Methylsiloxanes Release from One Landfill through Yearly Cycle and Their Removal Mechanisms (Especially Hydroxylation) In Leachates

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1. INTRODUCTION

Cyclic volatile methylsiloxanes (cVMS)—including octamethylcyclotetrasiloxane (D4), decamethylcyclopentasiloxane (D5), and dodecamethylcyclohexasiloxane (D6)—were the impurities of polydimethylsiloxane (PDMS), which have vast usage in both industrial and personal consumer products as lubricant, defoamer and electrical insulating agent, etc.1-3 In addition, some cVMS (especially D5) could be directly applied in personal care products.5-8 Because usage of cVMS-containing products could lead to their environment emission,1-3 environment occurrence and fate of cVMS had attracted attention of scientists.

During usage by general population, cVMS in some consumer products would be directly released to air (about 90%) and wastewater (about 10%).1-5 Hence they could exist in air, water, and biota samples close to municipal areas.4-10 Different from those in the above products, large amount of cVMS in other PDMS-based products such as elastomers and sealants may, via solid waste disposal, enter municipal landfills,1-3 where up to 95% of municipal solid waste collected worldwide was disposed.11 For example, according to reported emission rates of PDMS-based products and residual monomer contents in these products,1-3 D4–D6 in elastomers and sealants may account for 50.1–65.6% of their total emission from all PDMS-based products in European Union. In addition, despite lack of relevant data, cVMS in personal care product, through disposal of cVMS-containing municipal sludge, may also transfer to landfill. Overall, cVMS emission and fates in landfills are important for understanding the total exposure of cVMS in the global environment.

Until now, there have been some literatures reporting cVMS in landfill gas and leachates in different countries, such as Germany, Sweden, Finland and Canada, etc.12-15 One study reported that the yearly average total VMS emission to air were 80 and 250 kg/y respectively from two different landfills in Canada.16 However, due to limited samples (n = 1–3 for each landfill) or short sampling periods (~two months), the seasonal variation of

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Supporting Information

ABSTRACT: In one yearly cycle (2016), D4 and D5 were detected in biogas samples (n = 36, 0.105–2.33 mg/m³) from a Chinese municipal landfill, while D4–D6 were detected in influents/effluents of leachate storage pond (n = 72, < LOQ–30.5 μg/L). Mass loads of cVMS in both biogas (591–6575 mg/d) and leachate influents (659–5760 mg/d) increased from January to July (summer), and then decreased from July to December (winter). Removal experiments indicated that 1) hydrolysis and volatilization were predominant removal mechanism for D4 and D5, respectively, in leachate storage pond, responsible for their more significant removal (94.5–100%) in August; 2) indirect phototransformation (τ_1/2 = 25.5–87.0 days), such as hydroxylation by OH radical generated in leachates, was the predominant (50.0–75.5%) removal pathway for D6, which led to the largest removal efficiencies (65.2–73.7%) in June, the month with the largest sun light intensity and highest photosensitizer (e.g., Fe^{2+} and NO_3^-) concentrations. Monohydroxylated products of D5 and D6, D4TOH and D5TOH, were detected in leachate effluents (39.6–187 ng/L) during May-July. Compared to D5 and D6, volatilization half-lives of D4TOH (86.3 days) and D5TOH (177 days) in leachates were 2.9 and 1.4 times longer, while their hydrolysis half-lives (7.50 days for D4TOH and 21.5 days for D5TOH) were 7.1 and 10 times shorter, respectively.

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either cVMS concentrations or mass loads were not analyzed in landfill gas and leachates through yearly cycle.

Another data gap is related to cVMS fate in landfill leachates. Before emission to aquatic environment, leachates are always treated on site or transferred to WWTPs. At present, the dominant leachate/wastewater treatment techniques are based on biological treatment processes, where main removal mechanisms of cVMS—volatilization and sorption to sludge—have been determined. However, before being transferred to treatment processes, leachates in most landfills are stored in opened storage pond, where long hydraulic retention time of the wastewater (>24 h ~ 10s of days) and non-neutral (acidic or basic) pH of leachates may lead to significant removal (volatilization and hydrolysis) of cVMS. More important, landfill leachates often contain high concentrations of dissolve organic matter (DOM) and NO_3~−, photosensitizers that could generate hydroxyl radicals (•OH) via photochemical reaction. Furthermore, some acid landfill leachates contained high levels of transition metal ions (such as Fe^{2+}), which could accelerate •OH production rates via photo-Fenton reaction with hydrogen peroxide (a common constituent of natural waterbody). As cVMS in air could be hydroxylated by •OH, the indirect phototransformation (hydroxylation) of cVMS in leachates of the storage pond may also be possible. To the best of our knowledge, up to now, there is no research on the fate of cVMS in leachate storage ponds of any landfills.

