Ultraviolet degradation of methyltins: Elucidating the mechanism by identification of a detected new intermediary product and investigating the kinetics at various environmental conditions

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Abstract

The photodegradation of methyltins, as environmental pollutants, has scarcely been studied so far because of the shortage of rapid and sensitive speciation methods, even though they have very simple structures. The photodegradation of monomethyltin trichloride (MMT), dimethyltin dichloride (DMT) and trimethyltin chloride (TMT) was studied with our new developed HPLC-FPD hyphenated system, which enables rapid and sensitive detection of methyltins. The half-life times and kinetic rate constants of their degradation at different pH were calculated. The results suggest that MMT, DMT and TMT can be degraded under the UV irradiation rapidly at different pH, with a degradation rate sequence of TMT < DMT < MMT. An unknown intermediary product, which is more stable and has higher concentration at pH 8 for MMT and DMT, of methyltin photodegradation was detected for the first time. This unknown intermediary product was identified as methyloltin with electrospray mass spectrometry, and the possible mechanism was proposed based on the intermediary product. The effects of some environmental parameters such as salinity and humic acid on the degradation rate of methyltins were also investigated. Results suggest that salinity and humic acid have strong effect on their degradation, especially for TMT, which was almost never degraded in the solutions containing NaCl and humic acid.

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1. Introduction

Organotin compounds are omnipresent in the environment (Burton et al., 2006; Rüdel et al., 2007; Gómez-Ariza et al., 2006) because of their extensive applications as biocides, antifouling agents and polyvinyl chloride heat stabilizers over the past decades. Although their uses have been banned by many countries (Fent, 1996), some organotin compounds such as methyltin compounds continue to be produced and used as polymer stabilizers nowadays. On the other hand, unlike the other organotin compounds that might disappear following the prohibited use, the methyltin compounds will exist in the environment for a long time because of biotic and abiotic tin methylation (Ashby and Craig, 1988). Generally, the shorter chain of alkyl groups, the higher toxicity of the organotins. For methyltins, the toxicity follows the sequence: trimethyltin (>tetramethyltin) > dimethyltin > methyltin (Pettibone and Cooney, 1988). For example, methyltins, especially trimethyltin, demonstrate more toxic to fungi and insects than other organotin compounds (Ebdon et al., 1998).

Due to the high toxicity of organotin compounds, their fate in the environment (biotic and abiotic degradation) is an important issue in the study of their environmental behavior (Burton et al., 2005; Lee et al., 2006). The disappearance of organotins from the environment is mainly
attributed to their biodegradation, photolysis and chemical degradation. They can be degraded through stepwise dealkylation to form inorganic tin under various environmental conditions. The biodegradation of organotin compounds, such as degradation by bacteria (Harinoa et al., 1997; Inoue et al., 2000), has been studied in detail. The abiotic degradation mainly includes the gamma irradiation (Allen et al., 1987) and photolysis (Maguire et al., 1983; Blunden, 1983; Mailhot et al., 1999). Some investigations have been conducted to study the process of photodegradation of organotin compounds because photolysis is an important process for organotin compounds in the environment. But these studies mainly focus on the degradation of butyltin and phenyltin compounds rather than methyltins. Mailhot et al. (1999) investigated the iron (III) photoinduced degradation of tributyltin chloride (TBT). They proved that the degradation of TBT is due to the attack of OH radicals, and the main route to degradation is a stepwise debutylation of TBT to di- and mono-butyltin with final formation of inorganic tin. In order to accelerate the degradation rate in aqueous solution, TiO2 was used for photocatalytic degradation of butyltin compounds under UV illumination (Navio et al., 1996). The photodegradation of triphenyltin in water was also studied (Soderquist and Crosby, 1978, 1980).

