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#### **Contaminants of Emerging Concern in Fresh Leachate from Landfills in the Conterminous United States**



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Landfills are the final repository for a heterogeneous mixture of waste from residential, industrial, and commercial sources; and thus, have the potential to produce leachate containing a complex mixture of contaminants of emerging concern (CECs). This leachate is often discharged to pathways that ultimately lead to the environment (e.g. groundwater, streams, and receiving waters such as wastewater treatment plants). To provide the first national-scale assessment of CECs in landfill leachate across the United States, leachate samples from 19 landfills in 16 states were collected and analyzed for 202 CECs. This work summarizes the frequency of detections and concentration of CECs, and describes relations between CECs and selected landfill characteristics (*e.g*. waste composition, location, age of waste, waste load, and leachate production).

# **Contaminants of Emerging Concern in Fresh Leachate from Landfills in the Conterminous United States**

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#### **ABSTRACT**

To better understand the composition of contaminants of emerging concern (CECs) in landfill leachate, fresh leachate from 19 landfills was sampled across the United States during 2011. The sampled network included 12 municipal and 7 private landfills with varying landfill waste compositions, geographic and climatic settings, ages of waste, waste loads, and leachate production. A total of 129 out of 202 CECs were detected during this study, including 62 prescription pharmaceuticals, 23 industrial chemicals, 18 nonprescription pharmaceuticals, 16 household chemicals, 6 steroid hormone chemicals, and 4 plant/animal sterols. CECs were detected in every leachate sample, with the total number of detected CECs in samples ranging from 6 to 82 (median = 31). Bisphenol A (BPA), cotinine, and N,N-diethyltoluamide (DEET) were the most frequently detected CECs, being found in 95% of the leachate samples, followed by lidocaine (89%) and camphor (84%). Other frequently detected CECs included benzophenone, naphthalene, and amphetamine, each detected in 79% of of the leachate samples. CEC concentrations spanned six orders of magnitude, ranging from ng/L to mg/L. Industrial and household chemicals were measured in the greatest concentrations and composing more than 82% of the total measured CEC concentrations. Maximum concentrations for three household and industrial chemicals, *para*-cresol (7,020,000 ng/L), BPA (6,380,000 ng/L), and phenol (1,550,000 ng/L), were the largest measured, with these CECs composing 70% of the total measured CEC concentrations. Nonprescription pharmaceuticals represented 12%, plant/animal sterols 4%, prescription pharmaceuticals 1%, and steroid hormone chemicals <1% of the total measured CEC concentrations. Leachate from landfills in areas receiving greater amounts of precipitation had greater frequencies of CEC detections and concentrations in leachate than landfills receiving less precipitation.

#### **INTRODUCTION**

Landfills are commonly the final repository for heterogeneous mixtures of municipal solid and liquid waste composed of discarded materials from residential, commercial, and industrial sources. Use of landfills as a means of waste disposal will likely increase as the global 5 population increases and nations develop.<sup>1,2</sup> Whereas the number of active landfills in the United States decreased from about 7,900 in 1988 to 1,900 in 2009, the average landfill size has 7 increased.<sup>3</sup> Despite advancements in recycling, source reduction, and composting, the amount of municipal solid waste discarded in U.S. landfills increased from 150 million tons in 1985 to 165 9 million tons in 2010.<sup>4</sup> Because of the complexity and heterogeneity of such waste, landfills receiving such waste have the potential to produce leachate containing numerous organic chemicals including contaminants of emerging concern (CECs) such as pharmaceuticals, plasticizers, personal care products, and steroid hormones. Current criteria for landfill monitoring in the Code of Federal Regulations, Part 258, does not include analysis for many CECs, such as pharmaceuticals, personal care products, and steroid hormones. Such CECs are receiving growing attention as mounting evidence documents their presence in aquatic and terrestrial ecosystems from a variety of urban, industrial, agricultural, and other anthropogenic sources.<sup>5</sup> Although the environmental occurrence of CECs is now recognized as a global phenomenon.<sup>6</sup> 18 much is yet to be understood regarding the fate and effects of these chemicals.<sup>7, 8</sup> Nevertheless, there is a growing body of evidence indicating that exposure to CECs can result in deleterious 20 effects to ecosystem health. $9-17$ 

Although the chemical composition of leachate from landfills has been extensively studied, most research to date has focused on inorganic constituents and, to a lesser extent, some 23 xenobiotic organic chemicals.<sup>18-29</sup> More recently, however, studies characterizing the

1 composition of CECs in landfill leachate have been conducted.<sup>30-35</sup> In general, such research was limited in the number of landfills being investigated and/or the number of CECs analyzed. For example, leachate sampled from three landfill cells containing waste of different ages had 28 of 69 targeted CECs detected in one or more samples with concentrations ranging from 110 to 5 114,000 ng/L.<sup>30</sup> In another study of leachate from three landfills of unspecified location, CECs 6 were detected at concentrations up to  $6,230$  ng/L.<sup>33</sup> Fluorochemicals (used as coatings on paper, packaging, textiles, and carpets) were detected in leachate samples from four municipal landfills 8 with concentrations ranging from 2,300 to 2,800 ng/ $L^{34}$  Pharmaceuticals were measured at concentrations of up to 8.1 mg/kg in municipal solid waste sampled from one waste transfer station.<sup>35</sup> These studies indicate that landfills can be sources of CECs, but much remains unknown regarding the occurrence of broader suites of CECs in landfill leachate on a national scale.