In the present study, based on a whole year sampling, occurrence of cVMS in biogas and leachates from one municipal landfill located in China was systematically investigated together with their removal in the coupled leachate storage pond. The main objectives of this study included: (1) to characterize seasonal trends of cVMS emission in landfill; (2) to test the hypothesis that cVMS could be hydroxylated by •OH generated in situ in leachate storage pond; (3) to determine the further removal of hydroxylation products of cVMS in the leachate storage pond.

2. MATERIALS AND METHODS

2.1. Field Sample Collection. The studied municipal landfill, located in the east of Shandong Province, China, was opened at 2012, and processed approximately 220 000 kg/d of solid waste. The catchment area of the landfill covers about 171 km², with a population of 300 000. In this landfill, the leachate from the drainage layer was pumped to one opened storage pond (square = 3000 m², average leachate depth = 0.4 m) with an average hydraulic retention time of ~18 days. Subsequently, the leachate in storage pond was pumped off-site for treatment at one municipal WWTP in the same catchment. Detailed information on seasonal change, physical and chemical characteristics of landfill matrices were provided in SI Table S1.

Thirty six sampling events were carried out from January to December 2016, that is, three events per month. In each event, 24 h composite influent leachate of the storage pond (i.e., the effluent of drainage layer) and coupled effluent leachate were collected using automatic samplers (with silicone-free tubing) at flow proportion mode, respectively. Before pretreatment, the leachate samples were stored in sealed glass tubes (4 L) without headspace at 4 °C. Meanwhile, one 24 h air sample was collected from the exhaust gas pipe of the landfill by using a pump to pull air (100 L/min) through one particle filter followed by precleaned PUF/XAD-2/PUF (10 g, Supelco) cartridge.

The breakthrough of cVMS was determined by mounting a backup cartridge in series with the primary cartridge. At the same dates as landfill sampling (January to December 2016, three events per month), 24 h composite influent (n = 36) and effluent (n = 36) samples were also collected and stored forming one municipal WWTP containing anaerobic-anoxic-oxic processes, which had same catchment with the landfill and treated the leachate effluents from the studied landfill. Notably, the influents of the WWTP were sampled at the site before leachate mixing, that is, leachate was not part of the influent sample of the WWTP.

Wind speeds and light intensities were measured on-site by TP6252A air velocity meter and HI97500 lux/light meter, respectively. Dissolved oxygen (DO), pH and electrical conductivity (EC) values of leachate were determined by Hach HQ40d portable DO/pH/conductivity multi-parameter meter. Dissolved organic carbon (DOC) of leachate were determined by TOC analyzer (TOC-VCPH, Shimadzu, Japan). NO_3~− and total suspended solid (TSS) were monitored using standard methods for waters and wastewater analysis. Total nitrogen (TN) and Fe were measured with persulfate and spectrophotometric methods, respectively.

2.2. Standards and Chemicals. Cyclic methylsiloxanes (D_4, D_5, D_6, purity >98%) were purchased from Sigma-Aldrich (St. Louis, MO). 13C-labeled D_4, D_5, and D_6 were purchased from Cambridge Isotope Laboratories (Andover, MA). 1-Hydroxy-nonamethylcyclotetrasiloxane (D_4TOH, purity >95%) and 1-hydroxy-undecamethylcyclohexasiloxane (DSTOH, purity >95%) were custom-synthesized in Toronto Research Chemicals (Toronto, Ontario) and purified in Shandong University, China. Isopropanol, methanol, acetone, ethyl acetate, and n-hexane were purchased from Fisher Scientific (Fair Lawn, NJ).

2.3. Field Sample Pretreatment and Analysis. Aqueous Samples. Cyclic methylsiloxanes (D_4, D_5, D_6) and hydroxylated methylsiloxanes (D_4TOH, D_5TOH) in aqueous samples were extracted by liquid–liquid extraction. 13C–D_4, 13C–D_5 and 13C–D_6 were used as internal standards for D_4, D_5/D_4TOH and D_6/DSTOH, respectively. 100 mL of sample spiked with acetone solution of internal standards (40 μL, 500 μg/L) was extracted with 25 mL of n-hexane followed by 20 mL of n-hexane/ethyl acetate (1:1 by volume). Subsequently, the organic layer was transferred to a flat-bottom flask and concentrated to 8–10 mL by evaporation at 30 °C under a stream of nitrogen. The remaining extract was purified by passing through 1.0 g of anhydrous sodium sulfate cartridge, and then concentrated to 0.5 mL with the same way mentioned above, and diluted to 1 mL with n-hexane/ethyl acetate (1:1 by volume).