The degradation kinetics and mechanism of methyltin compounds, including photodegradation, was scarcely studied. Blunden (1983) studied the ultraviolet degradation of methyltin chlorides in carbon tetrachloride and water using NMR technique, but the adopted concentration of these methyltin compounds was 0.05 M (as Sn), which is 5935 mg l⁻¹ (as Sn) and far higher than their real concentration in environment. One of the main reasons for the scarce research on the degradation of methyltins is the shortage of a rapid and sensitive method for the speciation of methyltin compounds. Gas chromatography (GC) and high performance liquid chromatography (HPLC) based hyphenated techniques, such as GC-AAS (atomic absorption spectrometry) (Ritsema, 1992) and GC-ICP-MS (inductively-coupled plasma mass spectrometry) (Tao et al., 1999) are the general methods to determine methyltin compounds. The complicated pretreatment step (Pellegrino et al., 2000) and derivatization step (Morabito and Massanisso, 2000) in GC methods limited their applications in the study of methyltin degradation because a large number of samples during degradation process have to be analyzed rapidly in 30 min. For the HPLC methods, the difficulties originate mainly from the scarce of sensitive detector to measure methyltin compounds. Though ICP-MS (Chao and Jiang, 1998) is a sensitive detector for methyltin compounds, it is difficult to couple it with reversed phase-HPLC for the speciation of methyltins because of its incompatibility with the higher content of organic solvents (methanol or acetonitrile) that have to be used to separate methyltins. High levels of organic solvents in the mobile phase of reversed phase liquid chromatography causes the extinction of ICP which restricts the applications of ICP-MS hyphenated with reversed phase HPLC.

Recently, we have developed a HPLC-HG-MQLS-QSIL-FPD (HPLC-hydride generation-microporous membrane gas–liquid separator-quartz surface-induced luminescence flame photometric detector) system for convenient speciation of methyltin compounds with high sensitivity and efficiency (Zhai et al., 2007).

In this work, the degradation kinetics and mechanism of methyltins under the UV irradiation was studied with the new developed HPLC-HG-MQLS-QSIL-FPD method, and the newly detected intermediary product was confirmed using ESI-Q-TOF-MS. The influenced mechanisms of some environmentally relevant parameters such as pH, salinity and humic acid on the degradation of methyltins were studied in detail.

2. Materials and methods

2.1. Chemicals and reagents

All reagents were used as purchased without further purification. Three methyltin compounds, including trimethyltin chloride (TMT, 98%), dimethyltin dichloride (DMT, 97%) and monomethyltin trichloride (MMT, 97%), were obtained from Aldrich Chemical Co. (USA). Standard stock solutions were prepared by dissolving methyltin compounds in deionized water and stored at 4 °C. The working solutions were prepared daily by appropriate diluting the stock solutions with water and adjust to pH 3.5 using hydrochloric acid. Methanol was HPLC grade solvent (J.T. Baker, Phillipsburg, NJ, USA). Tropolone (98%) and humic acid (sodium salt, tech. 50–60% as humic acid) were purchased from Acros Organics (NJ, USA). The 600 mg l⁻¹ humic acid as stock solution in the experiments was filtered using 0.45 μm membrane. Acetic acid (AR, 99.5%), hydrochloric acid (AR, 36–38%), sodium chloride, (AR, 99.5%), potassium hydroxide (GR) and anhydrous calcium chloride were obtained from Beijing Chemical Factory (Beijing, China). Potassium borohydride (>95%) was purchased from Institute of Geophysical and Geochemical Exploration (Langfang, China). The deionized water (18.3 MΩ) was from an ultrapure water system (Barnstead International, Dubuque, IA, USA).

2.2. Apparatus

The hyphenated HPLC-HG-MQLS-QSIL-PFD system was the same as described previously (Zhai et al., 2007). In brevity, methyltin compounds were first separated by the HPLC system which consists of a DGU-12A degasser, a LC-10A pump (Shimzdu, Japan), an injector valve (Rheodyne 7725i with a 200 μl sample loop), and a C₁₈ separation column (Agilent Zorbax Eclipse XDB-C₁₈, 5 μm particle size, 4.6 × 150 mm). The analytes from the HPLC column reacted with the derivatization reagents to form volatile methyltin hydride. These volatile methyltin hydride compounds were separated from the liquid matrix with a
microporous membrane gas–liquid separator (MGLS). When the volatile methyltin hydride compounds pass through the gas–liquid mixture channel of the MGLS, they diffused through the micropores of the PTFE membrane into the gas channel and were transported to the dryer by the high-purity nitrogen from GC injector used as carrier gas. After dried by the anhydrous calcium chloride in the dryer, the methyltin hydride compounds were introduced into the QSIL-FPD for detection. The QSIL-FPD was installed on a gas chromatography (BeiFen-3420, China) with BF-2002 chromatographic workstation software. The operating conditions of this hyphenated system are displayed in Table S1 (see supplementary material).

