

Open access • Journal Article • DOI:10.1021/ES3045289

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Quantification of Triclosan, Chlorinated Triclosan Derivatives, and their Dioxin Photoproducts in Lacustrine Sediment Cores

A THESIS SUBMITTED TO THE FACULTY OF THE GRADUATE SCHOOL OF THE UNIVERSITY OF MINNESOTA BY

Cale Thomas Anger

IN PARTIAL FULFILLMENT OF THE REQUIREMENTS FOR THE DEGREE OF MASTER OF SCIENCE

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September 2012

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ACKNOWLEDGMENTS

I would first like to offer my utmost gratitude to my advisor, Bill Arnold. Bill has been an excellent mentor during my tenure in the Department of Civil Engineering. It has been an honor to work in his lab. His dedication, support, and enthusiasm for research have been an inspiration.

I also wish to thank the members of the environmental research group for their helpful comments and assistance during the last two years. Special thanks are due to David Tan and Pat McNamara for help with troubleshooting the LC/MS and providing valuable insight into method development. Additional thanks are due to Noah Hensley and Hao Pang for many productive discussions on triclosan analysis and their help with collecting sediment cores.

Thank you to my committee members, Dan Engstrom and Paul Capel, for their critical review of this thesis. Special thanks are due to Dan for his dedication to the project, help with fieldwork, and knowledge on sediment coring and dating.

This research would not have been possible without the cooperation of numerous scientific organizations around Minnesota. First, I would like to thank the St. Croix Watershed Research Station (SCWRS) for their assistance with radiometric dating and freeze drying. Special thanks to Dylan Blumentritt and Jill Coleman-Wasik for their efforts in the radiometric dating process. The Limnological Research Center (LRC) also deserves recognition for allowing us to use their facilities, providing field supplies, and assisting with the magnetic susceptibility measurements. Additionally, I would like to thank the Large Lakes Observatory (LLO) and the crew of the RV Blue Heron for their hospitality during the collection of the Duluth Harbor and Lake Superior cores.

Thank you to Peter Villalta and Brock Matter at the Masonic Cancer Research Center Mass Spectrometry lab for their help with maintaining and troubleshooting the LC-MS-Q³. Both Pete and Brock were great resources when working through sample analysis issues.

I would like to acknowledge the Minnesota Natural Resources Trust Fund, the Department of Civil Engineering, and AMEC for funding this project and my graduate education. This project would not have been possible without their support.

Finally, I would like to thank friends and family that have supported me through my educational journey. Thank you to the Astashinskys, Mikhelsons, and the Westerbergs for always being there for me. Last, but certainly not least, I would like to thank my wife Yelena for her love, patience, and support throughout life.

ABSTRACT

Triclosan (TCS) is an antimicrobial agent commonly detected in wastewater effluent. During water and wastewater disinfection with free chlorine, TCS can be transformed to a series of chlorinated triclosan derivatives (CTDs). When discharged into surface waters, TCS and CTDs may be photochemically transformed via a cyclization reaction to a series of polychlorinated dibenzo-*p*-dioxins (PCDDs). Because PCDDs are a class of compounds known to be toxic and carcinogenic in the environment, tracking their formation from TCS and CTDs in aquatic ecosystems is necessary.

To evaluate the historical exposure of surface waters to TCS, CTDs, and their derived PCDDs, sediment cores were collected from an array of wastewater-impacted Minnesota lakes. Following radiometric dating, TCS and CTDs were extracted from core sections and quantified by liquid chromatography-tandem mass spectrometry (LC-MS-Q³). PCDDs were extracted from the same core sections and quantified by high resolution gas chromatography-high resolution mass spectrometry (HRGC-HRMS).

The concentrations and temporal trends of TCS, CTDs, and their PCDDs in aquatic sediments were found to be a function of historical wastewater treatment operations and lake system scale. Cores collected from large-scale riverine systems with many wastewater sources recorded increasing concentrations of TCS, CTDs, and their derived PCDDs since the patenting of TCS in 1964. In small-scale lakes with a single wastewater source, the depositional trends of these analytes were directly attributed to increased TCS use, local improvements in treatment, and changes in wastewater disinfection since the 1960s. Overall, concentrations of TCS, CTDs, and their PCDDs were higher in small-scale systems, reflecting a greater degree of wastewater impact. Although specific PCDD congeners are known photoproducts of TCS and CTDs, the same PCDDs were also found at very low concentrations in northern Minnesota lakes with little or no wastewater impact. Background levels of these PCDDs were attributed to a secondary, region-specific source in the 19th and 20th centuries.

