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Dendrochemistry of Multiple Releases of Chlorinated Solvents at a Former Industrial Site

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ABSTRACT: Trees can take up and assimilate contaminants from the soil, subsurface, and groundwater. Contaminants in the transpiration stream can become bound or incorporated into the annual rings formed in trees of the temperate zones. The chemical analysis of precisely dated tree rings, called dendrochemistry, can be used to interpret past plant interactions with contaminants. This investigation demonstrates that dendrochemistry can be used to generate historical scenarios of past contamination of groundwater by chlorinated solvents at a site in Verl, Germany. Increment cores from trees at the Verl site were collected and analyzed by energy-dispersive X-ray fluorescence (EDXRF) line scanning. The EDXRF profiles showed four to six time periods where tree rings had anomalously high concentrations of chlorine (Cl) as an indicator of potential contamination by chlorinated solvents.

INTRODUCTION

Annual growth rings can preserve past environmental changes within the biological requirements of tree survival. The relationship of one ring representing 1 year of tree growth was recognized by Theophrastus in ancient Greece, and Leonardo da Vinci in Renaissance Italy acknowledged year-to-year changes in rings as being related to climatic changes. The assignment of absolute calendar dates for the year of wood formation, the hallmark of dendrochronology, enables the dating of wood samples and artifacts as long as an unbroken record from the current time exists. Chiment and Chiment reviewed over 300 research articles on dendrochemical records derived from elemental isotopes covalently bound to wood constituents; essential “fertilizer elements” associated with ion-exchange sites in wood such as nitrogen (N), potassium (K), and magnesium (Mg); semivolatile or volatile organic compounds, including halogenated volatile organic compounds (SVOCs, VOCs, and HVOCs, respectively). Inferring changes in the external environment from dendrochemical patterns requires an understanding of tree biology, tree physiology, and tree infections. Dendrochemical interpretations are strengthened when combined with traditional analyses of wood chemistry and dendrochronology and can be used effectively to date environmental events.

Dendrochemistry, the chemical analysis and interpretation of precisely dated tree rings, can provide a date-specific record of chemical changes and expands on phytoscreening, in which the chemical analysis of plants is used to monitor and detect plant-chemical exposure. Phytoscreening methods have documented the presence of contaminants or their metabolic products from chlorinated solvents and fossil fuels. When
combined, phytoscreening and dendrochemical investigation can increase the understanding of current and past environmental events, termed phytoforensics. 10 Dendrochemistry can indicate a chemical change in the annual rings, and this information can be combined with groundwater data and modeling, knowledge of contaminant source type and location, and knowledge of tree species physiology to collectively generate scenarios of past events leading to contaminant release and transport and also potential exposure pathways.

Dendrochemistry has successfully utilized several analytical techniques. Examples include energy-dispersive X-ray fluorescence (EDXRF), proton-induced X-ray emission (PIXE), analysis of wood extracts or digests (e.g., inductively coupled plasma-mass spectroscopy, or ICPMS), and atomic emission spectroscopy (AES). EDXRF methods tend to be favored largely because of the capacity for simultaneous analysis of 31 target or marker elements and compounds at a high spatial resolution of 50 μm across the tree-ring sample. 6,12 Regardless of analytical method, it is important that the dendrochemist determine which observations are related to processes involved with tree physiology and species relative to the interaction with pollutants. 9,12,13

In this study, we describe the dendrochemical investigation methods and results in a groundwater pollution case, where the tree chemistry appears to reflect four to five asynchronous releases of HVOCs, along with other environmental influences and physiological anomalies. For this study, four trees were used that were located along a plume (see Figures 1-5). These trees (see Table 1) belong to three genera (two oaks, one sycamore, and one fir).

<table>
<thead>
<tr>
<th>Tree Number</th>
<th>Tree Species</th>
<th>Oldest Ring</th>
<th>Years of Unique Cl Anomalies</th>
</tr>
</thead>
<tbody>
<tr>
<td>5</td>
<td>Quercus sp.</td>
<td>1978</td>
<td>1989, 2000</td>
</tr>
</tbody>
</table>

