C heteroatoms [27, 28, 43, 77], especially $^{15}$N and $^{31}$P [50, 54] have been investigated. Dissolved samples and solid state material were used.

In solid state NMR, magic angle spinning (MAS) and cross polarization (CP) have significantly improved the results obtained for HS samples [44]. Recently, the distribution and molecular form of heteroatoms have been visualized by two dimensional modes [31], and derivatisation reactions like silylation in combination with NMR have led to information on structure reactivity relations [30]. This makes the NMR technique one of the most informative methods for the characterization of the composition of HS.

The relative amount of carbon assigned to structural areas and functional groups in brown water FA are given in Table III.

The NMR-data are well suited for correlation with the results from other analytical methods, and as a consequence for intercalibration and quality control of analytical determinations.

<table>
<thead>
<tr>
<th>Class of compounds</th>
<th>Identified compounds</th>
</tr>
</thead>
<tbody>
<tr>
<td>alkenes</td>
<td>$C_4$-$C_8$ alkenes, $C_4$-$C_8$ alkadienes</td>
</tr>
<tr>
<td>furans</td>
<td>methylfuran, hydroxyfuran, methylfurural</td>
</tr>
<tr>
<td>alkylbenzenes</td>
<td>toluene, $C_2$-alkylbenzene, $C_6$-alkylbenzene</td>
</tr>
<tr>
<td>alkylphenols</td>
<td>cresol, xylene, methoxyphenol</td>
</tr>
<tr>
<td>dihydroxybenzenes</td>
<td>dihydroxybenzene, dihydroxymethylbenzene</td>
</tr>
<tr>
<td>methoxyphenols,</td>
<td>2,6-dimethoxyphenol, coniferyl alcohol, 2,6-dimethoxy-phenyl-phenol, sinapyl alcohol</td>
</tr>
<tr>
<td>dimethoxyphenols</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Table III. Relative amounts of differently assigned C of HO13 FA as revealed by $^{13}$C-CPMAS-NMR [51].</th>
</tr>
</thead>
<tbody>
<tr>
<td>Functionality</td>
</tr>
<tr>
<td>-----------------------</td>
</tr>
<tr>
<td>Aliphatic</td>
</tr>
<tr>
<td>Overlap Region</td>
</tr>
<tr>
<td>O-Allyl-C</td>
</tr>
<tr>
<td>Aromatic</td>
</tr>
<tr>
<td>Unsubstituted</td>
</tr>
<tr>
<td>O, N-Substituted</td>
</tr>
<tr>
<td>Carboxyl</td>
</tr>
<tr>
<td>Carbonyl</td>
</tr>
</tbody>
</table>
6. Molecular size estimation

Among the principles for determination of molecular sizes or masses, gel permeation is an attractive method. With powerful detection systems it can be applied to original aquatic samples. Keeping in mind the limitation of interpretation due to undefined sorption effects and the lack of authentic calibration substances, the method is especially suited for comparative studies. In former years the most common stationary phase was Sephadex, and only UV detectors provided a satisfactory power for the direct investigation of aquatic samples [24]. In recent years, TSK columns with higher resolution have become available. TSK-gels of the HW and PW type are hydrophilic copolymers made from ethylene glycol and methacrylate [74]. Furthermore, a sensitive detection system for organic carbon was developed. This system can quantify fractions representing a few µg·L⁻¹ OC and by this is well suited for exact mass balances. Figure 6 shows the scheme of the chromatographic column and the UV/vis-detector followed by the OC detection system. It contains a thin-film UV-reactor which after the exclusion of the inorganic carbonate species oxidises the organic substances to CO₂ which is quantified in a non-dispersive IR-detector. UV leach reactors are used for the production of a low carbon water as a mobile phase [34]. The system works with original water samples of a volume up to 2 mL.

A typical gel chromatogram of a brown water is given in Figure 7. It shows the high molecular mass fraction assigned to HS at short retention time (t_R) followed by small mono- and diprotic acids. At higher t_R, hydrophobic smaller molecules appear.

There have been several authors addressing the errors involved in deriving molecular weight data for HS from gel chromatography. Perminova et al. [53] suggested a method which uses the additional structural data of HS to obtain more reliable molecular weight numbers for HS. Besides its suitability for weight estimation, gel chromatography is well suited for a general characterization of the DOC of aquatic systems and for the comparison of NOM from different sources [32]. The running conditions guarantee minimized influences on the identity of the samples.