Air Samples. After sampling, the PUF/XAD-2/PUF sandwich was spiked with acetone solution of internal standards (100 μL, 500 μg/L, 13C–D_4, 13C–D_5, and 13C–D_6), and then extracted three times with 30 mL of ethyl acetate/n-hexane mixture (1:1 by volume) by sonication for 1 h. The total extract was concentrated to 1.0 mL with a gentle stream of nitrogen before analysis.

Full Scan analysis. Full scan chromatograms (m/z range 50–600) for extracted compounds in leachate were analyzed by Agilent 7200 Q-TOF GC/MS, which was equipped with HP-5MS 30 m × 0.25 mm × 0.25 μm. Ion source temperature was 230 °C for both electron-impact ionization (EI) and positive chemical ionization (PCI) mode. The EI and PCI mass spectra of D_4TOH and DSTOH, obtained by Q-TOF GC/MS, were described in SI Section 2.
**SIM Analysis.** The concentrations of target compounds in leachate/air were acquired by selected ion monitoring (SIM) of GC-MS analysis using Agilent 7890A gas chromatograph -5975C mass spectrometric detector equipped with HP-5MS column (30 m x 0.25 mm x 0.25 μm). The MS was operated in electron-impact ionization mode (EI) at 70 eV, ion source temperature of 230 °C and quadrupole temperature of 150 °C. MS parameters for compounds were summarized in SI Table S2.

2.4. Quality Assurance/Quality Control (QA/QC). Similar to our previous studies, we made some efforts to reduce siloxanes contamination during sample collection and analysis. First, the analyst did not use consumer products containing siloxanes. Second, prior to use, all glass tubes were cleaned with hexane, and then heated at 300 °C for 10 h. Third, prior to use, both anhydrous sodium sulfate and PUF/XAD-2/PUF cartridges were immersed in hexane for 4 h, and subsequently rinsed with 10 mL of hexane. After rinsing, the cartridges were dried using purified nitrogen and stored in capped glass tubes. Fourth, only steel pipes, but not silicone pipes, were used for nitrogen blow process. Fifth, all field blanks were collected to assess potential ambient contamination—the field blanks of air were collected by exposing PUF/XAD-2/PUF cartridges for 1 min at the sampling site, while the aqueous field blanks (leachate and wastewater) were prepared with a water solution of siloxane-free commercial humic acid (Sigma-Aldrich, St. Louis, MO). Finally, for every five air or aqueous samples, a procedural blank was prepared and analyzed.

D4TOH and D5OH were not found in any field or procedural blanks of the aqueous samples. Their limits of quantitation (LOQs), determined as 10 times the standard deviation of signals of procedural blank samples (n = 7) spiked with target compounds (100 ng/L), were 4.5–7.5 ng/L in aqueous sample (SI Table S3). Recoveries of D4TOH and D5TOH in aqueous samples were 88–94% (SI Table S3).

Mean concentrations of D4, D5 and D6 were 1.2–3.9 ng/L in aqueous field blanks, and 0.9–3.2 ng/m³ in air field blanks, 0.6–2.4 ng/L in aqueous procedural blanks and 0.3–2.2 ng/m³ in air procedural blanks, respectively (SI Table S3). LOQs for D4-D6, determined as 10 times the standard deviation of the procedural blank signals (n = 7), were 1.7–2.8 ng/L in aqueous samples, while 2.2–4.5 ng/m³ in air samples. Recoveries of cVMS in aqueous and air samples were 88–94% and 83–94%, respectively (SI Table S3). All reported concentrations of D4, D5 and D6 in this study were reported with their field blank values subtracted.

2.5. Simulated Experiments. 2.5.1. Elimination of Methylsiloxanes in Leachate. As all effluent leaches of storage pond contained high levels (>100 ng/L, SI Table S4) of D5 and D6, their standard solutions were not added in leachates during this experiment. However, 40 μL of acetone solution of D4 (1.00 mg/L) were added into 24 samples (collected during April 6 to November 29) due to its low original concentrations (<LOD-95.7 ng/L). For each of effluent leachate samples (n = 36) collected from the storage pond during 12 months, one set of 15 opened glass vials (40 mL) containing 40 mL of leachate were incubated under the similar illumination intensity, wind speed and air/leachate temperature as the corresponding values measured when the effluent leachate sample was collected (SI Table S1). The light source was solar simulator with 500 W xenon Lamps, and simulated illumination intensities were obtained by adjusting electric current of light source. At each of the predetermined time intervals (0, 2, 10, 24, and 48 days), three vials were taken to determine concentrations of D4, D5, and D6. Meanwhile, two additional sets of 15 glass vials - one capped and another open set - were also incubated under the same conditions except they were not illuminated. Furthermore, for evaluating methylsiloxanes hydroxylation arose from hydroxyl radical, the fourth set, opened under illumination and spiked with 100 μg/L of isopropanol as OH scavenger was prepared for each of the three leachate effluents collected during June. Meanwhile, the full scan chromatograms for transformation products of D5 and D6 in leachates under illumination were analyzed by Agilent 7200 Q-TOF GC/MS with method mentioned in Section 2.3.

