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Pyrolysis-gas chromatography-mass spectrometry with electron-ionization and resonance-enhanced-multi-photon-ionization for the characterization of terrestrial dissolved organic matter in the Baltic Sea†

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With the effects of global warming the input of terrigenous material into the oceans is increasing, with unknown consequences for the ecosystem. The Baltic Sea is an ideal research object and observed effects can be transferred to the oceans. This paper combines the influence of biotic and abiotic factors especially for terrigenous dissolved organic matter (tDOM). The study is focused on specific lignin target molecules and reflects the influence of salinity and microbial activity. Samples were taken along the salt gradient. In addition, an incubation experiment, mixing of tDOM-rich river water with Baltic Sea water from three different stations, was carried out. A newly developed pyrolysis gas chromatography mass spectrometry (Py-GC/MS) method using two different mass selective analyzers in one measurement cycle was established for the analysis of tDOM species. It enables the characterization of natural samples by a universal (electron ionization quadrupole MS) as well as an aromatic fingerprint (resonance-enhanced-multi-photon-ionization time-of-flight MS). By thermal desorption (TD) and subsequent pyrolysis the free volatile and high molecular weight structures are accessible. A huge part of the chemical species exists as high molecular structures. The salt content has a high influence on the composition of DOM. Generally, under TD conditions greater changes were observed, especially for the incubation experiment. Under pyrolysis and the chosen experimental conditions, the lignin apparently is hardly degraded by microorganisms.

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1. Introduction

The climate change and the associated negative consequences for many ecosystems, particularly the oceans, are discussed worldwide. As a result of the global warming, the temperature in arctic regions increases and the rivers located there have shown trends of a higher water discharge.¹ With the enhanced inflow by Nordic rivers, the input of terrigenous material in the oceans

rises as well. The structural characterization of this material such as dissolved organic matter (DOM) and the understanding of its alterations while being transported to the oceans has been an increasing area of research over the past few decades.² Large fractions of terrigenous dissolved organic matter (tDOM) are removed from the river inflow areas towards the open oceans. Possible reasons are abiotic as well as microbial degradation.^{3–5} While abiotic factors such as salinity or solar radiation are studied in detail,^{6–10} the microbial degradation is poorly understood and previous experiments showed contrasting results.^{1,4,11–14}

Terrestrial DOM is a ubiquitous and dynamic mixture with a characteristic organic chemical signature for every individual region.^{15,16} A distinction between particulate organic matter (POM) and DOM is operationally defined. The latter is assumed to pass a 0.45 µm filter pore whereas POM is retarded. It is also a combination of refractory and labile marine and terrigenous compounds. The riverine organic matter can be soil-derived and mostly has a terrigenous origin.⁶ The input of tDOM by freshwater is considerably high,^{8,17} but only a small fraction (0.7–2.4%) of this introduced total DOM in the oceans is detectable.¹⁸

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The reason for this could be various elimination processes that are still unclear in detail. This is remarkable, because arctic tDOM seems to be relatively stable over time.^{19,20}

Lignin, as a part of tDOM, is a tracer for terrestrial organic matter, since it is source specific and produced only by vascular plants.^{2,13,16} The degradation products of lignin are phenolic monomers, which can be divided into four different structural families: *p*-hydroxy (H building block), syringyl (S building block), vanillyl and cinnamyl family.²¹ For simplification the last two are taken together as V building blocks, because they have the same aromatic base bodies. The S/V value is often used to distinguish between gymnosperm (non-flowering vascular plants) and angiosperm (flowering vascular plants) sources.^{13,21} A high amount of H building blocks often represents grass lignins,⁴ but the free forms can also have a marine origin.²²

The influence of an increasing salinity on high-molecular-weight-DOM and the alterations by flocculation, adsorption to suspended matter,²³ photochemical transformation^{7,9,24} and microbial decomposition¹⁰ are important removal mechanisms for tDOM on its way from the freshwater input to the oceans. Often a combination of biotic and abiotic factors is responsible for the degradation of DOM. Salinity can play an important role in enhancing the decomposition rate by bacterial communities.^{4,5,10} In general, the degradation rates of DOM vary tremendously, between 6 and 70%,^{1,14,25,26} depending on the experimental approach. Moreover, the temperature has a significant influence on the degradation of DOM.²⁷ It is claimed, that colloidal, high-molecular-weight-DOM is more biological and chemical reactive than under low temperature conditions.²⁸

In order to describe the role of tDOM, especially of lignin, its input, transformation and degradation, specific methods are needed to get more detailed information on the structure at a molecular level. In the ocean the concentration of DOM is extremely low with <1 mg L⁻¹.²⁹ Therefore the isolation and purification of DOM is important and often linked to the subsequent analysis. Retention-based methods are enriching the DOM and remove inorganic material (*i.e.* salt),^{28,30} which is essential for chromatographic methods. Generally gas chromatography (GC) in combination with alkaline cupric oxidation to yield specific lignin monomers^{11,13,21} or conversional pyrolytic degradation^{28,31} are established for characterizing lignin-deriving organics. By using a derivatisation reagent the number of compounds that are identifiable can be increased.^{4,8,21,32}

In this study, a GC separation followed by two distinct MS-methods in parallel was applied for the characterization of tDOM for the first time. The analytical system consists of a quadrupole and a time-of-flight MS (QMS and ToFMS) in combination with electron ionization (EI) and photo ionization (PI), respectively. EI, a hard ionization technique, leads to high fragmentation of the organic molecules. Despite the fact that the characteristic fragmentation pattern can be used for substance identification, sometimes the loss of the molecular ion aggravates the identification. The lack of the molecular ion signal can be prevented by using photo ionization such as

resonance-enhanced-multi-photon-ionization (REMPI), a soft ionization method. For a successful ionization *via* an intermediate state two photons have to be absorbed.^{33–35} REMPI has a high selectivity and sensitivity for aromatic hydrocarbons, depending on the used photon wavelength and additionally on the ionization energy threshold.^{36–38} The developed Py-GC/MS method with two different mass selective analyzers combined in one measurement cycle, enables the characterization of natural samples by a universal (EI) as well as an aromatic fingerprint (REMPI) and was successfully applied by Otto *et al.* (2015) to crude oil samples.³⁹ Therefore the aim of the present study was to characterize riverine DOM by the newly developed Py-GC/MS combination³⁹ and its structural and semi-quantitative compositional changes along the gradient of salinity and during decomposition by marine bacterial communities.

2. Experimental section

The experimental part is roughly divided into the sampling along the salt gradient of the Baltic Sea and the incubation experiment on three different stations.