**EXPERIMENTAL SECTION**

**Study Area.** Verl is a relatively small city in Germany with an area of 73 km² and a population of about 25000. The aquifer at Verl consists of heterogeneous glacial deposits, resulting in uncertainties in determining the aquifer characteristics. The aquifer contains an elongated plume of groundwater contaminated by chlorinated aliphatic solvents (Figure 1). An earlier investigation using phytoscreening methods demonstrated that the trees take up groundwater contamination and could be used to map the plume. 10,14 The phytoscreening analysis also provided evidence of multiple TCE sources, subsequently supported by the discovery that fill material at the site was also contaminated by HVOCs. 14

**Sample Preparation.** Increment cores (10 mm in diameter) were extracted from nine trees (Figures 1 and 2) on February 22, 2010. After 5 days of drying in ambient air between grooved wooden blocks, cores were surfaced with a graded series of silicon carbide sandpapers (50, 100, 200, and 400 grit). The sanded cores were observed under a digital microscope (40–100× magnification) to evaluate the presence of branch anomalies, infection, and tree response to injury and infection that could have affected the dendrochemical pattern.

**Figure 1.** Location of study area, groundwater sampling, and trees sampled for dendrochemistry in 2010. Groundwater concentrations were measured in 2010 by the consulting firm HPC Envirotec. The plume map used in figure is for the total of PCE, TCE and dDCE.

**Figure 2.** Sampling of tree 2 using a 10-mm Pressler borer.
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Stockholm University, using an Itrax Multiscanner (www.cosysys.se). All sample processing and EDXRF analyses were conducted under conditions of constant air temperature (20°C) and relative humidity (50%). The sample cores were first cut along their full length parallel to the transverse wood surface with a twin-blade saw to produce 2-mm-thick laths with planar surfaces. The laths were then mounted vertically in the Itrax Multiscanner to allow for radiographic density and EDXRF analyses along the length of the planar transverse surface of the samples.

**EDXRF Analysis.** The Itrax Multiscanner is a “flat-beam” X-ray scanner where samples are exposed to a precisely defined, thin X-ray beam with a vertical dimension of 50 μm. An X-ray line camera records the transmitted radiation that passes through the sample. Hence, a full-scale two-dimensional radiographic image with 50-μm resolution is obtained by moving the sample relative to the beam in 50-μm steps. At each step, a silicon drift chamber detector (SDD) records the characteristic fluorescent X-ray radiation (XRF) from specific alkaline earth and other metal elements with the atomic weight of Mg and higher. The collimator system within the SDD confines XRF recording to a sample area of 0.05 × 2 mm². The SDD functions as an energy-dispersive X-ray detector, with a capacity of 100,000 counts per second (cps) and a resolution of about 135 eV at 5.9 keV. Equipped with the standard 1.9-kW chromium-tube X-ray source, the SDD records about 20,000 cps for wood tissue. An exposure time of 10 s is normally sufficient to record element concentrations above 10 ppm.

With the resolution of 50 μm, a tree ring with a width of 1 cm provides 200 XRF data points for elemental concentrations. High-resolution profiles of elemental concentrations were established over a large number of rings within a few hours (about 3.3 min/mm) and in a nondestructive way. The raw EDXRF data, made available as an Excel file, provide relative concentrations in counts for 31 elements along with physical parameters (locations, counts, mean squared errors). The raw data were computerized using an xls template, called Scanchem, that automatically generates density-corrected concentrations and time-weighted centered moving averages (TWCMAs). For the Verl site, Cl associated with chlorinated solvent degradation was used as a monoelemental marker that was detectable by XRF. HVOCs are naturally degraded by bacteria within groundwater and soils in reducing environments in reductive dechlorination or “dehalorespiration” (i.e., PCE to TCE to dDCE) and can generate large plumes of Cl ions that can be taken up by root systems of trees. The parent HVC can also be taken up by trees. HVOCs within trees can spread through the porous wood and can partly evaporate through the stem and leaf crown. HVOCs can also degrade in plant tissues if reducing conditions dominate (i.e., in a decaying tree) or through aerobic plant defense pathways to form trichloroacetic acid and other metabolites that can be further degraded.