Macromass Electrospray ionization-Quadrupole-Time of flight-mass spectrometry (ESI-Q-TOF-MS) with Masslynx 4.1 workstation software was from Waters Company. The major voltage conditions are listed as follows: negative ionisation mode (ESI−), 2500.0 V capillary, 30.0 V sample cone, 2.0 V extraction cone, 1.0 V ion energy, 5.0 V collision energy, 5630.0 V TOF flight tube and 1780.0 V reflectron. The high-pure nitrogen is as the carrier gas with flow rate of cone of 50 l h\(^{-1}\) and desolvation of 350 l h\(^{-1}\). The temperatures of desolvation and source are 150.0 and 80.0 °C, respectively. The syringe rate is 30.0 µl min\(^{-1}\) and the scan mass range is from \(m/z\) 100 to 500.

The ultraviolet light (UV) source for photodegradation is a 40 W low-pressure mercury vapor lamp with the wave length of emission of 253.7 nm and lamp length of 38 cm.

2.3. Procedure

Into a 250 ml beaker was added 200 ml of 500 ng ml\(^{-1}\) (as Sn) MMT, DMT or TMT solution, and the UV lamp was adjusted to 6 cm over the liquid surface of beaker. Typically, four samples can be simultaneously processed. After irradiation for the prescribed time with the UV lamp, 200 µl of the sample solution was taken from the beaker and directly injected into HPLC-HG-MGLS-QSIL-PFD system, and the methyltin compounds can be completely separated and determined within 20 min. To investigate the effect of pH, methyltin solutions was adjusted to pH 4, 6 and 8, using hydrochloric acid and potassium hydroxide. Methyltin solutions with 0, 0.5, 1.5 and 3.0% (m/v) of sodium chloride and pH 6 were used to study the effect of salinity, and solutions with 0, 3, 6 and 9 mg l\(^{-1}\) of humic

![Fig. 1](image_url). Effect of pH on the photodegradation rates of methyltins. (A) MMT, (B) MMT containing 0.05% methanol (v/v), (C) DMT, and (D) TMT.
acid and pH 6 were used to evaluate the effect of humic acid.