2.5.2. Phototransformation of D5 and D6 in Aqueous Environment. Aqueous samples for phototransformation experiments were prepared by diluting the stock solution of DOM (DOC = 10.0 g/L), which was concentrated from the leachate effluent of the studied storage pond (as described in the section 2.2), with Milli-Q ultrapure water. Since dissolved oxygen - the important participating substance for DOM photochemical reaction - may be reduced with the increase of illumination time in caged leachates, the phototransformation experiments of D5 and D6 were performed in opened systems. Aqueous samples (40 mL), with different levels of DOM (DOC = 0–1000 mg/L, pH 7.0, NO3⁻ = 0, Fe²⁺ = 0), NO3⁻ (0–2000 mg/L, NaNO3), FeSO4, DOC = 1000 mg/L, pH 7.0, Fe²⁺ = 0) and Fe²⁺ (0–2000 mg/L, FeSO4, DOC = 1000 mg/L, NO3⁻ = 0, pH 7.0), were spiked with acetone solution (40 μL, 500 μg/L) of D5 and D6, respectively, and then incubated in 15 opened glass vials under different light intensities (0–80000 lx). At each of the predetermined time intervals (0, 2, 10, 24, and 48 days), three vials were taken to determine concentrations of D5 and D6. To subtract volatilization and hydrolysis rates of D5 and D6, another paired set of 15 opened glass vials were incubated under non-illumination.

2.5.3. Elimination of Hydroxylated Methylsiloxanes (D4TOH and D5TOH) in Leachates. The stock solution of DOM (Section 2.5.2), spiked with FeSO4 and NaNO3, was diluted to simulate leachates (DOC = 2000 mg/L, temperature = 21.0 °C, pH 6.5, Fe²⁺=2000 mg/L, NaNO3 = 1400 mg/L) collected from effluent of the storage pond in June, when fastest productions of D4TOH and D5TOH were found, as discussed in Section 3.2.

For D4TOH and D5TOH, four sets of 15 glass vials (40 mL) - one set opened under illumination (light intensity = 65 000 lx), one set opened under nonillumination, one set capped under nonillumination, and one set (sterilized by 1% NaN3) capped under nonillumination - were filled with simulated leachate (40 mL) and acetone solutions of hydroxylated methylsiloxane (100 μL, 200 μg/L), respectively. For each of glass vials (n = 15), three vials were sacrificed for the determination of the remaining cVMS concentrations at each of the predetermined time intervals (0, 2, 10, 24, and 48 days). Meanwhile, in set capped under nonillumination, dimethylsilanediol [Me₂Si(OH)₂] and methylsilanetriol [MeSi(OH)₃] were also analyzed with methods described in the section 2.2.

3. RESULTS AND DISCUSSION

3.1. Methylsiloxanes in Landfill Gas and Influent Leachate. Occurrence. D4, D5 and D6 were detected in all influent samples (n = 36) of storage pond collected in 2016, with concentrations ranging from 0.338 to 7.18 μg/L, 4.23–30.5 μg/L, and 1.00–14.7 μg/L, respectively (SI Table S4), which were similar to their reported concentrations in leachates from landfills.
located in Sweden (1−2 μg/L for D4, 0.1−0.4 μg/L for D5) and Finland (3.9 μg/L for D5, 1.7 μg/L for D6).\(^{12,15}\)

In all landfill gas samples (n = 36), D4 (0.753−2.33 mg/m³) and D5 (0.105−0.738 mg/m³) were detected, while D6 concentrations were all below detection limits. D4 and D5 concentration in biogas were similar to those in biogas samples from Canada (1.27−5.03 mg/m³ for D4 and 0.395−3.33 mg/m³ for D5) and Germany (4.24−8.84 mg/m³ for D4 and 0.4−1.09 mg/m³ for D5), etc.\(^{12,15}\)

**Mass Load to the Local Environment.** The annual mass load of cVMS in leachate/biogas was calculated using the following equation:

\[
M = \sum_{i=1}^{12} \left\{ \left[ C_{i(1)} \times V_{i(1)} + C_{i(2)} \times V_{i(2)} + C_{i(3)} \times V_{i(3)}\right]/3 \right\} \times d_i
\]

Where \(i\) represented number (1−12) of sampling month; \(C_{i(1)}−C_{i(3)}\) represented the concentrations of target compounds in three leachate/biogas samples from the \(i\) month; \(V_{i(1)}−V_{i(3)}\) represents the daily volume rates of leachate/biogas during three sampling events in the \(i\) month; \(d_i\) represents the total days of the \(i\) month. It was too difficult to collect leachate/biogas samples in all days (n = 366) of the tested year, and the sampling times in the present study were relatively limited - 3 days for each month. Therefore, there may be some deviations between our estimated annual mass loads of cVMS and the actual values.