At relatively low concentrations, Cl is usually stable in woody tissues after fixation, presumably through ionic binding, and is not readily remobilized/retranslocated. The potential for HVC movement across ring boundaries and for HVC dechlorination within trees leads to the possibility that Cl generated from in-tree degradation might be fixed in growth rings not corresponding to the year of HVC uptake. In situations where there is no substantial dechlorination activity in the groundwater and soils, the Cl signature in tree cores is unlikely to provide information on the timing of HVC uptake. When there is substantial underground dechlorination within the capture zone of the root system, however, it is likely that the groundwater Cl signature would overwhelm in-tree

effects and would predominantly be located in the growth ring corresponding to the year of uptake. In the tree cores examined at Verl, there was no distinctive change in Cl chemistry from one side of the heartwood/sapwood boundary to the other, implying that there was no significant translocation across the boundary, as is sometimes seen for translocated elements.25

RESULTS AND DISCUSSION

The EDXRF procedure allows for the direct comparison of two or more elements at once (Figures 3 and 4). Bielemental profiles for Ca (in white) and Cl (in green) are superimposed on an X-ray image of the core from tree 7 (Figure 3). The direction of growth is from left to right, with the transition between annual growth rings marked by the contrast between the more physically dense late wood (light-colored in the X-ray image) and the less dense early wood of the following year (dark-colored in the X-ray image). Note the characteristic annual, intra-ring patterns, often showing the lowest Cl concentrations in the early wood.

![Figure 3](image-url)

**Figure 3.** Relative concentrations of chlorine (green) and calcium (white) determined by energy-dispersive X-ray fluorescence overlaid on an X-ray image of the tree 7 core (Verl, Germany). Also, note that, at the 1990/1991 rings, the Cl impact occurred during the period of late wood formation, as evidenced by the peak having its highest amplitude in the 1990 late wood, and the subsequent early wood recorded a Cl minima significantly higher than in previous rings. Note the characteristic annual, intra-ring patterns, often showing the lowest Cl concentrations in the early wood.

The Cl-only anomalies are interpreted as being associated with the chlorinated solvent plume. Also, in the 1990/1991 rings of tree 7, the Cl uptake appears to have occurred during the late wood growth period. Consistent with this interpretation, the minimum Cl concentration in the subsequent early wood growth period was substantially higher than in previous rings (Figure 3).

EDXRF shows distinct bielemental anomalies for Ca and Cl in tree 2 (Figure 4), peaking in 1908, 1918, 1931, and 1976. The dual Ca and Cl signature was interpreted as possibly being the result of drought or other climatic influences not related to contamination; the 1976 drought had a similar Ca/Cl signature in tree 2 as the three earlier anomalies. Tree 2 also shows four to five asynchronous Cl-only impacts (Table 1), potentially indicating the presence of HVOCs.26 The 1948/1949 Cl peak in tree 2 does not have a directly corresponding Ca peak, but there are other Ca anomalies in close proximity (Figure 4), imparting some level of uncertainty as to whether this was a Cl-only event. It is of interest, however, that, while searching for a potential cause of the 1948/1949 Cl anomaly in tree 2, investigators found that the industrial plant had been active since 1945 and that two buildings containing PCE were destroyed in 1948 and on April 22, 1949, potentially resulting in a contamination release.

It is highly likely that the chemistry in tree 2 is influenced by groundwater chemistry because the depth to groundwater at tree 2 is only about 50 cm. Cl anomalies in tree 2 are present in growth rings from two years that the PCE-containing buildings were destroyed and might represent HVOCs related to those events. If so, then the HVOCs would have been rapidly transported from the source area to tree 2. A potential explanation for rapid transport of HVOCs is that they could have moved by overland flow through a drainage ditch or creek. There is, in fact, the remnant of a former ditch or stream near trees 1 and 2, over 15 m in length and oriented in the direction of groundwater flow (the ditch is shown in the background of Figure 2). Construction and landscaping have eliminated much of the stream and obscured its original path; however, proximity to trees 1 and 2 and orientation in the direction toward the source area imply the possibility that the ditch once provided surface drainage from the source area to the vicinity of tree 2. The subsequent Cl anomaly in tree 2, 11–12 years later (1960/1961), could potentially be explained by groundwater transport of contamination and delayed impact from the same 1948 and 1949 releases. Modeling simulations (MODFLOW) produced time-of-travel estimates consistent with groundwater transport of contaminants to the four trees at dates corresponding to the Cl anomalies (see Figure 5).