To provide the first national-scale assessment of CECs in landfill leachate across the United States, fresh leachate samples from 19 landfills in 16 states (Figure 1) were collected in 2011 and analyzed for 202 CECs. The analyzed CECs included 100 prescription pharmaceuticals, 33 industrial chemicals, 30 household chemicals (includes ten pesticides), 19 nonprescription pharmaceuticals, 16 steroid hormone chemicals, and 4 plant/animal sterols. The targeted CECs were selected for analysis because they were expected to be persistent in the environment; are used, excreted, or disposed of in substantial quantities; may have human or environmental health effects; or are potential indicators of environmentally relevant classes of chemicals or source materials. This paper summarizes the frequency of CEC detections and concentrations and describes relations between occurrence and selected landfill characteristics

(*e.g*. waste compositions, geographic settings, ages of waste, waste loads, and leachate production).

Figure 1. Near here…

#### **LANDFILL SITES**

For this study, fresh leachate samples (leachate from the beginning of the liquid-waste stream emanating from the waste source before any storage or treatment processes) were collected during the summer and fall of 2011. The criteria used to select landfills for sampling were that the landfills be: (1) active (non-closure or post-closure status), (2) permitted to accept municipal solid waste, (3) receivers of non-hazardous commercial and industrial waste, and (4) equipped with leachate-collection and recovery systems. Fresh leachate was collected from composite or discrete landfill cells prior to on-site pretreatment or off-site discharge. The sampling network consisted of 12 municipal and 7 private landfills representative of landfills across the country that contained a heterogeneous mixture of municipal, construction debris, wastewater sludge (biosolids), and non-hazardous commercial and industrial waste (Figure 2). Common materials in municipal waste typically consist of paper products (28%), food scraps (14%), yard trimmings (14%), plastics (12%), metals (9%), rubber, leather, and textiles (8%), 18 wood (6%), glass (5%), and other miscellaneous waste (4%).<sup>3</sup> The sampling network consisted of a range of landfill sizes, both in terms of amount of annual leachate produced and waste load (Figure 3). The average age range of waste in all sampled landfill cells was 2 - 19 years (Table 1S). In addition, a variety of leachate handling and disposal practices were used at the sampled landfills. Thirteen landfills discharged leachate to a waste water treatment plant (WWTP) through direct plumbing to sewer lines or transport by tanker truck. Five landfills recycled

leachate by (1) spraying on the landfill cap, (2) pumping leachate into injection wells in the landfill, (3) evaporating leachate stored in lagoons, or (4) evaporating leachate by injection into gas flares. One landfill used on-site facultative aerobic treatment prior to leachate discharge to a river. Landfill characteristics were compiled for each of the 19 landfills (Table 1S).

**Figure 2.** Near here…..

**Figure 3.** Near here….

## **SAMPLING, ANAYLTICAL, QUALITY ASSURANCE, AND STATISTICAL METHODS**

Leachate samples were collected using standardized protocols and procedures by environmental sampling staff from the U.S. Geological Survey, State environmental agencies, County and municipal governments, and environmental firms on contract by private solid waste companies. Samples were collected from 13 landfills equipped with sump pumps that were part of leachate-collection systems and 6 landfills equipped with gravity-fed leachate-collection systems with access to the leachate stream by a manhole (Figure 2). For 11 of the 13 landfills equipped with sump pumps, the pumps were run approximately 5 minutes to remove stagnant leachate stored in the lines prior to field rinsing a pre-cleaned container at least twice before collecting samples for chemical analysis. The remaining two landfills with sump pumps were equipped with barbed spigots from which tubing was connected. Leachate for chemical analysis was acquired directly from the spigot at those landfills. For the remaining 6 landfills, leachate was collected directly from the gravity-fed leachate stream with the use of a peristaltic pump and tubing. At least one liter of leachate was pumped through new tubing as a field rinse prior to

1 sample collection. All samples were immediately chilled to 4<sup>o</sup>C after collection and shipped overnight to the participating analytical laboratories.

Additional samples were collected for determination of alkalinity (by incremental 4 titration using a TIMM 900 Titration Manager auto Titrator), ammonium concentration ( $NH<sub>4</sub>$ +) using colorimetric CHEMets kits (CHEMetrics Inc., Calverton, VA), anion concentrations (by ion exchange chromatography using a Dionex Ion Chromatograph 120), and non-volatile dissolved organic carbon (NVDOC) concentration (by high temperature combustion using a Shimadzu TOC-Vcsn Analyzer (Shimadzu Corporation). Samples also were collected for determination of cation concentrations (by inductively coupled plasma-optical emission spectroscopy using a PerkinElmer Optima 4300), trace metal concentrations (by inductively coupled plasma mass spectrometry using a PerkinElmerElan 9000) and organic acid concentrations (by ion exchange chromatography using a Dionex Ion Chromatograph 600). Samples collected for determination of alkalinity, anions, cations, and NVDOC were filtered in 14 the field through a 0.45-µm filter. Cation samples were field acidified to a pH of 2 with HNO<sub>3</sub> 15 and NVDOC samples were field acidified with  $40\%$  H<sub>3</sub>PO<sub>4</sub> to a pH of  $2^{21}$ 