2.1. Sample size and sampling

Samples were collected on a research cruise with the Meteor (M87/3a) between Skagerrak (S1 station) and the Gulf of Bothnia (At4 station) in May/June 2012. One part of the sampled water was taken for the salinity gradient measurements (see Fig. 1a).

Another important part of the study was the incubation experiment at three representative salinities. The stations are labeled with M1, M2 and M3 (identical with At1, At3 and At4). The selection of the relevant stations was based on the publication of Herlemann *et al.* (2011), where three suitable bacterial communities in the Baltic Sea, a marine/brackish group (salinity 8–32), brackish group (salinity 3–8) and a freshwater-brackish group (salinity < 3) were identified.⁴⁰ The microorganisms are able to use tDOM as the carbon source. In order to investigate the degradation of tDOM by microorganisms, tank experiments were conceived.

Kalix river water (located in North Sweden), which is rich in terrigenous dissolved organic material, was mixed with the sampled Baltic Sea water. The river water was sampled during the spring flood in May 2011 (see Herlemann *et al.* (2014) for detailed information), sterile-filtered and stored at –20 °C. For each sampling station, the river water was at first mixed with artificial seawater to reach the salt content of the current sample station, in order to limit the stress for the microorganisms. Finally, it was filled up with 50% surface water of the current station. This mixing tank contains 120 L water volume and is abbreviated as uRW-tank. In this study the mixed uRW tanks as well as a control tanks (cBS, Baltic Sea water only) were examined for three representative salinities. The reason for utilizing the cBS tank is the fact that Baltic Sea water contains natural DOM and microbial processes are still running. The composition of DOM is also changed. Therefore, the investigation of such changes by added artificial DOM through river water becomes necessary. Three biological replicates from each



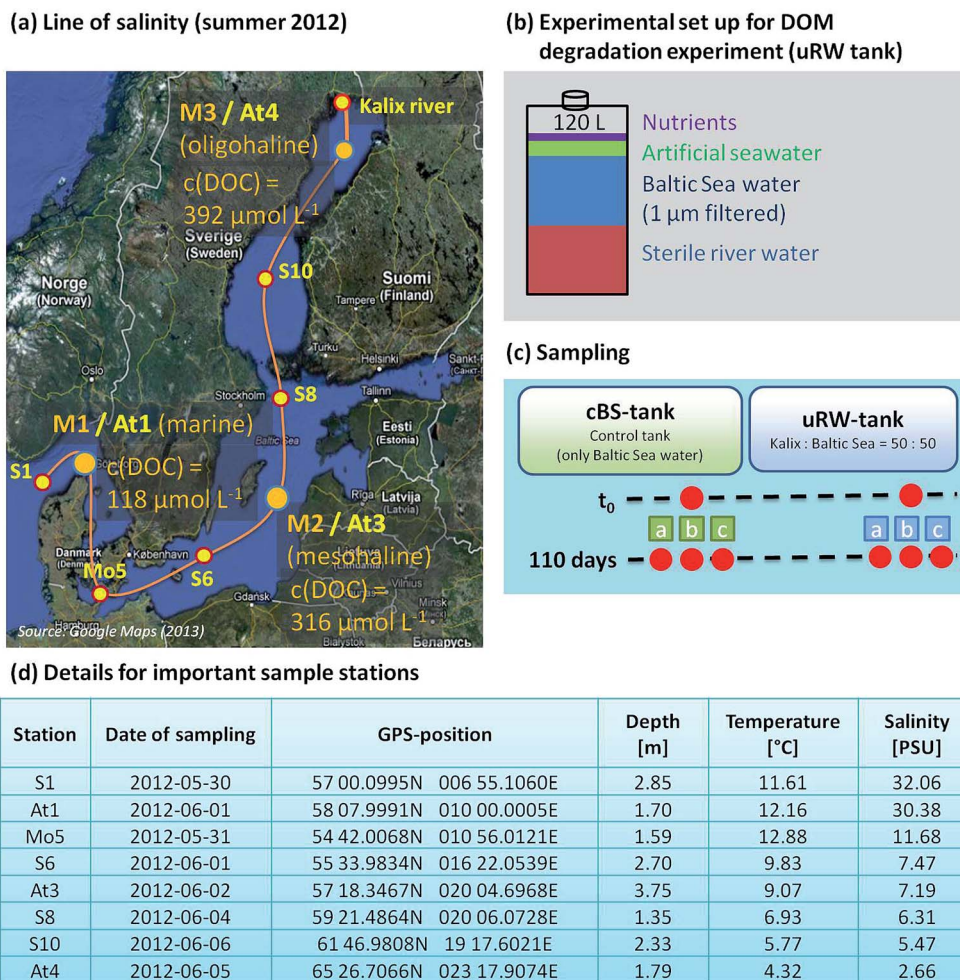


Fig. 1 (a) Map of the Baltic Sea displaying the sample stations for the line of salinity and the three sampling sites for the incubation experiment. The sampling of Kalix river water is described in detail by Herlemann *et al.* (2014).³ The dissolved organic carbon (DOC) concentration on At1, At3 and At4 is shown. (b) Experimental design of the tDOM decomposition culture experiment, (c) the tank setup and the time line for sampling and (d) location sites in detail.

selected station were stored for 110 days at 10 °C. Sampling was carried out at the beginning and after 110 days. The experimental set up for DOM degradation experiments is illustrated in Fig. 1b and c.

2.2. Sample preparation

All water samples taken along the gradient of salinity and for the degradation experiment were treated in the same way. 1 L was immediately filtered through pre-combusted (400 °C, 6 h) Whatman GF/F filters upon collection and the DOM fraction was concentrated by solid phase extraction (SPE) using reverse phase cartridges according to the protocol of Dittmar *et al.* (2008). A recovery rate of the reverse phase extraction for DOM is approximately 65 percent.³⁰ The water was acidified with HCl (37%, Carl ROTH, Karlsruhe) to pH = 2 and pumped through the cartridges at a low flow rate of ~30 mL min⁻¹ by a membrane pump. Then the cartridges were purged twice with 6 mL Milli-Q water (pH = 2) to remove salt residues. The cartridge adsorber material was dried under a stream of N₂,

extracted with 6 mL of methanol (ultra LC-MS, Carl ROTH, Karlsruhe) and rapidly stored at -20 °C after the extraction. All glassware used was acid-washed (0.1 M HCl) and pre-combusted at 350 °C for 6 h.