To evaluate the potential risk that TCS and CTDs pose to aquatic environments, the contribution of their derived PCDDs to the total PCDD pool, in terms of mass and toxicity, was determined for each sediment core. In heavily impacted systems, the PCDD contribution from TCS and CTDs accounted for up to 70% of total PCDD mass and 32% of total PCDD toxicity in recent sediment. Thus, the discharge of TCS and CTDs may pose a substantial threat to wastewater-impacted lakes in Minnesota.

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Chapter 1: Introduction

1.1 Pharmaceuticals and Personal Care Products in the Aquatic Environment

Pharmaceuticals and personal care products (PPCPs) represent a diverse group of micro-pollutants that are of increasing environmental concern. Ranging from prescription drugs and hormones, to antimicrobials and fragrences, PPCPs are designed to evoke a specific biochemical response during human use [1-3]. Unfortunately, many of these compounds are incompletely metabolized and/or are directly disposed of in regular household or commercial settings [1,4,5]. When released into the environment, the physiochemical properties of PPCPs and their metabolites may contribute to an array of ecological problems, including acute and chronic toxicity [6], endocrine disruption [7], and reproductive impairment in organisms [8,9].

The primary route of entry for PPCPs into an aquatic environment is through the discharge of municipal and industrial wastewater. Traditional wastewater treatment plants (WWTPs) are designed to remove suspended solids, pathogens, and nutrients, but not low concentrations of PPCPs. Consequently, many of these micro-pollutants are present in wastewater effluent at ng/L to low μ g/L levels [8,10,11]. From 1999-2000, the U.S. Geological Survey (USGS) found that 80% of 139 U.S. waterways impacted by urbanization or agriculture contained organic wastewater contaminants, including steroids, reproductive hormones, antibiotics, and prescription drugs [8]. Due to the biological activity of these compounds at low concentrations, coupled with the possible synergistic effects of chemical mixtures on aquatic toxicity, the presence of PPCPs in surface waters is of great concern [12].

The risk that individual PPCPs pose to an aquatic ecosystem is highly dependent on their fate and transport in wastewater treatment and aquatic environments. In traditional WWTPs, PPCPs may sorb to particles during primary and secondary settling, or be biologically transformed in the activated sludge process [13]. Abiotic processes, such as hydrolysis, may also play a role in PPCP transformation in wastewater treatment [13]. Reactions with disinfectants, including free chlorine (Cl₂, NaOCl), ozone (O₃), and ultraviolet light (UV), often reduce the flux of PPCPs into surface waters [14-17]. These reactions, however, may also produce harmful disinfection byproducts (DBPs) that are more toxic than the parent pollutant [18,19].

Once PPCPs and DBPs are discharged into the environment, their behavior will be dictated by their physical and chemical properties, including water solubility, lipophilicity, hydrophobicity, vapor pressure, and overall reactivity. Persistent pollutants may sorb to sediment or bioaccumulate in aquatic organisms. More reactive compounds will degrade rapidly by a host of transformation processes, including microbial degradation, photolysis, and redox reactions. In many aquatic systems, even the most reactive PPCPs can be "pseudo-persistent", primarly due to a constant source from wastewater effluent [1]. Consequently, a combination of persistence and biological activity leads to potential risks for environmental receptors.

1.2 Outline of Thesis

The primary goal of this research is to better understand the fate of triclosan (TCS), a well known PPCP, and its byproducts in aquatic systems. TCS is a common antimicrobial agent found in personal care products that are routinely washed down the drain. Previous research has shown that TCS is transformed during water and wastewater

disinfection with free-chlorine [20-23]. Specific disinfection byproducts, known as chlorinated triclosan derivatives (CTDs), are discharged with TCS into surface waters [24, 25]. TCS and its CTDs may subsequently be photochemically transformed to specific polychlorinated dibenzo-*p*-dioxins (PCDDs), a class of compounds that are known to be toxic and carcinogenic in the environment [26].