Totally, the annual mass loads were about 84.5 g for D4, 435 g for D5, and 160 g for D6 in influent of storage pond, as they were 664 g for D4 and 155 g for D5 in biogas, respectively. Through dividing their total mass loads in leachate and biogas by served population (300 000) of the landfill, the calculated annual emission to landfill (leachate+biogas) per capita was 2.50 mg for D4, 1.97 mg for D5 and 0.533 mg for D6, respectively.

For the total environment of any urban locations, cVMS may also be released to WWTP. cVMS in municipal WWTPs were mainly attributed to their residuals in cosmetics, personal care and household products used by general public, while 90% of which may already volatilized to the air compartment before they reach the wastewater stream.\(^{1−3}\) For the study location, we detected cVMS concentrations of 0.164−0.353 μg/L for D4, 1.33−2.90 μg/L for D5 and 0.412−0.941 μg/L for D6 (SI Table S5) in influents of one municipal wastewater treatment plant having the same catchment with the studied landfill. Their annual mass loads per capita through the influents to the WWTP were 10.5 mg for D4, 84.5 mg for D5 and 26.4 mg for D6. Therefore, we roughly estimated that annual total emission rates (to air before reaching WWTP and influent wastewater of WWTP) of D4, D5, and D6 per capita through using cosmetics, personal care and household products in the studied area were 105 mg for D4, 845 mg for D5 and 264 mg for D6, respectively—about 31−362 times larger than emission rates of cVMS from solid siloxane-containing wastes disposal.

**Temporal Trend.** Figure 1 showed that at 2016, the mass load of ∑D4−D6 had increasing trends both in leachates (from 1234 to 5760 mg/d) and biogas (from 591 to 6575 mg/d) from January to July (Summer), while decreasing trends both in leachates (from 5760 to 659 mg/d) and biogas (from 6575 to 680 mg/d) during July−December (Winter). Logarithm of emission rates of cVMS in both biogas and leachates had statistical correlations with several characteristics of influent leachate and air (SI Table S6), such as DOC, air/leachate temperature, EC, NO₃⁻, etc. However, kinds of characteristics had bivariate correlations (SI Table S6), and their principal components analysis was not suitable, cumulative percentage of variance explained by components with eigenvalues >1 were limited (74.65%, SI Table S7). Therefore, it was too difficult to comprehensively discuss influence of all kinds of leachate characteristics on cVMS emission rates. In this section, we merely evaluated influences of temperature and DOC on cVMS emission.

Due to seasonal change, air and leachate temperatures increased from −3.4 to 35 °C and from 5.43 to 28.9 °C during January to July, respectively, while decreased from 35 to 2.3 °C and from 28.9 to 5.02 °C during July−December, respectively. High temperatures should accelerate desorption rates of cVMS from solid waste.\(^{1−3}\) In the present study, the logarithm of mass load of D4 (slope = −2.802, R² = 0.855, p < 0.05) and D5 (slope = −3.343, R² = 0.7184, p < 0.05) in biogas had inversely relations with reciprocal of leachate temperatures (SI Figure S1). Meanwhile, higher leachate temperature would accelerate anaerobic degradation, which turn the solid organic matter with high molecular weight into DOC with low molecular weight.\(^{35}\) Logarithm of DOC mass load in influent leachates had inversely relations (slope = −1.957, R² = 0.6754, p < 0.05, SI Figure S2) with reciprocal of leachate temperatures. Dissolved organic matter also could elevate the release of methylsiloxanes from solid phase.\(^{35}\) In the present study, the logarithm of mass load of cVMS in both biogas and leachate had positive relations (slope = 0.0004−0.0008, R² = 0.5138−0.841, p < 0.05) with DOC concentrations (SI Figure S3).

**3.2. Time Trends of Methylsiloxanes Elimination in the Leachate Storage Pond.** Concentrations of cVMS were 3.21−459 ng/L for D4 [detection frequencies (df) = 83.3%], 0.121−5.97 μg/L for D5 (df = 100%) and 0.474−4.47 μg/L for D6 (df = 100%) in effluents of leachate storage pond, while 3.67−11.6 ng/L for D4 (df = 100%), 55.9−133 ng/L for D5 (df = 100%), and 12.3−38.1 ng/L for D6 (df = 100%) in effluents of the studied WWTP, respectively (SI Table S4 and S5).