3. Results and discussion

3.1. Degradation kinetics at different pH

The methyltin compounds can be degraded under the irradiation of ultraviolet light. As can be seen from Fig. 1, the degradation rates of MMT, DMT and TMT is strongly dependent on the solution pH. All the degradation curves of methyltins are in accord with the pseudo-first-order rate equation except for the MMT at pH 4 and 6, which are in accord with zero-order rate equation (see Table S2 in supplementary material). For all the studied methyltins the highest degradation rate was obtained at pH 6, while the lowest one was obtained at pH 8. Two major reasons might be responsible for this phenomenon. First, the species of the three methyltin compounds in water are dependent on the pH, which influences the energy needed to break the Sn–C bond. Various species of methyltin cations are formed over a wide pH range (Stefano et al., 1999). For example, $(\text{CH}_3)_2\text{Sn}^{2+}$ forms five species including $(\text{CH}_3)_2\text{Sn(OH)}^+$, $(\text{CH}_3)_2\text{Sn(OH)}_2^0$, $(\text{CH}_3)_2\text{Sn(OH)}_3^0$, $[(\text{CH}_3)_2\text{Sn}]_2^{2+}$ and $[(\text{CH}_3)_2\text{Sn}]_2^{2+}/[\text{OH}]^2$, while $(\text{CH}_3)_3\text{Sn}^+$ forms only $(\text{CH}_3)_3\text{Sn(OH)}^0$ and $(\text{CH}_3)_3\text{Sn(OH)}_2^0$. Monomethyltin(IV) cation undergoes hydrolysis and forms $\text{CH}_3\text{Sn(OH)}_2^0$, $\text{CH}_3\text{Sn(OH)}_3^0$ and $\text{CH}_3\text{Sn(OH)}_4^0$ species in the pH range 2–10.5. Moreover, the species would be hydrated to form more stable structures with a coordination number of 5 or 6. Second, the UV degradation of organic compounds is the free radical reaction, which is influenced by the various anions in aqueous solutions. For example, the degradation rate of 4-chlorophenol (Lipczynska-Kochany et al., 1995) in aqueous solutions was found to decrease with the following sequence: $\text{ClO}_4^->\text{NO}_3^->\text{Cl}^->\text{HPO}_4^{2-}>\text{HCO}_3^-$, since these anion can scavenge the hydroxyl radical in the reactions. As an anion, $\cdot\text{OH}$ can also function as a free radical scavenger in aqueous solutions, it can inhibit and reduce the formation of hydroxyl radical thus decrease the degradation rate of methyltins at pH 8. Since HCl was used to adjust the solution pH, the aqueous solutions at
pH 4 possessed higher concentration of Cl⁻ than that at pH 6 and thus lower degradation rate. Similar results were obtained in the degradation of trichloroethylene (TCE) with persulfate oxidation (Liang et al., 2007). The maximum degradation rate of TCE occurred at pH 7, while the degradation rate decreased with either an increase or a decrease of pH within pH 4–9 as SO₄²⁻ may play a role in scavenging hydroxyl radical and thus slowing down the TCE degradation rate.

As shown in Fig. 1 and Table S2, the degradation rate of methyltins has a sequence of TMT < DMT < MMT because of the reverse sequence of methyl groups which hinder the reaction. However, Blunden (1983) observed a degradation rate sequence of TMT > DMT > MMT, which probably is due to the adopted concentration of methyltins and the energy of UV lamp are different from this study. Another reason is that the NMR technique used by Blunden cannot distinguish the signals of the MMT and the intermediate product, which have very similar structure. Our study indicates that the degradation route of methyltins might be a stepwise demethylation to inorganic tin, as the detected degradation products were DMT, MMT and an unknown intermediate product for TMT; MMT and the unknown intermediate product for DMT; as well as only the unknown intermediate product for MMT, respectively. However, Blunden (1983) did not detect MMT during the TMT degradation process, which might be attributed to the low sensitivity of NMR spectroscopy as very low concentration of MMT was detected during the degradation process of DMT and TMT at pH 6 in this study. High concentration of the intermediate product was formed for MMT and DMT at pH 8, which means that the intermediate product can be easily formed at pH 8; whereas DMT degraded to MMT, then form the intermediate product and finally tin. For TMT, however, the degradation mechanism might be different from that of MMT and TMT as the intermediate product was not detected even at pH 8. The possible degradation mechanisms of methyltins and the confirmation of the intermediate product will be discussed later.

To prove the degradation to take place through the radical reaction, methanol (0.05%, v/v), a well-known hydroxyl radical scavenger (Marugan et al., 2006), was added to the MMT aqueous solution. It can be seen from Fig. 1B that methanol greatly inhibits the degradation of MMT at all the studied pH values. The half-life time at pH 4, 6 and 8 increased from 181, 93 and 210 min to 305, 815 and 1645 min, respectively (Table S2, supplementary material). Therefore, it can be concluded that the degradation of methyltins under the UV irradiation is through the radical reaction.