To determine concentrations of 202 CECs in leachate samples, four analytical methods were used:

**(1) LC/MS/MS Pharmaceutical Method.** This method was used to determine concentrations of 95 pharmaceuticals (includes both prescription and nonprescription) and 10 pharmaceutical degradates (Tables 1 and 2S) by analysis of a 100-µL aliquot from 1 mL of a filtered leachate sample. This method and associated validation results and performance characteristics are 22 described in detail elsewhere.<sup>36</sup> Upon receipt of each leachate sample at the laboratory, 10 to 30



A 100-µL aliquot of the filtered water sample was injected into a high-performance liquid chromatograph (HPLC) coupled to a triple quadrupole mass spectrometer (MS/MS) by using an electrospray ionization source operated in the positive ion mode. The 105 analyzed CECs were separated using a reversed-phase gradient of formic acid/ammonium formate-modified water and methanol. Multiple reaction-monitoring (MRM) of two fragmentations of the protonated molecular ion of each analyte to two unique product ions was used to specifically and sensitively identify each compound. The primary MRM precursor-product ion transition was quantified for each compound relative to the primary MRM precursor-product transition of the specific IDS chosen for that compound. The secondary MRM precursor-product ion transition was used to qualitatively confirm compound identity. The use of direct analysis without prior sample preconcentration and cleanup steps, combined with the separation provided by the HPLC and the selectivity and specificity of the MRM-MS/MS technique, results in reporting levels (RLs; determined in reagent water) that range between 2.2 and 198 ng/L; the median RL for all 20 pharmaceuticals using this method was 19 ng/L. The central tendency of RLs for this method, as 21 defined by the 25<sup>th</sup> and 75<sup>th</sup> percentiles of RL distribution, is between 8.9 and 57 ng/L.





**(4) GC/MS Household/Industrial Chemicals Method.** This method was used to determine concentrations of 60 household and industrial chemicals (Table 1 and Table 2S) in 1 L of filtered samples. The samples were filtered through 0.7-µm glass fiber filter (Whatman GF/F) and extracted by vacuum through disposable solid-phase extraction cartridges that contained modified polystyrene-divinylbenzene resin as the stationary phase. Cartridges were then dried with nitrogen gas, and the compounds were eluted with dichloromethane-diethyl ether (4:1). Each sample extract was diluted to a final volume of 400 µL, isotopically labeled internal standard compounds were added, and target analyte concentrations were determined by capillary column gas chromatography/mass spectrometry (GS/MS) in full-scan, electron impact mode. The target chemicals were qualitatively identified against an in-house mass spectral library of authentic standards, and compound concentrations were determined using the injection internal standard method. This method and the associated validation results and performance 15 characteristics are described in greater detail elsewhere.<sup>40</sup> The RLs for this method ranged between 20 and 4,800 ng/L. The median RL for all household and industrial chemicals was 80 17 ng/L. The central tendency of RLs for this method, as defined by the  $25<sup>th</sup>$  and  $75<sup>th</sup>$  percentiles of RL distribution, was between 40 and 320 ng/L. These RLs were based on a 1,000-mL sample volume; the complexity of leachate samples were such that samples of 100 mL or smaller were diluted to 100 mL and analyzed, with RLs adjusted upward in proportion to dilution.

**TIC Analysis.** In addition to 202 CECs, tentatively identified compounds (TICs) in the landfill leachate sample extracts were identified in analytical chromatograms using a National Institute of Standards and Technology (NIST) peak library (NIST05a mass spectral reference library)



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- blanks. ISDs and surrogates were added to all leachate and quality-control samples analyzed.





#### **Statistical Methods.**

Statistical methods were used to test for significant differences in distributions of frequency of CEC detections and total measured CEC concentrations with respect to waste composition (proportion of wastewater sludge, municipal waste, and industrial waste) deposited in the 19 landfills. To test these relations, analysis of variance using the nonparametric Kruskal-17 Wallis Rank-sum Test was used.<sup>41</sup> The Kruskal-Wallis Rank Sum Test also was used to test for significant differences in the distribution of frequency of CEC detection and total measured CEC concentrations with respect to landfill characteristics such as geographic location, ages of waste, waste loads, leachate production, and precipitation. The null hypothesis of these relations was 21 that frequency of CEC detections or total measured CEC concentrations for landfill groups would not differ in distribution with respect to the categorized landfill characteristics. A p-value of <0.10 was used to determine statistical significance, due to the small number of landfill sites, to indicate that the null hypothesis should be rejected, based on a significance level of 10%

 $1 - (\alpha = 0.10)$ . Rejection of the null hypothesis based on a test statistic for an individual landfill characteristic indicates that the landfill characteristic was significantly related to frequency of CEC detections and total measured CEC concentrations for the grouped landfills and that distribution in samples was significantly different with regard to a landfill characteristic.