2.3. Measurement system and sample injection

For the chemical analysis of tDOM, a pyrolysis-gas chromatography/mass spectrometry (Py-GC/MS) system enabling simultaneous application of hard and soft ionization techniques was used (for details see ESI† S1), which had been successfully applied for the investigation of crude oil samples.³⁹

Before sample injection, the methanolic extract was pre-concentrated under N₂ (2 mL to 60 μL approximately), so that 0.5 to 1.0 mg of organic material is present. 10 μL of this extract were injected into the pyrolyzer (Frontier Laboratories, Double-Shot-Pyrolyzer, model: PY-2020iD), which is working with helium as carrier gas (head pressure 1 bar). The samples first undergo a thermal desorption (TD) process at 250 °C for one minute. After the TD process, the same sample is introduced for



a second time after the oven was heated up to 500 °C. For analysis of the elevated chemical compounds, the pyrolyzer is coupled directly to a gas chromatograph (HP 5890 Series II), split 1 : 10, equipped with a non-polar column (SGE-BPX-5; 30 m × 0.25 mm × 0.25 μm). The temperature program was conducted as follows: 60 °C (held for 2 min) to 320 °C (held for 12 min) with 10 °C min⁻¹. After the separation process the gas flow was split (by a deactivated press fit 3-way Y-Splitter for FS tubing with 0.20 to 0.75 mm OD) and transferred simultaneously to two different mass spectrometers. One MS system is a QMS using EI with 70 eV. The other one is a ToFMS using [1 + 1]-REMPI (266 nm). The split ratio was 1 : 100 in favor of the EI-QMS. This is caused by the different column diameters. If the REMPI system gets the same amount of substance as the EI system, the multi channel plate (MCP) would be overloaded and detection of molecular ions would not be possible.

For calibration and optimization of the REMPI-ToFMS system a gas standard (Linde AG, Pullach, Germany) containing 1 ppm of each benzene, 1,2,4-trimethylbenzene, benzaldehyde and toluene in N₂ was applied. For generating REMPI, a Nd:YAG laser (BIG SKY ULTRA, Quantel, Les Uli Cedex, France) with two frequency doubling units yielding 266 nm photons was employed, which corresponds to a photon energy of 4.11 eV. The laser is operated with 10 ns pulse width and a 20 Hz repetition rate exhibiting a power density of approximately 7 × 10⁶ W cm⁻². A reflectron ToF-MS (Reflectron CTF10, Kaesdorf Geräte für Forschung und Industrie, Munich, Germany) is utilized for the detection of the molecular ions. Details about the coupling and more specific device parameters are given in Otto *et al.* (2015) and Fendt *et al.* (2012).^{36,39} The coupling of pyrolyzer and GC/MS with two distinct ionization methods was checked for functionality with an IHSS (International Humic Substances Society) reference sample (Nordic Reservoir NOM). The reference material was dissolved in Milli-Q water and the sample preparation was carried out in the same way as for the water samples. The samples were measured in a mass-to-charge-ratio (*m/z*) range from 10 to 400. On the one hand the total ion chromatograms (TIC) of EI-QMS were consulted to identify compounds with the help of different MS tools (NIST spectral library, mass spectral interpretation). On the other hand, if the data base information was inaccurate, then standard substances were dissolved (approximately 100 mg L⁻¹) and subjected to chromatographic analysis or a tentative assignment was performed, based only on analysis of the fragmentation patterns. For semi quantitative statements the peak intensity of molecular ions of the REMPI-ToFMS measurements was used.

3. Results and discussion

The high complexity of DOM in Baltic Sea water is evident from Fig. 2. It shows an aromatic fingerprint (detection by REMPI-ToFMS) of the sample under TD and pyrolysis conditions, respectively. The *m/z* ratio is plotted against GC retention time. The intensity of the mass signal is depicted as color display. The surface water of the northernmost station (At4) was analyzed. It has a much higher concentration of DOM in comparison to

stations of lower latitudes due to many boreal and subarctic rivers, which drain in the Gulf of Bothnia and influence the water chemistry *via* a high tDOM input, especially in spring time.¹⁵ The TD step shows a less complex spectrum of substances with low signal intensities. Aromatic compounds are present in a mass range between *m/z* 70 and 250. In the pyrolysis step a broad substance spectrum with higher signal intensities is visible. The aromatic substance spectrum starts with an *m/z*-value of approximately 60 going up to 330.

Fig. 2 illustrates that free evaporable molecular structures are barely present. In contrast, a lot of components are accessible only in the pyrolysis step. Hence, DOM mainly consists of large high molecular complex structures. Many of these are aromatic, especially phenolic compounds, hinting at lignin degradation products.³²

With the help of EI-QMS, referring to the fragmentation pattern and database matching, (Wiley, Nist) a classification was successful. Some representatives of possible lignin derivatives are shown in Fig. 3. The GC retention time is displayed against the signal intensity for specific molecular ion tracks. Guaiacol (*m/z* 124, *t*(R) = 468 s), *p*-hydroxyacetophenone (*m/z* 136, *t*(R) = 698 s), 2,6-dimethoxyphenol (*m/z* 154, *t*(R) = 800 s) and vanillin (*m/z* 152, *t*(R) = 152 s) are typical degradation products of lignin. Phenol (*m/z* 94, *t*(R) = 370 s) could be a pyrolysis fragment of *p*-hydroxybenzoic acid, an H-building block of the lignin structure (Fendt *et al.*, 2011; van Heemst *et al.*, 2000).^{32,41} The direct comparison of the two ionization techniques shows that with EI nearly all structures are detectable, but the chromatograms get very complex. Moreover, fragmentation makes peak evaluation even more difficult, which is obvious for the traces *m/z* 152 (vanillin) and 154 (2,6-dimethoxyphenol), respectively, where the relevant peaks are difficult to detect. The simultaneously recorded REMPI-ToFMS spectrum depicts only the molecular ions of aromatic structures. The chromatograms become less complex. REMPI shows clear peak intensities for the aromatic structures vanillin (*m/z* 152) and 2,6-dimethoxyphenol (*m/z* 154). Therefore REMPI data are used for the subsequent statistical analysis (see below), whereas the EI data are still consulted for structural identification.