The removal efficiencies of cVMS in storage pond (RERP) and WWTP (REWWTP) were calculated with the following equations:

\[
\text{RE}_{\text{RP}} = 100\% \times \left( M_{\text{inf-RP}} - M_{\text{eff-RP}} \right) / M_{\text{inf-RP}}
\]

\[
\text{RE}_{\text{WWTP}} = 100\% \times \left( M_{\text{eff-RP}} + M_{\text{inf-WWTP}} - M_{\text{eff-WWTP}} \right) / \left( M_{\text{inf-RP}} + M_{\text{inf-WWTP}} \right)
\]

Where \(M_{\text{inf-RP}}\) and \(M_{\text{eff-RP}}\) represent mass load of cVMS in influent and effluent of storage pond, whereas \(M_{\text{inf-WWTP}}\) and \(M_{\text{eff-WWTP}}\) represent mass load of cVMS in influent and effluent of WWTP, respectively.

Generally, removal efficiencies (SI Table S8) were for 81.1−100% (mean = 94.9%) for D4, 57.1−98.1% (mean = 73.3%) for D5 and 35.4−73.7% (mean = 53.2%) for D6 in leachate storage pond, while 96.3−98.1% (mean = 97.5%) for D4, 95.7−97.0% (mean = 95.9%) for D5 and 95.7−97.4% (mean = 96.6%) for D6 in the WWTP, respectively. As previous literatures had determined the main removal mechanisms of cVMS in WWTPs—volatilization and sorption to sludge,\(^{1−9}\) the main focus of the present study is the cVMS removal in leachate storage pond as discussed in the following paragraphs.

In storage pond, the most significant removals of D4 (RE_{RP} = 99.8−100%, 99.9% at mean) and D5 (RE_{RP} = 94.5−98.1%, 96.2% at mean) were found in August, while D6 was fastest removed (RE_{RP} = 65.2−73.7%, 70.0% at mean) in June (Figure 1, SI Table S8). Based on incubating all effluent leachate
samples ($n = 36$) collected from storage pond in the whole year under various conditions (e.g., capped or opened; illumination or nonillumination), we made semilog plots of cVMS concentrations versus incubation time (SI Figure S4), and then calculated their first-order rates of volatilization, hydrolysis, and phototransformation in these leachates: (1) hydrolysis rates of cVMS were their slopes of semilog plots in capped system under nonillumination; (2) volatilization rates were calculated by subtracting the slopes of semilog plots in the capped system under nonillumination from the values in paired opened system under nonillumination; (3) phototransformation rates were calculated by subtracting the slopes of semilog plots in the opened system under nonillumination from the values in paired opened system under illumination.

D4. Overall, hydrolysis ($k = 3.34 \times 10^{-2} \text{ d}^{-1}$ at mean) was the predominant removal mechanism for D4 in the whole year, followed by volatilization ($k = 2.93 \times 10^{-2} \text{ d}^{-1}$ at mean). Degradation rates of D4 in leachate under nonillumination were

![Figure 1. mass loads of cVMS in biogas/leachate, and their removal efficiencies in leachate storage pond.](image-url)
approximately equal to those under illumination, indicating its poor in-/directed phototransformation of D4. Higher air temperature (33.3 °C at mean) in August perhaps lead to larger usage of alkaline domestic products (such as soaps, body washes and detergent, etc.) in the studied area, which may explain the higher pH (mean = 8.0) of leachates in this month than those pH values (6.3–7.8, mean = 7.2) for the rest of the year. Hydrolysis rates (k = 4.43 × 10⁻² – 1.88 × 10⁻¹ d⁻¹) of D4 during July to September 7, especially in August (k = 1.04 × 10⁻¹ – 1.88 × 10⁻¹ d⁻¹), were faster than those (k = 9.71 × 10⁻⁴ – 3.67 × 10⁻² d⁻¹) in the rest of the year (Figure 2), perhaps because of both high leachate pH (7.6–8.1) and temperature (24.0–28.5 °C, mean = 26.7 °C) in this period. As leachate pH increased from 7.6 to 8.1, the hydrolysis rates of D4 increased 4.3 times, which was in the same order of magnitude as the calculated value (3.2 times) based on the reported acid- (k = 3.91 × 10⁵ at 25 °C) and base- (k = 7.10 × 10⁴ at 25 °C) catalyzed reaction rate constants of D₄. In addition, volatilization rates of D₄ in August (k = 5.78 × 10⁻⁵–6.78 × 10⁻² d⁻¹, that is, t₁/₂ = 4.44–52.2 days) were also faster than those in other months (k = 1.20 × 10⁻² – 5.25 × 10⁻¹ d⁻¹, that is, t₁/₂ = 5.73–25.0 days), which should be attributed to higher air/leachate temperature in this month.  