To assess the impact of TCS, CTDs, and their derived PCDDs on aquatic ecosystems, sediment cores were collected from a series of wastewater-impacted lakes in Minnesota (MN). Analyte concentrations and temporal trends were discerned in each core to provide insight into the historical exposure of each lake to these compounds. Furthermore, the influence of wastewater treatment practices (e.g., improvements in treatment or type of disinfection) and lake system scale were probed to evaluate how they influence analyte trends. Finally, the PCDD contribution from TCS and CTDs to the total PCDD pool, in terms of mass and toxicity, was determined for each core. Through this analysis, the potential risk that TCS, CTDs, and their derived PCDDs pose to an array of MN lakes could be examined. This thesis includes:

Chapter 1: Introduction.

Chapter 2: Background and Literature Review. This chapter provides background on triclosan (TCS) as a personal care product and its occurrence in the environment. Furthermore, information is provided on the transformation of TCS during wastewater disinfection and in surface waters.

Chapter 3: Quantification of Triclosan, Chlorinated Triclosan Derivatives, and their Dioxin Photoproducts in Lacustrine Sediment Cores. This chapter presents results on the occurrence of TCS, chlorinated triclosan derivatives (CTDs), and their derived polychlorinated dibenzo-*p*-dioxins (PCDDs) in a series of wastewater-impacted Minnesota lakes.

Chapter 4: Conclusions

Chapter 2: Background and Literature Review

2.1 Triclosan as a Personal Care Product

Triclosan (5-chloro-2-(2,4-dichlorophenoxy)phenol; TCS) is a broad-spectrum antimicrobial agent that is present in a wide array of medical and personal care products (Figure 2-1). After being patented in 1964, initial use of TCS was confined to hospital and health care settings [27,28]. Since 1980, however, the compound has been integrated into common household items, including liquid hand soaps (0.1 - 0.3% w/w), bar soaps, dental care products, textiles, and plastics [29-31].



Figure 2-1: Chemical structure of TCS.

With a continually increasing demand for products with antibacterial properties, production of TCS has increased dramatically over the last 30 years. In 2001, it was estimated that 76% of liquid hand soaps and 29% of bar soaps contained TCS (45% of all soaps available to consumers) [28]. In the United States and Europe, annual use is estimated at 300 and 350 metric tons/year, respectively [32,33]. Furthermore, recent estimates suggest that TCS usage rates may increase by 5% each year in the future, leading to greater environmental exposure from this compound [11].

2.2 Detection in Wastewater and the Environment

Removal in Wastewater Treatment. Approximately 96% of TCS used in personal care products is washed down the drain, leading to concentrations on the order of 1-10 µg/L in wastewater influents [11,24,25,33-37]. Removal rates of TCS can vary, depending on the secondary unit operations employed in a wastewater plant. Facilities with activated sludge, a secondary treatment process applied in 75% of U.S. WWTPs, tend to remove TCS at high rates (90-98%) [11,24,29,32,34,35]. Field-scale wastewater mass balances have confirmed that 50-79% of this removal is due to aerobic biodegradation in the sludge matrix, while the remainder is due to sorption to biosolids [11,33]. No evidence of anaerobic biodegradation has been reported in the timescale of wastewater treatment [38,39].

TCS removal rates in other, less common secondary treatment processes are poorly characterized. McAvoy et al. [24] found that facilities with trickling filters, a secondary treatment process employed by 5% of U.S. WWTPs, tend to remove TCS at variable rates (58-86%). The lower and more variable removal rates are attributed to short hydraulic retention times in the trickling filters, providing less time for aerobic biodegradation during secondary treatment.

Detection in Wastewater Effluent and Surface Waters. Although most secondary treatment facilities can remove TCS at high rates, a significant amount persists through treatment and into surface waters. Effluent concentrations ranging from 10-600 ng/L, to as high as 2.7 μ g/L, have been reported at WWTPs in the U.S., Europe, and Australia [7,24,25,33,34,36,37,40-42]. In the U.S., recent studies have estimated annual TCS

loading to surface waters. Based on an estimated per capita usage rate of 3-5 mg/day, loading rates on the order of 10-20 metric tons/year have been determined [32,43].