3.1. Trends with salinity

The changes of DOM with salinity as well as during the incubation experiments are discussed in the following on the basis of selected and frequently observed species representing the H- (phenol, vinyl phenol, *p*-hydroxyacetophenone), V- (guaiacol, 4-vinyl-2-methoxyphenol, vanillin) and S-building blocks (2,6-dimethoxyphenol, 4-vinyl-2,6-dimethoxyphenol, acetosyringone), respectively.^{2,16,21}

For data analysis and to obtain semi-quantitative results, the peak intensity from the REMPI-chromatogram of a molecule with a characteristic retention time is used. That was conducted for all stations from the Kalix river (tDOM source, lowest salinity) to the S1 station (highest salinity). On every station three replicates of the surface water samples were measured. The relative peak intensities were calculated and are illustrated



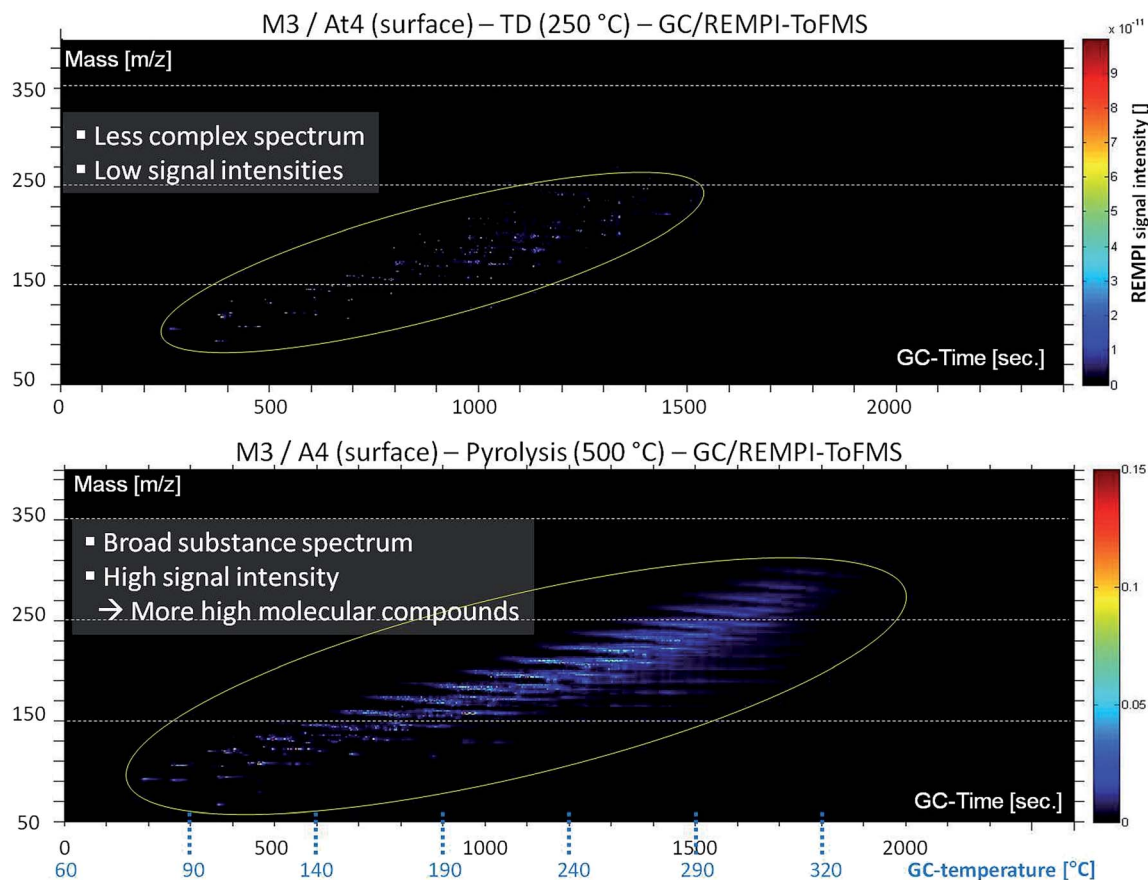


Fig. 2 Image plot (GC retention time vs. mass-to-charge-ratio vs. signal intensity) of At4 surface water (rich in tDOM) under TD (top) and Py (bottom) conditions resulting from the use of REMPI-ToFMS detection.

in Fig. 4–6. For this, the peak intensity of the REMPI chromatogram of a molecular ion was divided by the total intensities for all nominal masses from 90 to 450 m/z in a certain time window. Under the additional assumption that the North Sea input (*via* Skagerrak) constitutes no source for the components of tDOM its influence is not considered.¹⁸

Phenol (m/z 94, $t(R) = 370$ s) is observed to show a decreasing trend from the river mouth to the S1 station (under TD conditions). It is feasible that it was released by biotic or abiotic processes from high molecular weight structures (*e.g.* lignin). To decide about this, incubation experiments were carried out, the results of which are discussed later. Under pyrolysis conditions phenol is also observed, with a slightly falling trend towards the higher salinity. The main origin of pyrolytic phenol is lignin. Similar courses are obtained for other H building block species such as vinylphenol (m/z 120, $t(R) = 610$ s).

In contrast, *p*-hydroxyacetophenone (m/z 136, $t(R) = 698$ s) shows a different pattern. The concentration of the free component clearly increases up to the S6 station and then decreases towards the salt water inflow. However, under pyrolysis conditions, its content remains almost unchanged. This suggests that the free substance (registered under TD conditions) is additionally formed by an alternative mechanism,

which superimposes the general mechanisms forming other H-building blocks such as phenol. However, this only holds down to the latitude of the S6 station. The V-building blocks (Fig. 5), under TD as well as pyrolysis conditions, depict a similar decreasing trend along the Baltic Sea transect as the H-building blocks.

The S building blocks were taken into consideration (Fig. 6), although their input sources into the Baltic Sea should be very low, because there is mainly a freshwater surplus from Scandinavian rivers¹⁵ with typical vegetation of non-flowering vascular plants.^{13,21} Therefore, according to a signal-to-noise ≥ 3 , the representatives 2,6-dimethoxyphenol and 4-vinyl-2,6-dimethoxyphenol could not be analyzed semi-quantitatively under TD conditions. Acetosyringone as the only detectable relevant substance under TD is represented graphically and shows a constant distribution along the transect. Under pyrolysis conditions the concentration of S-building blocks is generally higher. For 2,6-dimethoxyphenol and 4-vinyl-2,6-dimethoxyphenol a slightly rising trend is visible with increasing salinity. A possible explanation is that lignin components are introduced *via* the river systems, having their origin in angiosperm tissues. In general, the relative distribution pattern of the S-building blocks along the line of salinity does not change.