#### **RESULTS AND DISCUSSION**

A total of 129 out of 202 analyzed CECs were detected in one or more leachate samples 8 collected during this study (Table 1). Detected CECs included 62 prescription pharmaceuticals, 23 industrial chemicals, 18 nonprescription pharmaceuticals, 16 household chemicals (included two pesticides), 6 steroid hormones, and 4 plant/animal sterols. CECs were detected in every leachate sample, with the total number of CECs in a single leachate sample ranging from 6 to 82 (median number of CECs = 31; Figure 4A). From the total 3,838 chemical measurements, the total number of detections included 231 prescription pharmaceuticals, 124 industrial chemicals, 114 household chemicals, 113 nonprescription pharmaceuticals, 32 plant/animal sterols, and 31 steroid hormone chemicals. Proportions of total measured concentration in this study refer to the sum of individual chemicals for a given chemical group divided by sum of all chemical concentrations measured. Although prescription pharmaceuticals were the most frequently detected chemical group (accounting for 35% of total measured detections), they only accounted for 1% of the total

measured CEC concentration (Figure 5). Household and industrial chemicals combined

accounted for 37% of the total detections, had the highest concentrations, and contributed to

more than 82% of the total measured CEC concentration, primarily dominated by *para*-cresol,

BPA, and phenol concentrations (Table 1). Nonprescription pharmaceuticals were detected at 2 similar frequencies as the household and industrial chemicals but accounted for just 12% of the total measured CEC concentration (Figure 5). Plant/animal sterols and the steroid hormone 4 chemicals were the least frequently detected chemicals and only accounted for 4% and  $\leq$ 1% of the total measured CEC concentrations. Summaries of total measured concentrations by chemical group show general concentration patterns, but the concentrations do not take into account variations in potency and bioactivity among individual CECs. **Figure 4. Near here….. Figure 5. Near here…..**  Twenty-one CECs including 5 household chemicals, 5 industrial chemicals, 4 nonprescription pharmaceuticals, 4 prescription pharmaceuticals, 2 plant/animal sterols, and 1 steroid hormone were measured in 50% or more of leachate samples (Figure 6, Table 1). BPA (a component of plastics), DEET (insect repellant), and cotinine (nicotine degradate) were the most frequently detected chemicals, being measured in 95% of the leachate samples. BPA, DEET, and nicotine are widely used chemicals in household/industrial products. The high frequency of detection of those compounds is consistent with results described in other studies of landfill 17 leachates, as many discarded household/industrial products end up in landfills.<sup>26-33</sup> The topical anesthetics lidocaine and camphor also were frequently detected in leachate samples in 89 and 84% of samples, respectively. Lidocaine is a medication used to relieve pain and itching and is often applied as a patch to the skin. Camphor is a natural product found in certain trees and plants that is also used as a fragrance, flavoring, plasticizer, anesthetic, and topical ointment applied to the skin to relieve pain and reduce itching. Both lidocaine and camphor have been Found in other studies to be part of the municipal waste stream.<sup>42</sup>



CEC concentrations ranged over six orders of magnitude, from (ng/L to mg/L) in leachate samples. There were 645 measurements that had concentrations of >1 ng/L, 545 of > 100 ng/L, 390 of > 1,000 ng/L (1 µg/L), 197 of > 10,000 ng/L, 62 of > 100,000 ng/L, and 9 concentrations 5 of  $> 1,000,000$  ng/L (1 mg/L). Household and industrial chemicals were measured in the highest concentrations followed by nonprescription pharmaceuticals, plant/animal sterols, prescription pharmaceuticals, and steroid hormone chemicals (Figure 6).

8 Household and industrial chemicals with maximum concentrations of  $> 1,000,000$  ng/L range included *para*-cresol (7,020,000 ng/L); BPA (6,384,000 ng/L); and phenol (1,550,000 ng/L) (Figure 6, Table 1). Measurement of *para*-cresol and BPA in landfill leachates at the µg/L 11 and mg/L concentration range has been reported in previous studies.<sup>24,27,33</sup> Combined concentrations of *para*-cresol, BPA, and phenol accounted for 70% of the total measured CEC concentrations in leachate samples collected from the 19 landfills, with samlpes from 5 landfill sites (LF 8, LF 2, LF 3, LF 10, and LF 15) accounting for 83% of the total measured CEC concentrations (Figure 4B).

Concentrations of nonprescription pharmaceuticals and the plant/animal sterols commonly were in the µg/L range and included maximum concentrations for the following frequently detected (>50%) chemicals: ibuprofen (705,000 ng/L), acetaminophen (333,000 19 ng/L), lidocaine (147,000 ng/L), and pseudoephedrine (44,100 ng/L). The frequently detected plant/animal sterols, cholesterol and 3-*beta*-coprostanol, were measured in concentrations as large as 23,400 and 834,000 ng/L, respectively. Prescription pharmaceuticals generally were measured in smaller concentrations than the nonprescription pharmaceuticals. Concentrations for the frequently detected prescription pharmaceuticals, amphetamine, carbamazepine,

carisoprodol, and tramadol, were generally in the 100's to 1,000's ng/L (Table 1, Figure 6).

2 Estrone was the only steroid hormone chemical detected in 50% or more of samples, with

concentrations in the 1's to 100's ng/L (Figure 6).