3.2. Confirmation of the unknown intermediary product

It is very important to elucidate the structure of the intermediate product to understand the mechanism of methyltin degradation. According to the chromatograms of methyltins (Fig. 2), the retention times of TMT, MMT, DMT and the intermediate product are 9.6, 13.7, 17.3 and 11.2 min, respectively. Because MMT, DMT and TMT have the same intermediate product and MMT has the simplest structure, MMT was used to deduce the structure of the intermediate product. Fig. 3 shows that the peak height of the intermediate product increases at first and then decreases during the degradation process of MMT, which means that the intermediate product is an organotin compound because organotin compounds have higher sensitivity using the sulfur interference filter (390 nm) in this study. However, the maximum emission wavelength of Sn–H bonds is at 610 nm (Dagnall et al., 1968) and the sensitivity of FPD for inorganic tin is poor at 390 nm, even no signal for the end product, inorganic tin. The retention time of the intermediate product is shorter than that of MMT in the reversed phase HPLC chromatogram, suggesting that the polarity of the
Fig. 4. Spectra of MMT and the intermediate product using ESI⁻-Q-TOF-MS. (A) the intermediate product, (B) standard MMT, (C) and (D) are the magnification spectra of (A) and (B) from m/z 235 to 360.

Table 1
Ions observed in the negative ESI mass spectra of MMT and the intermediate organotin compound and their structures

<table>
<thead>
<tr>
<th>Major ion (m/z) and structure</th>
<th>Intermediate product: (CH₂OH)Sn⁺³⁻</th>
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<tbody>
<tr>
<td>MMT</td>
<td>[Sn(OH)₄Cl]⁻</td>
</tr>
<tr>
<td>221, 223, 225, 227, 229, 231</td>
<td>[Sn(OH)₃Cl]⁻</td>
</tr>
<tr>
<td>271, 273, 275, 277, 279, 281</td>
<td>[Sn(CH₃)Cl₄]⁻</td>
</tr>
<tr>
<td></td>
<td>[Sn(CH₂OH)(OH)₃Cl]⁻</td>
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<tr>
<td></td>
<td>[Sn(CH₂OH)(OH)₂Cl₂]⁻</td>
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<td></td>
<td>[Sn(CH₂OH)(OH)Cl₃]⁻</td>
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<td>[Sn(CH₂OH)(OH)Cl₄]⁻</td>
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<tr>
<td></td>
<td>[Sn(CH₂OH)(OH)Cl₅]⁻</td>
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<td></td>
<td>[Sn(CH₂OH)Cl₆]⁻</td>
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<td>[Sn(CH₂OH)Cl₇]⁻</td>
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<td>[Sn(CH₂OH)Cl₈]⁻</td>
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<td>[Sn(CH₂OH)Cl₉]⁻</td>
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<tr>
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<td>[Sn(CH₂OH)Cl₁₀]⁻</td>
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<td></td>
<td>[Sn(CH₂OH)Cl₁₁]⁻</td>
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<td></td>
<td>[Sn(CH₂OH)Cl₁₂]⁻</td>
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intermediate product is stronger than that of MMT. Thus, the intermediate product might be \( \text{CH}_2\text{OH}–\text{Sn}^{3+} \), \((\text{CHO})–\text{Sn}^{3+}\) or \(\text{CH}_3\text{O}–\text{Sn}^{3+}\). As \(\text{Sn–C}\) bond is detected at 390 nm (Aue and Flinn, 1977) by QUIL-FPD, the intermediate product should not be \(\text{CH}_3\text{O}–\text{Sn}^{3+}\). During the degradation process, the amount of the formed intermediate product was strongly dependent on the solution pH because its content at pH 8 is greatly higher than that at pH 4 and 6. This suggests that this intermediate product is easier formed and more stable at alkaline solutions, thus methylol tin compound (\(\text{CH}_2\text{OH}–\text{Sn}^{3+}\)) has higher possibility.