Concentrations of TCS in impacted surface waters vary greatly, depending on the scale of the water body and the extent of contamination from treatment facilities. Typical surface water concentrations range from 1-200 ng/L in riverine and lacustrine systems impacted by WWTPs [8,29,33,35,37,40,44,45]. In the landmark study by Koplin et al. [8], TCS was detected in 58% of 85 impacted U.S. waterways at median and maximum concentrations of 140 ng/L and 2.3 µg/L, respectively. In arid regions, where wastewater effluent may dominate stream flows, TCS contamination may be more severe. The Metropolitan Water District of Southern California (MWDSC), for example, imports water from the Sacramento-San Joaquin River (SSJR). During the 2001 dry season (August-September), 70% of the SSJR flow was comprised of wastewater, leading to TCS concentrations as high as 734 ng/L in raw MWDSC drinking water [46].

Detection in Sediments. Due to its low solubility (10 mg/L at 20°C), low vapor pressure (4×10⁻⁶ mmHg at 20°C) and high hydrophobicity (log $K_{ow} = 4.8$, neutral pH), TCS has the potential to persist in sediments or bioaccumulate in aquatic organisms. Several studies have confirmed long-term preservation of TCS in lake and estuarine sedimentary environments. Buth et al. [47], for example, investigated two sediment cores from a depositional zone along the Mississippi River. Dated sediment cores provided evidence of TCS accumulation, reflecting consumer usage since the 1960s.

In a similar study, Cantwell et al. [31] investigated four urbanized estuaries along the U.S. Atlantic coast. Spatial and temporal trends of TCS in these cores reflect increased consumer usage and improvements in wastewater treatment processes (e.g. activated sludge treatment) over the last 40 years. Singer et al. [33] reported similar depositional trends when conducting a TCS mass balance in Lake Greifensee, Switzerland.

The temporal trends of TCS in dated sediment cores suggest that this antimicrobial is recalcitrant in anaerobic environments. Halden and Paull [32] estimated the environmental half-life of TCS in anaerobic sediments to be 540 days, based on a quantitative structure-activity relationship (QSAR) analysis. To date, no experimental evidence has been presented to confirm this estimate. A study by Miller et al. [48] found that TCS levels directly parallel heavy metal concentrations in estuarine sediment near a WWTP in Jamaica Bay, New York. Treated wastewater effluent is known to be the principal source of heavy metals to this aquatic environment. Due to the conservative nature of heavy metals, along with the analogous depositional trend of TCS, Miller et al. assert that TCS is not being degraded to an appreciable extent in these sediments.

A recent study conducted by the USGS found that TCS is ubiquitous in the freshwater bed sediments of wastewater-impacted surface waters in Minnesota [49]. TCS concentrations ranging from 0.4 to 85 ng/g were found in the top 10 cm of sediment near WWTPs. TCS concentrations in bed sediments collected from lake systems were up to three times higher than observed in rivers and streams. Furthermore, TCS was detected both upstream and downstream of WWTPs, confirming the widespread occurrence and persistence of TCS in these environments.

Detection and Toxicity in Aquatic Organisms. As a lipophilic compound, TCS has been detected in an array of aquatic organisms, including algae and aquatic plants [50,51], snails [52], fish [53-55], and even dolphins [56]. Coogan et al. [50] specifically looked at algal bioaccumulation factors (BAFs) in a wastewater-impacted stream in Texas. The study found that TCS concentrations ranged from 50-400 ng/g fresh weight in algae, yielding BAFs on the order of 1000 in this environment.

As primary producers, algae are essential components of aquatic food webs. Bioaccumulation in these organisms increases the potential for trophic transfer of TCS, potentially leading to high concentrations in aquatic consumers (e.g., fish) and humans. Adolfsson-Erici et al. [54], for example, found TCS concentrations as high as 4.4 μ g/g in wild fish bile in Sweden. The same study also collected random samples of human milk and recorded TCS concentrations up to 300 ng/g lipid weight.