Trees 5–7 also show, depending on their age and location, a minimum of two to four asynchronous Cl anomalies (Figures 5 and 6). Potentially, the anomalies might represent the transport

![Figure 4](image-url)

**Figure 4.** Detail of bielemental profiles for Ca in white and Cl in green, superimposed on an X-ray image of the tree 2 core. The direction of growth is from left to right, expressed by the transition from dark early wood (with low density) to light late wood (with high density). The chlorine-only anomalies are interpreted as a potential response to a chlorinated solvent plume.
of additional releases of contaminants. Trees 5 and 6 are at the edge of the present groundwater plume boundary (Figure 1); however, the historical extent of contamination might have been different than what is now present. Knowledge of groundwater and potential contamination sources and events is needed to provide corroborating lines of evidence.

Measured groundwater flow rates and model (MODFLOW) estimated plume migration rates (consulting firm HPC Envirotec, written communication, 2010) showed the probable plume migration rate to be about 20–30 m/year in the southern part and 2–3 times higher in the northern part of the plume. The plume migration rate, as estimated from a comparison of Cl anomalies in trees to the distance separating the trees, was about 30 m/year in the southern part of the plume and 70–120 m/year toward the northern half of the plume. The average plume migration rate based on the tree cores was about 54 m/year over the 820 m separating the source from the most distant tree, tree 5 (see Figure 6). The close agreement between the estimates of the plume migration rate supports the hypothesis that the Cl anomalies represent movement of groundwater contamination. Agreement between model-derived and tree-chemistry-derived groundwater flow rates was also observed in a previous study.26

It should be noted that there are uncertainties in both the groundwater- and tree-derived estimates of contaminant transport. The uncertainties in the groundwater-derived estimates involve uncertainties in both modeling formulations and porosity estimates, particularly in these heterogeneous glacial deposits. Uncertainties in the tree-derived estimates involve a possible half-year lag time during the nongrowing season when a dendrochemical signal might not be effectively recorded. If, however, dendrochemical signals can be shown to be a reasonable estimate of contaminant arrival dates for a particular site, then the information could have great supportive value as a complementary approach for estimating contaminant transport rates.

The black, red, blue, and yellow lines in Figures 5 and 6 identify the potential asynchronous releases of Cl, labeled A–E, respectively, and correspond to the suspected releases listed in Table 2. Based on the plume migration rates through groundwater estimated from the dendrochemical record and the distance between the southernmost trees and those closest to the source and historical records, the timing of the releases were estimated to be 1948–1949 (release A), 1975 (release B), 1985 (release C), 1990 (release D), and 1999 (release E), as reported in Table 2. Releases that took place after industry ceased its production activities in 1969 might be associated with

Figure 5. EDXRF data for the four analyzed cores from Verl, Germany, showing elemental profiles for Cl. They correspond, from top to bottom, to trees 7, 6, 2, and 5 and are listed in order of their distance from the plume. The red, blue, and yellow lines correspond to interpreted events B, C, and D, respectively, in Table 2. Note that tree 2 is the only tree old enough to have reflected potential releases identified from 1948/1949 and the subsequent 1960/1961 impact via groundwater; tree 6 is too recent to show the impact found in tree 7 in 1980 and later found in trees 2 and 5 (red arrow). The recent 2004 Cl anomaly found in tree 7 would not have impacted tree 6 in 2010. The orange arrows represent the location of the heartwood/sapwood boundary (HSB). The lack of a distinctive change in Cl concentrations across the HSB provides evidence that there is no significant chlorine lateral translocation in these trees.
progressive plant dismantling and remediation activities or with activities at a garage shop that began operation in 1980.

The data provided herein demonstrate that the high resolution of the EDXRF method allows comparisons of wood chemistry at a scale that can differentiate changes within a single growth ring. The data can be used to examine the timing of environmental events with approximately annual precision. Although the past can never be reconstructed with complete accuracy, the Verl study area shows the potential of dendrochemical methods to indicate multiple historical and asynchronous environmental impacts to trees, along with corresponding plume migration rates, and to age-date releases from the source. The combination of the dendrochemical data obtained using today’s methodologies, available information on historical activities at the industrial locations, and groundwater modeling data potentially can be used to project back possible scenarios of events that occurred in the past.

Table 2. Estimated Times of Pollutant Releases at the Site in Verl, Germany

<table>
<thead>
<tr>
<th>plume</th>
<th>time of contaminant release</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>1948, 1949</td>
</tr>
<tr>
<td>B</td>
<td>1975</td>
</tr>
<tr>
<td>C</td>
<td>1985</td>
</tr>
<tr>
<td>D</td>
<td>1990</td>
</tr>
<tr>
<td>E</td>
<td>1999</td>
</tr>
</tbody>
</table>

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