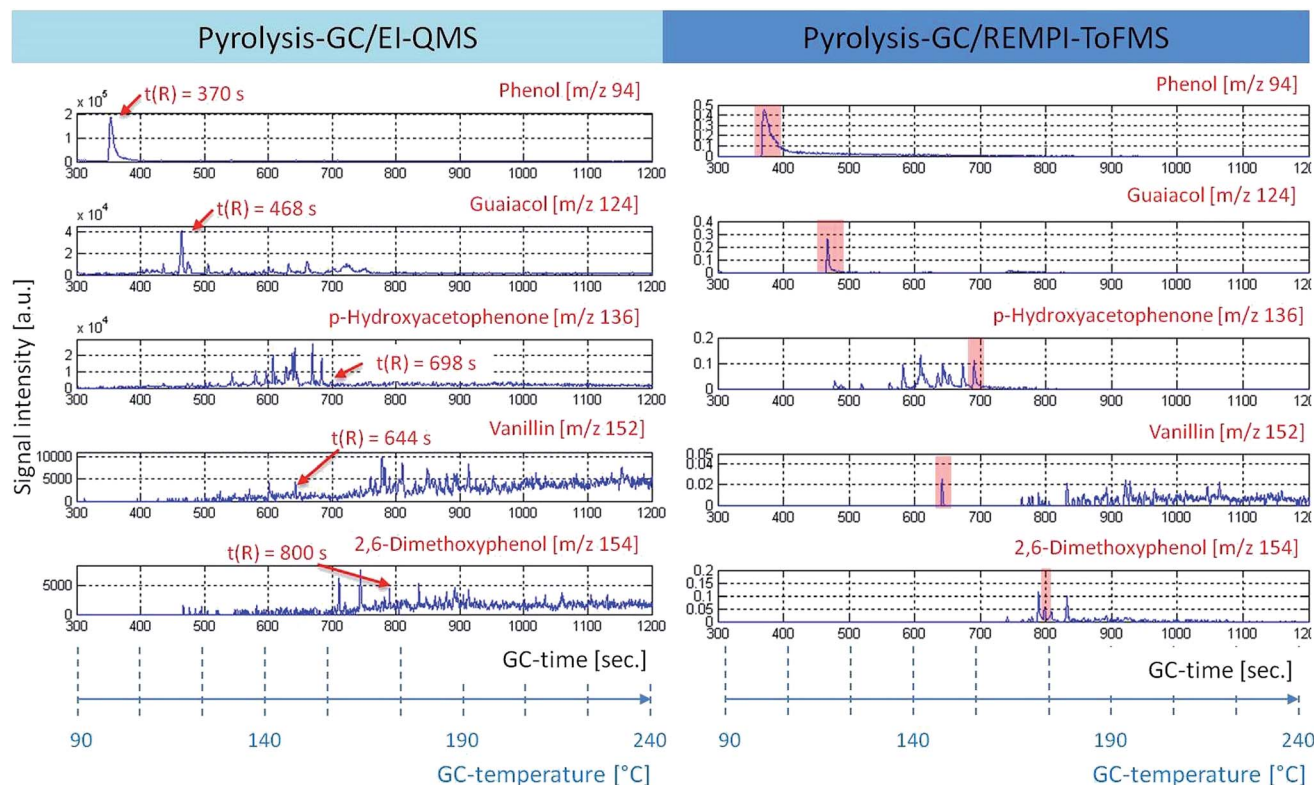


Fig. 3 The time course of ion traces under pyrolysis conditions (retention time or GC temperature vs. signal intensity). It allows a direct composition between the two mass-selective detectors, EI-QMS (left) and REMPI-ToFMS (right). In particular, m/z 94, 124, 136, 152 and 154 can be related to possible lignin degradation products, which can be used as tracers for tDOM.³²

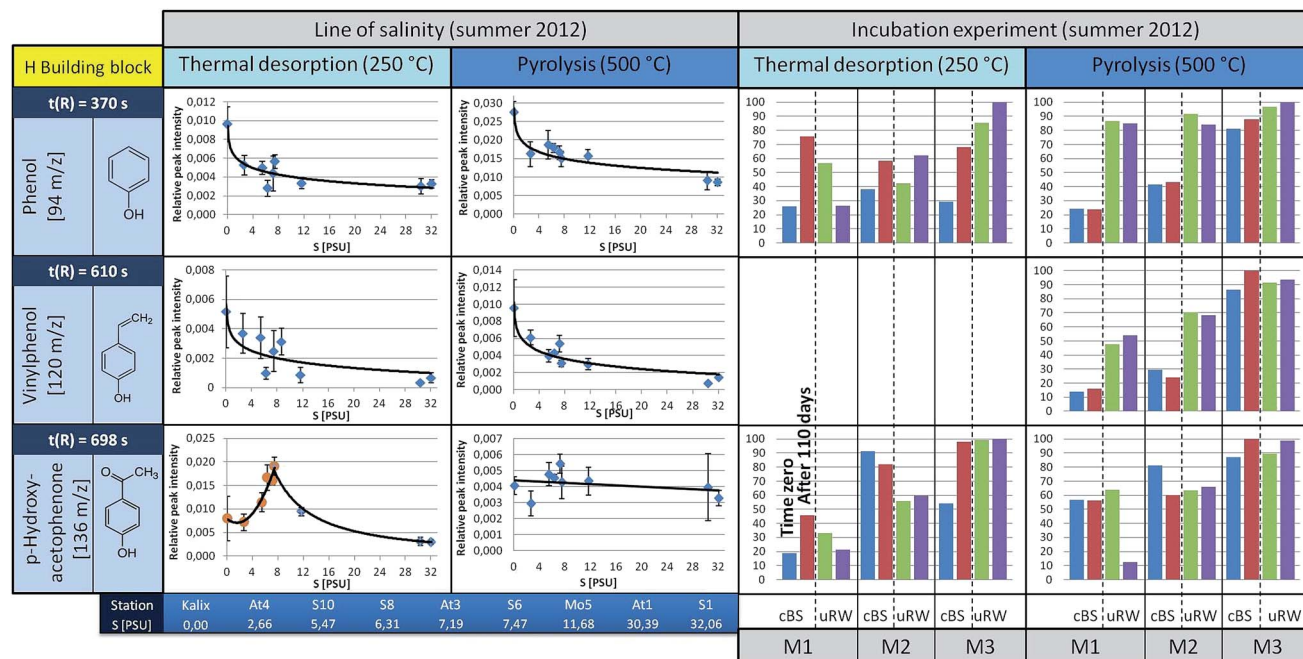


Fig. 4 Summary of three representatives of the H building blocks is shown (left: the curves along the transect of the Baltic Sea, right: the results of the incubation experiments, under TD and pyrolysis conditions). In the diagrams for the line of salinity the relative peak intensities are given for each compound vs. the salinity S . The error bars being the standard derivation of the three measurements. The black lines are the trend curves for the compound concentration. The latter of the respective stations is noted below. For the incubation experiments, the relative peak intensities are shown for a component (specific retention time). The control cBS (pure Baltic Sea water) at time zero (blue) and after 110 days (red) and the mixing tank uRW (Baltic Sea water + river water) at time zero (green) and after 110 days (purple) were compared for every station M1-3. For M1 surface water of At1 station was used, for M2 water of At3 and for M3 water of At4. Only REMPI-ToFMS data were used for the images.