#### **Geochemistry**

The pH of leachate samples were near neutral, ranging from 6.0-7.6 (Table 1S). In 6 general, chloride (Cl<sup>-</sup>) and sulfate  $(SO_4^2)$  were the most abundant anions; Cl<sup>-</sup> concentrations 7 ranged from 167 mg/L to 3,040 mg/L and  $SO_4^2$  concentrations ranged from 0.39 mg/L to 3,430 mg/L. Bromide (Br) concentrations were relatively small in leachate samples, although Br concentrations in excess of 20 mg/L were measured in leachate samples from four landfills. NVDOC concentrations varied greatly, from 13.0 mg/L to 6,110 mg/L. Four landfills produced leachate with NVDOC concentrations greater than 1,000 mg/L; these landfills were from geographic regions with greater annual precipitation (>50 centimeters annually) (Table 1S). Several leachates with the highest NVDOC concentrations also contained relatively large Br- concentrations (Table 1S). Sodium was the most abundant cation in all leachates, with a maximum concentration of 1,890 mg/L. Samples from the group of landfills producing leachate with the highest concentrations of NVDOC also had the greatest frequency of detectable CECs and highest CEC concentrations, which may be related to more concentrated leachate or enhanced transport of CECs due to the nature of the dissolved organic matter, although the mechanisms controlling aqueous transport of polar pharmaceuticals are complex and not well 20 understood.<sup>43-45</sup> Metals measured in concentrations greater than 50  $\mu$ g/L included: Fe, Li, Al, V, Cr, Mn, Co, Ni, Cu, Rb, Zn, Sn, As, and Se. The leachates with the highest NVDOC concentrations generally contained the highest metals concentrations (excluding Ba, Mn, Sr, and

1 Li), possibly indicating the importance of organic complexation for increasing metals 2 concentrations in leachate (Table 1S).

3 Relations between geochemical concentrations and frequency of CEC detections were 4 evaluated using linear regression analysis. Only nine chemical constituents had significant ( $p =$ 5 <0.05) positive correlations based a significance level of 5% ( $\alpha$ =0.05) with frequency of CEC 6 detections. The frequency of detection of CECs in leachate samples increased as concentrations 7 of NVDOC and inorganic analytes (Br, B, K, Si, Co, Cr, and V) increased. Dissolved organic 8 carbon has been shown to decrease sorption of steroidal hormones to soil solids<sup>46</sup> and was 9 proposed as a possible facilitator of transport of steroidal hormones in groundwater affected by 10 dairy waste lagoons.<sup>47</sup> The importance of dissolved organic matter fractions from sewage sludge have been demonstrated for transport of carbamazepine in soils.<sup>48</sup> Understanding of the effect of 12 dissolved organic matter on the transport of metals and other inorganic ionic species is well-13 developed and modeled.<sup>49</sup> These observations indicate that the associations between frequency of 14 CEC detections, NVDOC, and inorganic analytes may be related to the role of NVDOC in 15 solubilizing organic CECs and these inorganic analytes in leachate.

#### 16 **Table 4. Near here…..**

#### 17 **Potential Relations with Landfill Characteristics**

Previous studies have shown that landfill characteristics such as waste composition, ages of waste, precipitation, and landfilling technology can substantially affect leachate composition (*e.g.* concentrations of dissolved organic carbon, major ions, metals, and organic compounds) at 21 Iandfills.<sup>19,24,35</sup> Discarding wastewater sludge in landfills helps to solve an important disposal need for wastewater treatment plants (WWTPs) and has been shown to reduce chemical oxygen 23 demand in leachates,<sup>50</sup> but may increase concentrations of pharmaceuticals in leachates. Recent

guidance for disposal of unused pharmaceuticals recommends mixing with kitty litter or coffee a grounds and discarding them in household trash.<sup>51</sup> Chemicals used in household and personal care products also are commonly disposed in this manner. Therefore, the composition of CECs in leachate may be affected by the types of waste they receive. For example, landfills that accept only municipal waste (household trash) or large proportions of municipal waste may produce leachate with greater frequency of detection and concentrations of pharmaceuticals or household 7 chemicals than landfills that accept larger amounts of industrial waste.<sup>51</sup>

**Landfill Waste Composition.** The relation between waste composition and CEC detections and total measured CEC concentrations by chemical groups (pharmaceuticals, household chemicals, and industrial chemicals) was determined by grouping landfill sites into multiple waste-composition categories determined from proportions of wastewater sludge, municipal waste, and industrial waste composition categories (Table 9S). The landfills were grouped by: (1a) landfills that did not accept wastewater sludge, (1b) landfills that accepted wastewater sludge; (2a) landfills that contained <70% municipal waste, (2b) landfills that contained 70–80% municipal waste, (2c) landfills that contained >80% municipal waste; (3a) landfills that did not accept industrial waste, and (3b) landfills that accepted industrial waste. Analysis of waste composition showed no significant difference in the distribution of frequency of detection and total measured pharmaceutical concentrations between leachate from landfills that: (1) accepted wastewater sludge and landfills that did not accept wastewater sludge; (2) accepted mixed proportions of municipal waste; and (3) accepted industrial waste and those that did not accept industrial waste (Table 9S).