In conjunction with the potential for bioaccumulation in aquatic systems, TCS can have a range of toxic effects on specific organisms. A study by Orvos et al. [53] found that typical TCS surface water concentrations have little effect on specific invertebrates and fish. For example, the chronic no-effect concentration (NOEC) of TCS on rainbow trout over a 61 day period is 34 μ g/L. Similarly, the NOEC for *Daphnia magna* over a 21 day period is 40 μ g/L. Several studies, however, have confirmed that TCS is chronically toxic to algae and macrophytes at environmentally relevant concentrations. The 96 hr chronic NOEC for a species of green algae (*Scenedesmus subspicatus*) has been estimated at 690 ng/L [53]. Furthermore, TCS concentrations ranging from 15 ng/L to 1.5 μ g/L have been shown to induce major shifts in algal community structure [51] and effect macrophyte growth [57]. These community shifts have the potential to modify the trophic structure and function of aquatic ecosystems that receive WWTP effluent.

Detection in Biosolids and Soils. The incorporation of activated sludge treatment into conventional WWTPs has greatly reduced the flux of TCS into aquatic environments. Biosolids derived from this treatment process, however, typically contain high concentrations of TCS. Even after aerobic biodegradation during wastewater treatment, TCS concentrations have been documented on the order of 1-50 μ g/g dry weight in digested sludge [11,24,34,42,58-62]. In the U.S., approximately 50% of biosolids derived from wastewater treatment are land-applied [63]. Consequently, elevated concentrations of TCS can leach into soils and impact surface waters that receive agricultural runoff.

The persistence of TCS in biosolid-amended soils is directly a function of biosolid application, soil properties, biodegradation rates, and runoff in the surrounding watershed. Amended soil concentrations on the order of 1-100 ng/g, to as high as 833 ng/g have been documented [9,62,64]. Several studies have suggested that aerobic biodegradation is the most important mechanism for TCS dissipation in soils. Half-lives from 18 to 108 days have been reported in laboratory and field-scale studies [38,61,62,65,66]. Higher rates of aerobic biodegradation have been attributed to the mode of biosolids application. Namely, application of biosolids in liquid form tends to make TCS more bioavailable in soils. Dewatered biosolids, however, tend to have limited contact with soil, minimizing TCS diffusion into the surrounding environment [67].

Although application of liquid biosolids promotes biodegradation, this also increases the mobility of TCS, promoting runoff to surrounding surface waters. A study by Lapen et al. [68] documented TCS concentrations as high as $3.7 \ \mu g/L$ in drain tile effluent, shortly after the application of liquid biosolids. A similar study reported drain tile concentrations of 200 ng/L [69]. As a result, land application of biosolids can contribute to TCS contamination in impacted surface waters, at concentrations known to effect aquatic primary producers.

2.3 Fate and Transformation in Natural and Engineered Waters

The chemical fate of TCS during wastewater treatment and in the environment has been well documented. In addition to removal via activated sludge treatment, TCS has been shown to be chemically transformed during chlorine disinfection. TCS and its chlorinated DBPs can subsequently be discharged into surface waters, where they are susceptible to photochemical transformation.

2.3.1 Wastewater Chlorination

Chlorination is the most commonly practiced mode of wastewater disinfection. Historically, the use of chlorine as a disinfectant has been favored, primarily for its low cost, effectiveness at inactivating most pathogens, and ability to control odors [70]. Today, approximately 72% of WWTPs in the U.S. continue to use forms of chlorine in wastewater treatment [71].

Chemistry of Chlorination. In modern wastewater treatment, chlorine gas $(Cl_{2(g)})$ and hypochlorite salts (e.g. NaOCl) are commonly used for disinfection [72]. $Cl_{2(g)}$ hydrolyzes rapidly when added to water to chloride (Cl⁻) and hypochlorous acid (HOCl) (Eq. 1).

$$Cl_2 + H_2O \rightleftharpoons HOCl + H^+ + Cl^-$$
(1)

HOCl in (Eq. 1) will simultaneously dissociate in aqueous solution to form an equilibrium solution of HOCl and hypochlorite (OCl⁻) (Eq. 2).

HOC1
$$\rightleftharpoons$$
 H⁺ + OCl⁻ pK_a (25 °C) = 7.54 (2)

Together, Cl_2 , HOCl, and OCl⁻ are known as free chlorine. At environmentally relevant pH values, HOCl and OCl⁻ represent the main free chlorine species in solution available for reaction [14].