The logarithm of volatilization rates of cVMS had been found in this month. As DOM concentrations increased from 10 to 200 mg/L, elimination rates of D₅ and D₆ in leachate were faster than those in other months (mean = 1733 mg/L for DOM, mean = 1412 mg/L for NO₃⁻) in June than those in other months (1125 mg/L for DOM, 797 mg/L for NO₃⁻, mean value), perhaps because high leachate temperature (mean = 21.2 °C) in June would elevacth analogous degradation of solid organic matter to dissolved organic matter and nitrification of ammonia nitrogen to NO₃⁻.  

Second, faster anaerobic degradation of solid organic matter may produce dissolved organic matter (including organic acids) in June and reduced leachate pH (pH 6.3–6.6), which, in turn, may lead to Fe²⁺ accumulate in leachate, Fe²⁺ concentrations (mean = 1976 mg/L) in June were higher than those in other months (mean = 670 mg/L). Reduced Fe ions (Fe²⁺) also could accelerate the photoproduction of hydroxyl radicals by reacting with H₂O₂, another photoproduc of DOM. Third, light intensities (mean = 686, unit: 100 lx) were highest in June in the studied area (36.58° N, 120.42° E), which could accelerate photo-oxidation of DOM and NO₃⁻ to OH radicals.

**Phototransformation of cVMS in Simulated Leachate.** To further test our speculation about factors accelerating methylsiloxanes phototransformation in leachate, we investigated the influences of DOM, NO₃⁻, Fe²⁺ and light intensities on hydroxylation rates of D₅ and D₆ through illumination experiments (Figure 3). As DOM concentrations increased from 10 to 200 mg/L, elimination rates of D₅ and D₆ in leachate increased 3.8 and 4.4 times, while decreased 1.5 times and 1.5 times as DOM increasing from 10 to 200 mg/L, respectively, perhaps because OH scavenging of DOM [39] with too high concentrations would be faster than its OH-generation. This result indicated that faster phototransformation of D₅ and D₆ in June should not be mainly attributed to high DOM concentrations (mean = 1733 mg/L) in this month. Logarithms of elimination rate constants for both D₅ and D₆ were positively related to the logarithms of NO₃⁻ (the slope = 0.062 for D₅ and 0.064 for D₆), Fe²⁺ (the slope = 0.0779 for D₅ and 0.0816 for D₆) and light intensity (the slope = 0.2414 for D₅ and 0.3616 for D₆), indicating that their faster phototransformation rates in June may arise from higher concentrations of NO₃⁻ and Fe²⁺ and light intensities in this month.
3.3. Simulated Experiments for Further Elimination of Hydroxylation Products of Methylsiloxanes in Leachate.

Section 3.2 showed that D5 and D6 in leachates could undergo hydroxylation to D4TOH and D5TOH, respectively. Overall, D4TOH and D5TOH were not detected in any leachate in influent samples, but detected in six leachate effluents collected during May–July, with concentrations in the ranges of 39.6–58.2 ng/L and 104–187 ng/L, respectively (SI Table S4). Based on their phototransformation rates in leachates during May–June (Figure 2) and hydraulic retention time of storage pond (18 days), we estimated that Si mass ratios of phototransformation products of D5 and D6 to their paired cVMS should be 10.6−15.0% (mean = 13.1%) and 55.1−82.4% (mean = 66.8%), respectively, which were 13−19 times larger than Si mass ratios of D4TOH.
and D5TOH (3.01–6.39%, 4.41% at mean) to parent cVMS in the collected leachates. The above phenomenon could be caused by several reasons, for examples: (1) phototransformation products of D5 and D6 included other compounds besides D4TOH and D5TOH; (2) D4TOH and D5TOH were unstable and would undergo further removal in storage pond. In this section, we focused on removal mechanisms of D4TOH and D5TOH in leachates through simulated experiments (Section 2.5.4). Based on semilog plots of their concentrations versus incubation time (SI Figure S8), the first-order volatilization and degradation rates of D4TOH and D5TOH were calculated with the same methods for cVMS (Section 3.2).