There was however, a significant difference in the distribution of frequency of detection 23 ( $p = 0.092$ ) and total measured concentration ( $p = 0.087$ ) for household chemicals in leachate

between landfills accepting mixed proportions of municipal and industrial waste (Table 9S). The median number of detections and total measured concentrations for household chemicals was greater in landfills that accepted between 70–80% municipal waste than from landfills that accepted more homogenous mixtures of municipal waste (Table 9S). There also was a significant 5 difference  $(p = 0.10)$  in the distribution of total measured concentrations but not detections of household chemicals between landfills that accepted industrial waste and landfills that did not accept industrial waste. The median total measured concentrations of household chemicals was more than two times greater in leachate from landfills that accepted industrial waste than landfills that did not accept industrial waste. Comparison of CEC detections and total measured CEC concentrations of industrial chemicals in leachate indicated no significant difference between landfills that accepted industrial waste and landfills that did not accept industrial waste (Table 9S).

Results from the analysis of waste composition indicate that: (1) addition of wastewater sludge at levels of 10% or less did not significantly affect the frequency of detection and total measured concentration for pharmaceuticals in landfill leachate, and (2) leachate from landfills that received a heterogeneous mixture of 70–80% municipal waste tended to have greater frequency of detection and total measured concentration for household chemicals. The lack of statistical difference for distributions in CEC detections and total measured concentrations of pharmaceuticals and industrial chemicals between landfills grouped by waste composition may be related to site-specific variability of waste received, small sample size, analyzed CECs (Table 1 and Table 2S), or other landfill characteristics that promote leachate generation. **Region.** Total measured CEC detections and concentrations varied by regions of the United

States (as defined by the U.S. Census Bureau) (Figure 1). The seven sampled landfills in the

Pacific West and Northeast regions produced leachate with the greatest number of CEC detections and total measured CEC concentrations. Combined, more than half of the total detections and 66% of the total measured concentrations were in leachate samples collected at landfills in the Pacific West and Northeast regions (Table 4). Nine landfills in the Midwest, Central Southwest, and Mountain West produced leachate that contained only 36% of the CEC detections and 10% of the total measured CEC concentrations. Comparison of regional rankings by CEC detections and total measured CEC concentrations were similar in that leachate samples from the Pacific West and the Northeast regions contained the greatest frequency of detections and total measured concentrations. Leachate collected from landfills in the Mountain West region had the fewest CEC detections and the smallest total measured CEC concentrations. The Central Southwest ranked  $3<sup>rd</sup>$  for CEC 12 detections but ranked  $5<sup>th</sup>$  for total measured CEC concentrations, whereas the Southeast ranked  $5<sup>th</sup>$  for detections and  $3<sup>rd</sup>$  for total measured concentrations. The Midwest region ranked 4<sup>th</sup> for 14 detections and 4<sup>th</sup> for total measured concentrations. Due to the small number of landfills in the

six regions, data from samples from within regions were not evaluated for significant differences in distribution of CEC detections and total measured CEC concentration.

**Age of receiving waste.** Landfills were grouped into age-of-waste categories based on 18 similar age groupings used in a previous study of CECs in landfill leachate<sup>30</sup> and transitional 19 stages of landfill evolution.<sup>21</sup> Three age-of-waste categories were used: landfills containing waste of 'young' age (4 to 11 years), 'moderate' age (11 to 20 years), and 'old' age (>20 years). Landfill sites containing waste of moderate age produced the greatest frequency of detection of CECs and total measured CEC concentrations (Table 4). Landfill sites containing young and old waste had similar frequencies of detection of CECs, but landfills containing young waste



**Figure 7. Near here….** 

**Leachate production.** Landfills were grouped into three annual leachate-production categories based on the range of reported annual leachate produced, landfills producing: (1) small amounts of leachate (<5 million gallons), (2) moderate amounts of leachate (5 to 12 million gallons), or (3) large amounts of leachate (>12 million gallons). Landfills that produced moderate and large quantities of leachate annually produced leachate with similar frequencies of detection of CECs, but landfills that produced large quantities of leachate produced leachate with substantially greater total measured CEC concentrations compared to landfills that produced moderate quantities of leachate (Figure 7 and Table 4). Landfills that produced small quantities (<5 million gallons) of leachate produced leachate with the smallest frequencies of detection of CECs and the smallest total measured CEC concentrations. Even though there were substantial differences of total measured CEC concentrations between these landfill groups, the Kruskal Wallis Rank Sum test indicated no significant differences in the distributions for frequency of detection and total measured concentrations in leachate from landfills that produced differing amounts of leachate. **Precipitation.** Four of 19 landfills that produced the largest amount of leachate were in

areas that received >100 centimeters (cm) of precipitation annually and two landfills that produced the smallest amount of leachate were in areas that received <50 cm of precipitation. The two landfills that produced the smallest amounts of leachate (LF 16 and LF 19) received the largest waste load, indicating that waste load is not the sole factor in leachate production in landfills (Figure 3). Landfills were grouped into three categories based on the amount of annual 22 precipitation received: dry  $(< 50 \text{ cm})$ , moderately wet  $(50 \text{ to } 100 \text{ cm})$ , and wet  $(> 100 \text{ cm})$ . Landfills in wet environments produced leachate with substantially greater frequencies of