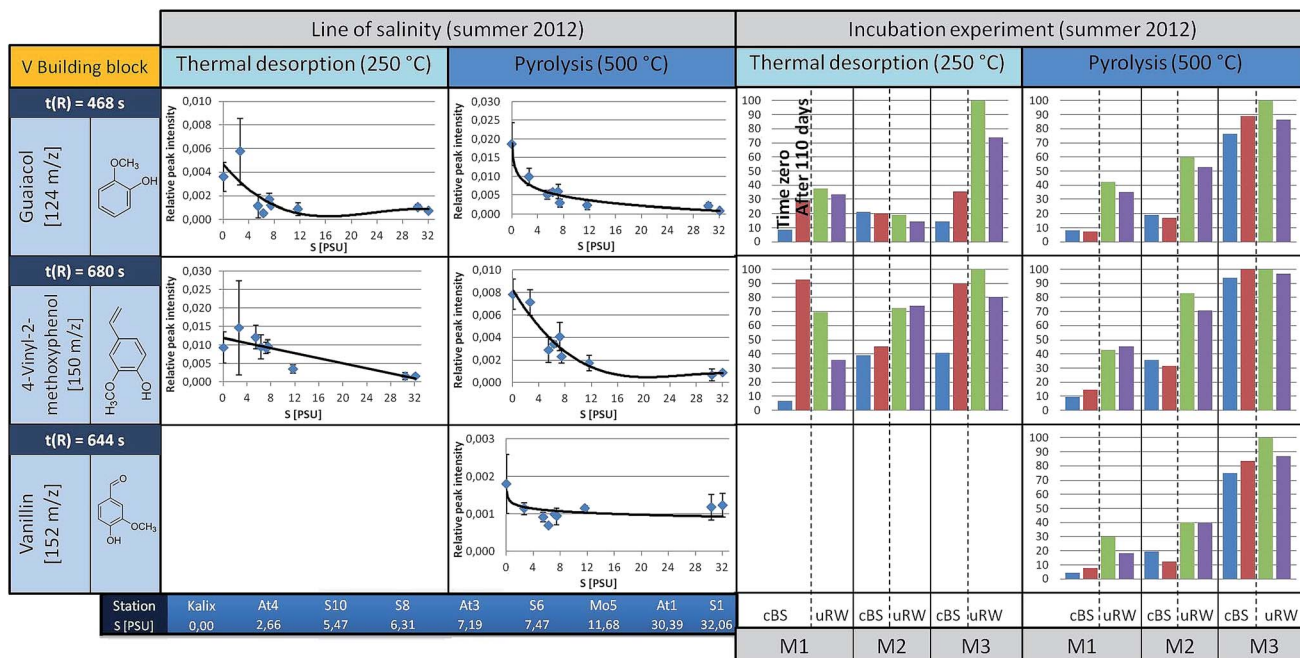


Fig. 5 A summary for three representatives of the V building blocks is given (left: the courses along the line of salinity, right: the results of the incubation experiments, under TD and pyrolysis conditions). For detailed information see Fig. 4. In the free areas (the incubation experiment at 250 °C, component *m/z* 120) no insufficient data for a clean presentation was available.

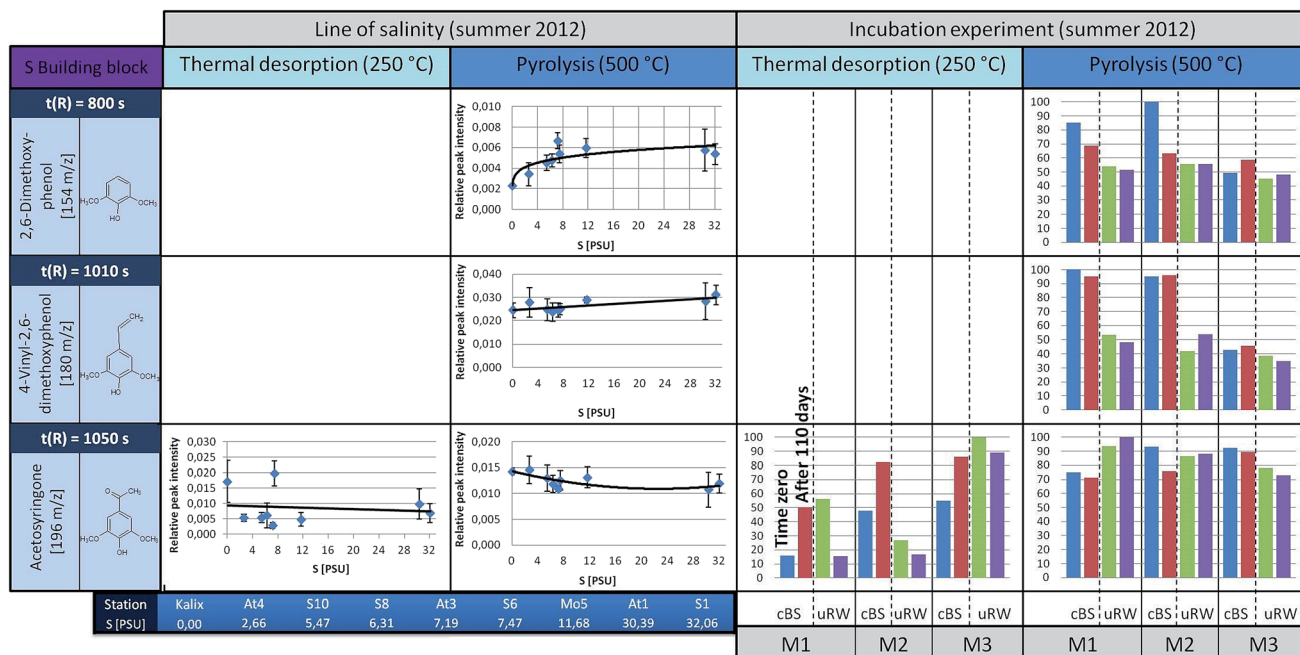


Fig. 6 A summary for four representatives of the S building blocks is illustrated (left: the courses along the line of salinity, right: the results of the incubation experiments, under TD and pyrolysis conditions). For detailed information see Fig. 4 and 5.