In non-nitrified wastewater, ammonia (NH_3) concentrations are generally on the order of 10 - 40 mg/L. At these NH₃ levels, free chlorine can react to form a series of chloramines (Eqs. 3-5):

$$HOCl + NH_3 \rightarrow NH_2Cl + H_2O$$
 (3)

$$HOC1 + NH_2C1 \rightarrow NHCl_2 + H_2O$$
(4)

$$HOC1 + NHCl_2 \rightarrow NCl_3 + H_2O$$
(5)

Mono- (NH₂Cl), di- (NHCl₂), and tri- (NCl₃) chloramines tend to be less reactive and more substrate specific, when compared to free chlorine [14].

Disinfection Byproducts. Although chlorination has been instrumental in protecting public health and aquatic ecosystems, there are disadvantages to its use. Over the last several decades, a suite of harmful DBPs have been characterized in chlorinated wastewater and drinking water. Trihalomethanes (THMs) and haloacetic acids (HAAs), for example, are DBPs formed when free chlorine non-selectively reacts with dissolved

organic matter (DOM). Several THMs and HAAs have been implicated as carcinogens to humans and wildlife, leading to their regulation by the U.S. Environmental Protection Agency (EPA) [18].

More recently, formation of DBPs from micropollutants, including PPCPs and pesticides, have been of increasing concern. Due to the diverse structural moieties of many micropollutants, DBPs can be formed through an array of chemical mechanisms. For PPCPs like TCS, a common reaction mechanism is electrophilic substitution of chlorine on its phenolic moiety [14]. This reaction has been attributed to the formation of chlorinated triclosan derivatives (CTDs) in chlorinated aqueous solutions [21].

Chlorinated Triclosan Derivatives (CTDs). Several studies have investigated the primary factors that influence CTD formation during water and wastewater chlorination [20-23]. When reacting with free chlorine, the phenolic moiety on TCS activates its *ortho* and *para* positions. Electrophilic addition of chlorine at these positions forms three CTD congeners: 4,5-dichloro-2-(2,4-dichlorophenoxy)phenol (4-CI-TCS), 5,6-dichloro-2-(2,4-dichlorophenoxy)phenol (4-CI-TCS), 5,6-dichloro-2-(2,4-dichlorophenoxy)phenol (6-CI-TCS), and 4,5,6-trichloro-2-(2,4-dichlorophenoxy)phenol (4,6-CI-TCS) (Figure 2-2). Depending on the amount of excess free chlorine present in aqueous solution, ether or ring cleavage can occur on the TCS and CTD molecules, forming toxic chlorophenols and chloroform [22,23].



Figure 2-2: Chlorination of triclosan (TCS) to form three chlorinated triclosan derivatives (CTDs): 4,5-dichloro-2-(2,4-dichlorophenoxy)phenol (4-Cl-TCS), 5,6-dichloro-2-(2,4-dichlorophenoxy)phenol (6-Cl-TCS), and 4,5,6-trichloro-2-(2,4-dichlorophenoxy)phenol (4,6-Cl-TCS) (adapted from [43]).

The electrophilic substitution of TCS during free chlorination is highly dependent on pH. Rule et al. [21] found maximum pseudo-first order rate constants for TCS substitution at neutral pH, with markedly lower rates at low and high pH. This trend in reactivity is attributed to the speciation of free chlorine and TCS in natural waters. Namely, HOCl tends to be the more reactive free chlorine species in solution. Furthermore, the phenolate (O⁻) form of TCS is better at activating sites on the aromatic ring for electrophilic substitution than its phenol (OH) conjugate [21]. Thus, with pK_a values of 7.54 and 7.6 for free chlorine (HOCl/OCl⁻) and TCS (OH/O⁻), the formation of CTDs is maximized at pH values relevant to water and wastewater treatment. Similar reactivity trends are observed in chloraminated waters. CTD formation rates, however, are 2-4 orders of magnitude slower when compared to free chlorine [73]. Recent studies have characterized the removal and generation of CTDs in wastewater treatment [24,25]. Buth et al. [25] systematically investigated the presence of TCS and CTDs in a large WWTP that disinfects with free chlorine. Low levels of CTDs were detected in wastewater influent, presumably due to reaction of TCS with residual chlorine in tap water. Due to their hydrophobic nature, the CTDs were removed at high rates during activated sludge treatment and subsequently reformed during disinfection. Up to 14% of the TCS present in the disinfection stage was degraded, yielding total CTD concentrations as high as 35 ng/L in wastewater effluent.