detection of CECs and total measured CEC concentrations compared to landfills in drier environments (Figure 7 and Table 4). Total measured CEC concentrations also were greater in wet environments than in drier environments for individual chemical groups (Figure 8). There 4 was a significant difference ( $p = 0.079$ ) in the distribution in frequency of detection and total measured concentrations for pharmaceuticals between landfills located in dry, moderately wet, and wet environments (Table 5). Previous studies have shown that the amount of precipitation a 7 landfill receives is an important component of water input and leachate composition.<sup>21,51,52</sup> Results from this study indicate that precipitation is an important factor in distribution of pharmaceuticals, and landfills located in areas receiving greater amounts of precipitation are likely to produce leachate with greater frequency of detection and concentrations of pharmaceuticals. Other types of CEC groups measured in leachate were not significantly different in frequencies of detections or total concentrations with differences in precipitation, perhaps related to slightly weaker trends from the small sample size (N=19) made even smaller when divided into subgroups and/or because of the characteristics of these groups of chemicals. The PRISM (Precipitation-elevation Regressions on Independent Slopes Model) grid of average 16 annual precipitation was the source of precipitation data used for landfill sites.<sup>53</sup>

- **Figure 8. Near here….**
- 

#### **Observations Between Landfill Characteristics**

Patterns of distribution of CECs varied according to waste composition, geographical location, ages of waste, waste loads, leachate production, and annual precipitation, but other patterns were observed between some landfill characteristics. Landfills that accepted large amounts of waste tended to be younger (mean maximum age of waste 10.8 years) compared to

landfills that accepted moderate and small amounts of waste (mean maximum age of waste 21.3 and 31.3 years, respectively), supporting the contention that landfill sizes have increased over time (Table 4). Landfills that produced large quantities of leachate tended to be older (mean maximum age of waste of 26 years) compared to younger landfills (mean maximum age of waste 5 of 10 years). Landfills that produced the largest quantities of leachate (>12 million gallons annually) were in areas that received the most precipitation (mean-annual precipitation >100 cm) 7 compared to landfills producing moderate and small quantities of leachate ( $\leq 6$  million gallons annually) that were in areas that received less precipitation. Landfills located in dry environments tended to be large in terms of amount of waste load, receiving about 3 times as much waste as landfills located in moderately wet and wet environments (Table 4). Four of the five landfills that recycled leachate were in dry environments (<50 cm of precipitation annually), whereas the 13 landfills that disposed of leachate to WWTPs were in moderately wet and wet environments (Table 1S).

#### **Identification of Organic Chemicals through Tentatively Identifiable Compound Analysis**

A total of 85 TICs were identified in one or more of the 19 leachate samples by mass spectral matching to a NIST library. Because authentic standards were not used to generate calibration curves for these chemicals, no attempt was made to estimate concentrations and only the presence or absence of each compound is reported. TICs were detected in every leachate sample, with the total number of TICs in each sample ranging from 4 to 18 (median = 11). Some TICs were detected in multiple landfills although most were detected infrequently (Table 10S). Fifteen TICs were detected in leachate from four or more landfills, 12 TICs were detected in leachate from three landfills, 17 TICs were detected in leachate from two landfills, and 41 TICs were detected in leachate from one landfill.

The most commonly detected TICs included PAH derivatives, aromatic hydrocarbons, alkanes, amides, and carboxylic acids. However, most of those compounds were detected relatively infrequently and probably due to the variability of the waste composition and chemical conditions in a given landfill. Moreover, the data generated from analysis of TICs was not sufficiently detailed to draw conclusions about relations of TICs with waste compositions and chemical composition of sampled landfills. Some of these chemicals may have been degradation products of compounds that were present in waste delivered to the landfill formed through chemical or biological processes. The leachates were complex mixtures similar to those reported 9 by other studies of classes of xenobiotic organic compounds.<sup>18-29</sup> Many other organic compounds beyond those analyzed for this paper may have been present in the leachate samples collected for this study. These were not detected because the full scan GC/MS analysis was not sensitive enough to detect them or they were not amenable to GC/MS analysis.

#### **CONCLUSIONS**

Landfills are the final repository for heterogeneous mixtures of waste from residential, industrial, and commercial sources. Therefore, landfills have the potential to produce leachate containing complex mixtures of CECs found in a variety of consumer products. Our study supports this assumption as fresh leachate collected from 19 landfills contained 129 of the 202 analyzed CECs. Fresh landfill leachate was found to contain complex mixtures of CECs that 19 include household and industrial chemicals  $(\sim 1,000-1,000,000 \text{ ng/L})$ , 20 prescription/nonprescription pharmaceuticals and plant/animal sterols  $(\sim 100-10,000 \text{ ng/L})$ , and 21 steroid hormones  $(\sim]$  -100 ng/L). Leachate from landfills that received heterogeneous mixtures

- of municipal and industrial waste tended to have greater frequency of detection and total
- measured concentration for household chemicals than landfills containing more homogeneous