3.2. Incubation experiments

But what are the reasons that phenol and vinylphenol (free or macromolecular pyrolysis fragment) tend to decline with higher salt content? In addition to the degradation by sunlight or flocculation,^{9,23} microorganisms are possibly able to affect the

compound concentration along the transect. This hypothesis was studied in the already mentioned tank experiments carried out for three stations, designated as M1, M2 and M3, corresponding to salinities of approximately $S = 32$, 7 , and 3 , respectively.



The results for selected H-building blocks are also given in Fig. 4. The bars in the diagram represent the relative peak intensity of a molecule at the compound-specific retention time. For each station a control tank (cBS) and a mixing tank (uRW) are compared, both for time zero (blue or green bar) and after approximately 110 days of incubation, equivalent to the end of the experiment (red or purple bars).

The direct comparison between cBS- and uRW-tank at time zero shows that the phenolic fraction (under TD) increases significantly due to the addition of river water. This is obvious at all three stations. Comparing the cBS tank at time zero (cBS_{t0}) with the state at the end of the experiment (cBS_{tEnd}) illustrates that the proportion of phenol increases. A similar trend is evident for the mixing tanks uRW (M2 and M3). The differing behavior of the uRW tank cannot be explained with the available data. Nevertheless, the free phenol content shows the tendency to be higher after 110 days. A possible source for this increase could be phenol previously bonded within the lignin structure, which is then released by microbial degradation. In order to prove this hypothesis, pyrolysis measurements were performed. If lignin was indeed degraded during the incubation experiment, the released pyrolytic phenol should decrease significantly after 110 days of incubation time, because it has already been microbially degraded. However, the phenol content between the start and the end time of incubation in all tanks stays nearly the same. This implies that the lignin in the tDOM is not degraded by microbial processes after all. Another source for the increase of the free phenol concentration in the TD step of the sBS and uRW tanks cannot be clearly assigned. For *p*-hydroxyacetophenone, it is difficult to give a reliable statement. In tendency, there hardly occurs a change under TD and pyrolysis conditions in the tank between time zero and 110 days.

The incubation experiment measurements of the V-building blocks under TD conditions show no clear trend. Vanillin was not detected, because its concentration at all stations is below the limit of detection (LOD) for the TD setup. For guaiacol and 4-vinyl-2-methoxyphenol with the exception of the M2 station a significant increase in concentration is observed in the cBS-tank after 110 days. With the M1 and M3 tanks an opposite trend between cBS and uRW samples is observed. For unraveling these contradicting findings, a broader data basis from additional experiments will be needed. In general, the V-building blocks behave similar to the H-building blocks, especially the absence of significant changes in the pyrolytic experiments is striking, corroborating the findings of the H-building blocks.

In the incubation experiments of the S-building blocks samples under TD conditions exhibited too low peak intensities. Under pyrolysis conditions no clear significant changes are observed between the start and end point of sampling for both the cBS and the uRW tank.

The biggest effect on the chemical composition of the tDOM is caused by the added, salinated river water (see ESI,† S2) in the incubation. In order to take a closer look on these effects, in the next section PCA is applied, which can be a helpful tool for data interpretation.

3.3. Principal component analysis (PCA)

Principal component analysis on the basis of the REMPI-ToFMS data was used as a statistical tool in order to detect structural changes along the salt gradient. In addition, PCA is utilized to evaluate the repeatability of technical replicates. The peak intensities of nominal masses between 50 and 350 *m/z* of a GC run were taken. For the three technical replicates of each station and between each station the retention times for the peak have to be matched. For this a self-developed specific software routine (in Matlab) is used that allows an automatic correction of the time shifts between the GC runs. Finally, the performed PCAs are resulting in a matrix of 27 measurements (3 replicates × 9 stations), the nominal masses 50–350 *m/z* and the maxima peak intensities for every ion trace. The data were normalized by the sum peak intensities in the window of the observed mass-to-charge ratio. The reduction to one peak per nominal mass leads to a loss of information, but is necessary for handling and interpreting the huge amount of data.

The results are displayed in Fig. 7. The repeatability of technical replicates is quite recognizable, the three measurements from each station are grouped together. An axis assignment for PC1 and PC2 is not clearly possible. Both reflect the trend of salinity as well as of temperature. These parameters decrease accordingly from the southwest to the northeast of the Baltic Sea. This would explain the apparent diagonal course in the diagram. In the TD step the stations extend in a line from the QII to the QIV sector, according to the salinity or temperature pattern. Apparently, except for the stations S8 and S10, a straight relationship between the molecular structure and abiotic parameters exists. In this context, the separation of the stations is not linear and logically grouped with the salinity gradient. The reasons are unclear and difficult to explain by one parameter. The At4 replicates are located in sector IV, but not in line with the other stations. The replicates of Kalix even significantly differ and have an outstanding position in the plot. Roughly the PCA can be divided in two different areas, the Kalix/At4 and the remaining line of salinity from S1 to S10.

The appropriate loadings plot shows some significant *m/z* values under TD conditions. Those nominal masses are situated in sector QIV, which are present at a lower salinity, for example *m/z* 136 as a possible lignin building block. In addition *m/z* 146, 192, 196 and 206 are significant for the location of At4 in this sector, however, a structural characterization for these *m/z* values was not possible. In sector QII nominal masses are located, which are linked to higher salinities, for example *m/z* 128 (naphthalene) or 166 (fluorene). They represent typical markers for polycyclic aromatic hydrocarbons (PAH) as a possible result of massive ship traffic in the Kattegat/Skagerrak area.⁴² These two *m/z*-ratios and *m/z* 180 (unknown structure) strongly influence the course. The *m/z* values 156, 170, 172, 206 and 213 are detected in the Kalix river, which are responsible for the location of the replicates in QI. A structural characterization is difficult, but a contamination by a plasticizer cannot be excluded.