To detect the intermediate product and thus elucidate its structure, direct electrospray ionization-Quadrupole-Time of flight-mass spectrometry (ESI-Q-TOF-MS) was used to determine the samples taken from a completely degraded MMT solution which had a higher concentration of the intermediate product (Fig. 3B). The result suggests the intermediate product at pH 3.5 and 8 has equal stability as the same peak height and retention time were obtained from the chromatograms shown in Fig. 3. Therefore, the structure of intermediate product was elucidated at pH 3.5 in order to get stronger signal using ESI-Q-TOF-MS. Electrospray ionization (ESI) is one of the softest ionization techniques, which enables the determination of molecular weight of many organometallic, especially organotin compounds (Rosenberg et al., 2000; Rosenberg, 2003; Apek et al., 2006). Both the positive and negative ionization modes were investigated by ESI-Q-TOF-MS, and results suggested the negative mode has stronger signals than the positive mode (data not shown). Moreover, experiment shows that, with negative ionization mode, better signal was obtained at pH 3.5 than at pH 8 as organotin compounds are easier to complex with \(\text{Cl}^-\) at pH 3.5 to produce a series of charged species. In fact, the detection of a number of charged organotin species is one of the characteristics of electrospray mass spectrometry, which was applied in identification of reaction products organotin compounds \(\text{R}_3\text{SnCl}\) \((\text{R}=\text{Ph, Me or Bu})\) by negative mode (Henderson and Taylor, 1996). Solutions of \(\text{Ph}_3\text{SnX}\) \((X = \text{Cl or Br})\) yields \([\text{Ph}_3\text{SnX}_2]^-, [\text{Ph}_3\text{SnX(OH)}]^-, [\text{Ph}_3\text{SnO}]^-\) and \([\text{Ph}_3\text{SnOSnPh}_3]^-, \text{Me}_3\text{SnCl} \) yields \([\text{Me}_3\text{SnCl}_2]^-, [\text{Me}_3\text{SnOSnMe}_3(OH)]^-\), whereas \(\text{Bu}_3\text{SnCl}\) gives \([\text{Bu}_3\text{SnCl}_2]^-\) and \([\text{Bu}_3\text{SnCl(OH)}]^-\). Various charged species were also detected in this study with negative ESI-Q-TOF-MS at pH 3.5 (Fig. 4 and Table 1). It can be seen from Fig. 4 that MMT and the intermediate product exhibited stronger signals at pH 3.5 and the negative ionization mode. The characteristic isotopic distribution of the tin atom allows clear proof of the presence of tin in these fragment ions because element chlorine has two isotopes with the relative isotopic abundance of \(m/z\) 35 (75.78%) and 37 (24.22%), while element tin has 10 isotopes with relatively higher abundance isotopes of \(m/z\) 116 (14.54%), 118 (24.22%), 120 (32.58%), 122 (4.63%) and 124 (5.79%). Comparing the spectra of MMT and the intermediate product, it can be found that there are no spectra at \(m/z\) 239, 241 and 243, the molecular weight of monomethyltin trichloride (\([\text{CH}_3\text{SnCl}_3]^+\)), which is a neutral molecule and can not produce the signal at the negative mode. This is why there are strong signals at \(m/z\) 273, 275, 277 and 279 (\([\text{CH}_3\text{SnCl}_3]^+\))

Although the intermediate product and MMT have the same ratio of mass/charge at \(m/z\) 273, 275, 277 and 279, the isotopic intensity of \(m/z\) 275 is stronger than \(m/z\) 277 for MMT than for the intermediate product, suggesting they are different compounds and contain different numbers of chlorine atoms. According to the
calculation of a series of molecular weights (Table 1), the intermediate product can be concluded to be methyloltin compound \((\text{CH}_2\text{OH})\text{Sn}^{3+}\) and it is \([\text{Sn(\text{CH}_2\text{OH})(\text{OH})\text{Cl}_3}]\) that give \(m/z\) 273, 275, 277 and 279. In fact, the alcohol derivatives of tributyltin were also found to be the intermediary product in the study of tributyltin photodegradation (Soderquist and Crosby, 1980).

### 3.3. Photodegradation mechanism

As the result of ESI-Q-TOF-MS suggests the intermediate product be methyloltin compound, it is relatively easy to elucidate the possible photodegradation mechanism. MMT produces inorganic tin and methyloltin compound through free hydroxyl radical reaction at pH 4 and 6. However, the reaction mechanism is different at pH 8 as methyloltin compound is always the major product until the complete degradation of MMT. This is due to the production of the hydroxyl radical that was partly inhibited at pH 8 and the hydroxyl radical mainly attacks the methyl group to produce the methyloltin compound first, then the hydroxyl radical attack leads to the cleavage of Sn–C bond and finally produces the inorganic tin (Fig. 5). DMT has a similar photodegradation pathway to MMT, which produce MMT, methyloltin compound and inorganic tin at pH 4 and 6, as well as the same phenomenon as MMT at pH 8 (figure not shown). TMT has different degradation mechanism from MMT and DMT as almost no methyloltin compound was detected even at pH 8. Therefore, the mechanism for TMT is that more methyl groups were simultaneously demethylated for TMT degradation under the UV irradiation.