The primary leachate disposal mechanism for landfills in wet environments was discharge to WWTPs. In contrast, landfills in dry environments recycled or retained leachate on-

site. Analysis of fresh leachate is an important first step in understanding landfills as a source of CECs, but may not necessarily be representative of CEC concentrations in leachate discharged to areas surrounding landfills. Additional research is needed regarding the frequency of detection and concentration of CECs in final leachate effluent that has been stored in tanks, lagoons, or treated on-site and discharged to pathways that lead offsite (e.g. receiving waters such as WWTP, streams, and groundwater). Such research would provide information that could be used to evaluate risk and provide better understanding of the fate of CECs in leachate, and may lead to changes in treatment methods, regulations for disposal of unwanted/unused pharmaceuticals, landfill setting considerations, and better knowledge of potential ecological effects posed by landfill leachate. ACKNOWLEDGEMENTS Access to landfill sites and sampling was done through a collaborative effort that included private solid-waste companies, State environmental agencies, and County and municipal governments. The review comments of Robert Eganhouse, Don Tillet, Stan Paxton, and William Andrews greatly improved this manuscript. This project was supported by the USGS Toxics Substances Hydrology Program and USGS Oklahoma Water Science Center. Special thanks to Kevin Smith for helping with preparation of sampling equipment and supplies. Any use of trade, product, or firm names in this publication is for descriptive purposes only and does not imply endorsement by the U.S. government.

### **REFERENCES**













Figure 1. Map showing states and regions of U.S. in which leachate was sampled from 19 landfills in 2011. 215x166mm (300 x 300 DPI)



Figure 3. Annual leachate production (gallons) and annual waste load (tons) estimates shown with average annual precipitation (centimeters) amounts for landfill locations, sorted from greatest to least annual leachate production. 144x127mm (300 x 300 DPI)



Figure 4. Number of detected contaminants of emerging concern (CECs) in leachate (A) and percent of total measured CEC concentrations in leachate (B), sorted from greatest to least number of detections. Frequency of detection and total concentration does not include tentatively identified compounds (TICs). 147x126mm (300 x 300 DPI)



Figure 5. Frequency of contaminants of emerging concern (CEC) detection by chemical group (red bars) and the percent of total measured CEC concentrations by chemical group (blue bars). 90x46mm (300 x 300 DPI)



Figure 6. Distribution of concentrations for contaminants of emerging concern (CECs) for chemicals detected in 50% or more leachate samples from the 19 landfills. 205x261mm (300 x 300 DPI)



## **Ranges for Selected Landfill Characteristics**

Figure 7. Percent of total measured contaminants of emerging concern (CEC) concentrations for subcategories of annual waste load (million tons), annual leachate production (million gallons), and annual precipitation (centimeters). 98x107mm (300 x 300 DPI)



Figure 8. Distribution of total measured contaminants of emerging concern (CEC) concentrations by chemical group for samples collected from landfills located in dry (< 50 centimeters annually), moderately wet (≥50 to ≤100 centimeters annually), and wet areas (>100 centimeters annually). 112x100mm (300 x 300 DPI)



Figure 8. Distribution of total measured contaminants of emerging concern (CEC) concentrations by chemical group for samples collected from landfills located in dry (< 50 centimeters annually), moderately wet (≥50 to ≤100 centimeters annually), and wet areas (>100 centimeters annually). 112x100mm (300 x 300 DPI)

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**Table 1.** Summary of analytical results for 129 detected contaminants of emerging concern (CECs) out of 202 CECs analyzed in samples from 19 landfills, 2011.



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**Table 1.** Summary of analytical results for 129 detected contaminants of emerging concern (CECs) out of 202 CECs analyzed in samples from 19 landfills, 2011.



 $a$  Method: (1) = LC/MS/MS pharmaceutical, (2) = GC/MS pharmaceutical, (3) = GC/MS/MS steroid hormones, (4) = GC/MS household/industrial chemicals,  $b$  chemical abstracting service report number,  $c$  reporting limit (RL),  $d$  concentration: E = flagged due to concentration being less than the reporting limit or greater than highest point on calibration curve,  $v =$ flagged if compound was detected in laboratory blanks between 3 and 10 times the blank concentration

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**Table 2.** Relative percent differences (RPD) between replicate samples analyzed for pharmaceuticals, steroid hormone chemicals, and household/industrial chemicals.



*a* RPD =  $|A-B| / ((A+B)/2)^*100$ 

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PP9616749f18quency of CEC detection, total measured CEC concentration, difference of gecesses & Impacts up to stamples grouped by region, age of waste, waste load, leachate production, and precipitation.



" Contaminants of Emerging Concern, <sup>b</sup> million tons, c million gallons, <sup>d</sup> minimum, e maximum, f the PRISM (Precipitation-elevation Regressions on Independent Slopes Model) grid of average annual precipitation was the so data used for landfill sites.<sup>53</sup>

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**Table 5.** Effect of precipitation on CEC dections and total measured CEC concentrations for nonprescription and prescription pharmaceuticals, household, and industrial chemicals.



<sup>*a*</sup> Contaminants of Emerging Concern, <sup>*b*</sup> minimum, <sup>*c*</sup> maximum, <sup>*d*</sup> p-value of <0.10 was used to indicate a signifcant difference between one or more of the sample distributions