7. Reactions and interactions

Due to the complex composition of HS it is highly questionable whether the identification of the molecular structure in the classical sense of analytical
chemistry will be ever achieved. A feasible method for obtaining information on the structure related reactivity is to react HS under well defined conditions with well defined substances to well defined products. This approach matches also the needs of a row of technical applications which are strongly influenced by HS.

7.1. Formation of AOX and THM and other disinfection by-products

Since the discovery of trihalomethanes (THM) as disinfection by-products in the chlorination of HS-containing water by Rook [57], the reactions of HS with chlorine and bromine have become a major health concern [35]. A model for the reaction of HS with chlorine was developed (Fig. 8), and far more than 150 reaction products have been identified [55]. Some compounds turned out to be mutagenic. The mutagenicity of MX and EMX (Fig. 9) was investigated by Kronberg [40].

7.2. Metal complexation

HS can act as electron- (e)-donor. In the presence of metals as e-acceptor, complexes are formed. The complex formation can be quantified by measuring the fluorescence intensity. According to the hypothesis that HS only fluoresce as uncomplexed ligand and that complexes with paramagnetic metal ions show no fluorescence, stability constants can be calculated [76]. The investigation of systematically varied donor/acceptor ratios can be used for a Stern-Volmer plot (Fig. 10). From the Stern-Volmer relation it can be seen that the complex formation is a static process which fixes the metal ions to specific ligand functional sizes. It is also interesting to see that Al(III) as a diamagnetic metal ion does not quench the fluorescence of the ligand. For most metal ions the experiments can be done in a concentration range typical for aquatic systems.

7.3. Interactions with xenobiotics

The structure of HS combines hydrophobic and hydrophilic regions. The resulting ambivalent character is the reason for solubility enhancement of lipophilic xenobiotics in water. Chiou et al. [11] showed that NOM from different origins have a different influence on the apparent solubility of DDT. Isolated HAs caused a stronger effect than river water and isolated FAs.

The interaction of HS with pyrene was investigated by Kumke et al. [42]. There was a clear visible effect when the fluorescence of the PAH was decreased in the presence of HS (Fig. 11). The interaction gained strength at lower pH-values.
Numerous articles on the direct and indirect photolysis of xenobiotics mediated by humic substances have been published, for example by Aguer et al. [4], Mills and Sullivan [49], Fischer et al. [20], and Cooper et al. [15].

7.4. Structural models

Chemists think in terms of molecules. This brought some active groups to the attempt to draw detailed molecular structures for humic matter [14, 16, 26, 37, 60]. Most of the models have been developed from the specific view of the investigation and by this reflect only specific properties of the matter. Some confusion arose from over-interpretations of these structures which should be seen as models giving not more than a possible suggestion and a reasonable explanation for measured properties of the samples. The operational definition of HS, their complex constitution and the manifold isomeric and energetically equivalent possibilities of structural arrangements make it highly unlikely that molecules of the given structures ever existed in humic material. As a consequence, models have been developed which do not use a detailed molecular structure but instead use symbols for building stones building blocks and functional regions within humic substances [52, 79]. Despite the confusion which arose from the misunderstanding of molecular structure models the positive aspect of the attempt to present measured data in a transparent form remains.
Recently a molecular structure has been proposed on the basis of force field calculations of lowest energy structures [64]. The availability of powerful data systems and programs for molecular calculations in connection with carefully measured data on well defined samples can give new impulse to the understanding of structure property relations in HS.

8. Where to go

The understanding of the function of ROS in nature is of fundamental importance. The water soluble part of it (DOM) plays a key role in this respect, for it is involved in the major reactions and transport processes. Further work should focus on:

1. Analytical multimethod characterization of typical samples from different origins to give a
comprehensive image of the molecular building blocks of the DOM;
2. Structure and physico-chemical property relations to understand dynamic processes in ecosystems;
3. Biochemical interaction of DOM with cells and enzymes to learn about genesis and the fate of ROS; this includes the colloidal range;
4. Formation and stability of ROS-bound xenobiotics including changed toxicity;
5. Environmental influences on the production and character of DOM.

Most of these topics need interdisciplinary approaches and new views. This may become obvious from such questions as what “dissolved” means, and how can we explain “refractory”. Operational definitions may help at the first stage, but molecular size distributions and lifetimes will have to follow.

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