### 3.4. Effect of salinity on degradation

The degradation rates of methyltin compounds were strongly affected by the salinity of aqueous solutions (Fig. 6). For TMT and DMT, the degradation rates at pH 6 decreased with the increase of NaCl concentrations in the range of 0–3.0\%(m/v). For MMT, however, the degradation rate in solutions with NaCl was much higher than that without NaCl, though the degradation rate also decreased with the increase of NaCl concentrations in the range of 0.5–3.0\%(m/v). It is noteworthy that the NaCl...
almost completely inhibited the degradation of TMT within 500 min. The degradation rate of DMT is greatly reduced by saline. The half-life times in solutions containing NaCl are more than twice of those without NaCl, and it increased with the NaCl concentration in the range of 0.5–3.0% (m/v). This is because Cl\(^-\) can scavenge the free radical in solutions of TMT and DMT. Furthermore, TMT was firstly degraded to DMT and then to other products, thus TMT degradation was greatly inhibited with the increase of NaCl concentration. However, it is difficult to explain the increase of degradation rate for MMT in the solution containing NaCl in comparison to that without NaCl. It is possible that the chloride anion can form ion pairs with alkyltin(IV) cations and their hydrolysed species, such as the formation of (CH\(_3\))\(_3\)Sn(OH)Cl\(^+\) and other Cl\(^-\) mixed species (Stefano et al., 1999), which greatly decreases the bond energy of C–Sn in MMT and lead to the easier degradation of MMT, even though the scavenger function of chloride ion is taken into account.

3.5. Effect of humic acid on degradation

The degradation rates of methyltin compounds were strongly influenced by the humic acid at pH 6 in this study (Fig. 7). It is interesting that the degradation rate of MMT is the zero order at pH 6 and various concentrations humic acid (0–9 mg l\(^{-1}\)), and decreased with increasing of humic acid concentration with half-life times of 95, 113, 180 and 254 min at 0, 3, 6 and 9 mg l\(^{-1}\) of humic acid, respectively. While DMT had slight degradation at the range of 0–9 mg l\(^{-1}\) of humic acid, TMT was almost completely inhibited by the humic acid (3–9 mg l\(^{-1}\)) at pH 6. It is noteworthy that the degradation routes with humic acid are the same as that without humic acid because the same intermediary products were detected. Two major reasons might be responsible for the decrease of methyltin degradation rate by humic acid. First, methyltins can reacts with humic acid to form polymeric species (Omar and Bowen, 1982) as humic acid, a ubiquitous class of electrolytes, can form the complexes with organotin compounds in natural waters through chelation and hydrogen bonding (Oloughlin et al., 2000; Giacalone et al., 2006), which influence strongly the degradation rates of methyltins. For example, Arnold et al. (1998) found that triorganotin cation can form the complexes with negatively charged ligands (i.e., carboxylate and phenolate groups) of the humic acids over the pH range 5.7–9.3. Second, humic acid and its UV degradation products (Garbin et al., 2007) can bind with free radicals and thus function as a powerful free radical scavenger. However, it is not clear why the degradation rate orders are
different from each other for MMT, DMT and TMT. It may be due to form the different complex structures of methyltins and humic acid.

4. Conclusion

The photodegradation of methyltins was systematically studied under different environmental conditions by the application of the newly developed HPLC-FPD hyphenated system. The elucidation of the previously unknown product helps us to better understand the mechanism of the degradation process of methyltin compounds. At the same time, these results can forecast the behavior and fate of methyltins in the environment. It is also important to understand the environmental behavior of other organotin compounds, such as butyltins and phenyltins.

Acknowledgments

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Appendix A. Supplementary material

Supplementary data associated with this article can be found, in the online version, at doi:10.1016/j.chemosphere.2008.02.048.

References