Overall, half-lives of D4TOH (6.66 days) and D5TOH (15.4 days) in opened leachates under illumination were 2.4 and 1.4 times shorter than those of paired methylsiloxanes (16.0 days for D5, 22.0 days for D6), respectively (Table 1). However, volatilization rates of D4TOH ($k = 3.49 \times 10^{-3} \text{ d}^{-1}$, $t_{1/2} = 86.3$ days) and D5TOH ($k = 1.70 \times 10^{-3} \text{ d}^{-1}$, $t_{1/2} = 177$ days) were 2.9 and 1.4 times lower than those of paired D5 ($k = 1.02 \times 10^{-2} \text{ d}^{-1}$, $t_{1/2} = 29.4$ days) and D6 ($k = 2.33 \times 10^{-3} \text{ d}^{-1}$, $t_{1/2} = 129$ days), respectively (Table 1), which should be because $-\text{OH}$ has stronger hydrophilicity than $-\text{CH}_3$. These results meant that besides volatilization, there would be competing removal mechanism (degradation) for D4TOH and D5TOH, less significant for D5 and D6 under such conditions.

Under nonillumination, the removal rates of D4TOH and D5TOH in capped leachate were approximately equal to those in paired sterile samples (SI Figure S8), indicating their negligible anaerobic biodegradation - the dominant biodegradation pathway of organic chemicals in leachate. After incubation for 48 days in capped leachate under nonillumination, total silicon mass of dimethylsilanediol and methylsilanetriol approximately accounted for 51.5% and 40.4% of spiked D4TOH and D5TOH, respectively (SI Figure S9), suggesting that hydrolysis was one important elimination pathways of D4TOH and D5TOH in such condition. Because $-\text{OH}$ had stronger electrophilicity than $-\text{CH}_3$, $\text{Si}-\text{O}$ bond with one branch of $-\text{OH}$ would be more polar and easier to undergo bond cleavage via acid/alkaline catalysis than $\text{Si}-\text{O}$ bond merely linked with branches of $-\text{CH}_3$, which may explain why hydrolysis rates of D4TOH ($k = 4.01 \times 10^{-2} \text{ d}^{-1}$, $t_{1/2} = 7.50$ days) and D5TOH ($k = 1.40 \times 10^{-2} \text{ d}^{-1}$, $t_{1/2} = 21.5$ days) were 7.1 and 10 times faster than those paired methylsiloxanes D5 ($k = 5.62 \times 10^{-3} \text{ d}^{-1}$, $t_{1/2} = 53.6$ days) and D6 ($k = 1.41 \times 10^{-3} \text{ d}^{-1}$, $t_{1/2} = 214$ days), respectively (Table 1).

![Figure 3. Correlations between logarithms of the hydroxylation rate constants (unit: 1/d) of cVMS and logarithms of leachate chemical characteristics (DOC, Fe$^{2+}$ and NO$_3^-$) and light intensity](image)

Table 1. First-Order Removal Rates of Compounds in Simulated Leachate

| Compound | Total removal | Volatilization | Hydrolysis | Phototransformation |
|----------|---------------|----------------|------------|---------------------|
|          | rate constant ($d^{-1}$) | half-life (days) | rate constant ($d^{-1}$) | half-life (days) | rate constant ($d^{-1}$) | half-life (days) | rate constant ($d^{-1}$) | half-life (days) |
| D4TOH    | $4.49 \times 10^{-2}$ | 6.66          | $3.49 \times 10^{-3}$ | 86.3        | $4.01 \times 10^{-2}$ | 7.50          | $1.30 \times 10^{-3}$ | 231          |
| D5TOH    | $1.96 \times 10^{-2}$ | 15.4          | $1.70 \times 10^{-3}$ | 177        | $1.40 \times 10^{-2}$ | 21.5          | $3.87 \times 10^{-3}$ | 77.8          |
| D5       | $1.88 \times 10^{-2}$ | 16.0          | $1.02 \times 10^{-3}$ | 29.4        | $5.62 \times 10^{-3}$ | 53.6          | $2.98 \times 10^{-3}$ | 101          |
| D6       | $1.37 \times 10^{-2}$ | 22.0          | $2.33 \times 10^{-3}$ | 129        | $1.41 \times 10^{-3}$ | 214          | $9.96 \times 10^{-3}$ | 30.2          |
The calculated phototransformation rates of D4TOH ($k = 1.30 \times 10^{-3} \text{ d}^{-1}$, $t_{1/2} = 231$ days) and D5TOH ($k = 3.87 \times 10^{-3} \text{ d}^{-1}$, $t_{1/2} = 77.8$ days) were 2.3 and 2.6 times lower than those of their corresponding D5 ($k = 2.98 \times 10^{-3} \text{ d}^{-1}$, $t_{1/2} = 101$ days) and D6 ($k = 9.96 \times 10^{-3} \text{ d}^{-1}$, $t_{1/2} = 30.2$ days), respectively (Table 1), which could explain why no other hydroxylated product of D5 and D6, besides monohydroxylated products (D4TOH and D5TOH), was found in both EI and PCI full scan chromatogram for leachates under illumination (SI Figure S7).

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