The PCA under pyrolysis conditions shows a curve from sector 3 up to 1 along the line of decreasing salinity. Under high



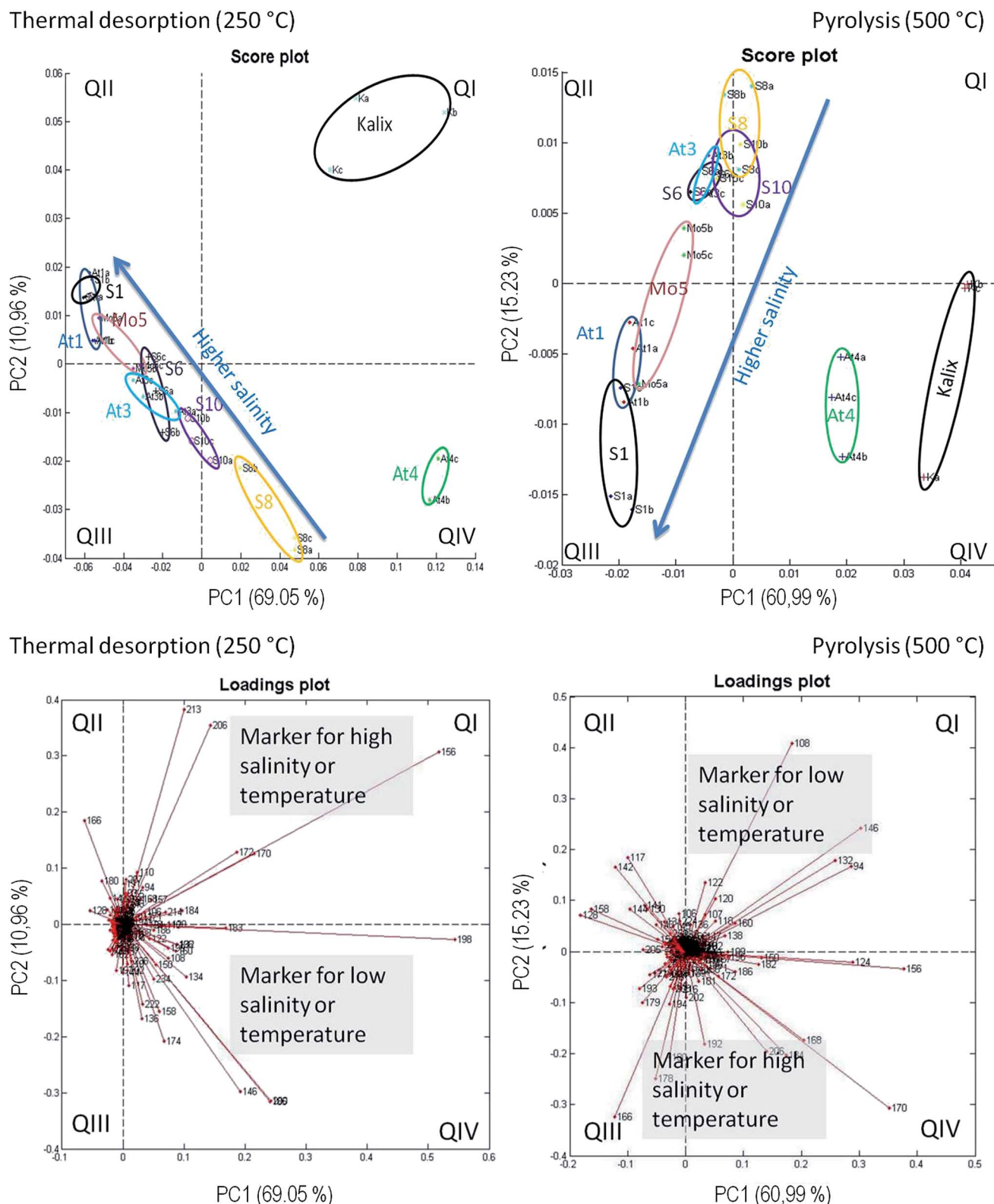


Fig. 7 Top: The score plot of the Principal Component Analysis (PCA) for the line of salinity for three technical replicates per station (grouped together in one cycle) under TD conditions (left) and pyrolysis conditions (right). PC1 and PC2 represent the salinity or the temperature gradient respectively. The trend arrow shows the gradient of salinity. Bottom: Loading plots for the score plots with nominal masses between 50 and 350 m/z .



salinity conditions, again PAH representatives are responsible for the shift. The loadings plot illustrates that m/z 166 (fluorene) or 178 (phenanthrene) are dominant at a high salinity (QIII). In QI (low salinity) mainly the nominal masses m/z 94, 108, 124, 132, 146, 156 and 170 – all of them representing lignin moieties – are responsible for the At4 and Kalix replicates having high positive PC1 scores and that is the reason why these two stations are differing from the others.

As a summary, the results of the PCA are consistent with those from the trend charts before. The abiotic factor salinity has a high influence on the DOM composition. A further important factor for the separation of sampling stations with respect to DOM mixture is the content of lignin derived species and PAH, indicating the influence of terrestrial material and anthropogenic input.

4. Conclusion

The compositional changes of DOM in the Baltic Sea were studied with respect to two different aspects: first, along a Baltic Sea surface water transect and secondly with a focus on the microbial degradation under various special salinity conditions. The chemical analysis of the water samples were carried out by a new coupled system, which has already been successfully tested on crude oil samples.³⁹

Generally along the salinity gradient, referring to the semi-quantitative measurements, a decrease of free and bonded lignin compounds was observed. The PCA underlines that, additionally showing a correlation between salt content and DOM composition. This trend of lignin along salinity is well described in the literature,⁴³ but the used method of TD and pyrolysis in combination with REMPI was applied for the first time for DOM characterization and allows the analysis of aromatic and macromolecular species in a relatively easy way.

Salinity has a crucial influence on DOM concentration and especially on the lignin derived structures. Other compound classes such as PAH are also affected, which is the subject of currently conducted additional analyses. In contrast, microorganisms are hardly breaking the lignin structure. This follows from the semi-quantitative analysis under pyrolysis conditions. Compared to this, under TD conditions changes are quite visible but difficult to interpret. It cannot be ruled out that on different conditions and longer time scales microorganisms would break the lignin structure. Especially several abiotic factors such as wind (continuous mixing, fresh air supply) and UV radiation were excluded in the current setup.

Nonetheless, the coupled system, consisting of a pyrolyzer, a gas chromatograph and two mass selective mass analyzers, has been proven useful to analyze water samples with respect to their DOM content. A sample separation and a combination of universal as well as selective and sensitive detection are advantageous for the characterization of the complex DOM. In particular trace levels of low molecular weight substances (free components or pyrolytic degradation products) can be detected. The analytical method enables a closer look at the molecular composition of natural samples.

The measuring setup could be optimized further by using varying GC columns or a GC \times GC system for a higher separation power.^{44,45} By the use of other relevant wavelengths for REMPI other substances should be accessible. Moving to higher wavelengths, larger aromatic species are ionized more efficiently.³⁷ By an increase of the pulse frequency (femtosecond LASER) short-lived intermediate states of molecules are better stabilized and the sensitivity thus increases.⁴⁶

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