Few papers have been published on the environmental occurrence and toxicity of CTDs. As chlorinated phenoxy phenols, CTDs are likely to have deleterious effects on aquatic environments, including chronic toxicity and endocrine disruption. The extent of these effects, however, have yet to be quantified.

2.3.2 Aqueous Photolysis

Fundamentals of Aqueous Photolysis. The photochemical transformation of organic chemicals can occur via two processes: direct and indirect photolysis. Direct photolysis occurs when a compound, or substrate, absorbs light within the UV-visible range (290-600 nm) and is electronically excited to a higher energy state [74]. The excited substrate may then be transformed to a different chemical (Figure 2-3). During indirect photolysis, however, the substrate of interest does not absorb light. In this case, a secondary solute, known as a sensitizer, becomes excited. The sensitizer may (1) directly react with the substrate, or (2) produce a series of reactive transients that degrade the compound of interest. Dissolved organic matter (DOM), and to a lesser extent, nitrate

 (NO_3^{-}) and iron (Fe^{III}), are common sensitizers in natural waters. Once excited, a number of transients can be derived from these species, including hydroxyl radicals ([•]OH), singlet oxygen (¹O₂), oxy and peroxy radicals ([•]RO, [•]ROO), superoxide radical anions (O₂[•]) and peroxides (H₂O₂) [72].



Figure 2-3: Fundamental pathways for direct and indirect photochemical degradation.

Aqueous Photolysis of TCS and CTDs. Direct photochemical degradation has been shown to be a major removal mechanism of TCS and CTDs in aqueous environments [26,33,37,75-78]. Buth et al. [26], for example, investigated the photochemical fate of these compounds in river and pure, buffered waters. Results from the study demonstrated that the pH-dependent speciation of TCS and CTDs greatly affects their photochemistry. Phenolate forms were found to degrade 44 to 584 times faster than their phenol conjugates, primarily due to a greater spectral overlap with sunlight irradiance. With pK_a values ranging from 5.9-7.6, a substantial fraction of TCS and CTDs will be in the phenolate form in surface waters (pH 6-9), leading to a high potential for photodegradation. Moreover, experiments conducted in river water (pH 8.2) indicated that indirect photochemical reactions with reactive transients are negligible when compared to direct photolysis.

Several studies have confirmed that photolysis is an important loss mechanism for TCS at the field scale [33,37,45,79]. Tixier et al. [45] found that direct photolysis accounted for 80% of total TCS removal from Lake Greifensee during summer and early fall. Similarly, Morrall et al. [79] documented a TCS half-life of 11 hrs in a wastewater-impacted river, leading to a 78% reduction over an 8 km reach. Photolysis half-lives for TCS on the order of 1-10 days were estimated for a number of global water bodies during summer months [45]. Degradation rates, in general, are greatly dependent on a series of environmental factors, including latitude, season, time of day, and water chemistry (e.g. pH, turbidity).

Formation of Polychlorinated dibenzo-p-dioxins (PCDDs). Concern has grown about potential deleterious photoproducts derived from TCS and CTDs in the environment. As polychlorinated phenoxyphenols, these compounds are susceptible to photochemical cyclization, leading to the production of polychlorinated dibenzo-p-dioxins (PCDDs) [80-83]. A number of studies have confirmed that the direct photolysis of TCS in pure and natural water yields a specific PCDD congener: 2,8-dichlorodibenzo-p-dioxin (2,8-DCDD) [26,75-78]. Similarly, 4-CI-TCS, 6-CI-TCS, and 4,6-CI-TCS have

been show to produce 2,3,7-trichlorodibenzo-*p*-dioxin (2,3,7-TriCDD), 1,2,8-trichlordibenzo-*p*-dioxin (1,2,8-TriCDD), and 1,2,3,8-tetrachlorodibenzo-*p*-dioxin (1,2,3,8-TCDD), respectively [26] (Figure 2-4). Photochemical yields of these PCDDs from TCS and CTDs vary with experimental conditions, but generally range from 0.5-3% [26,75].



Figure 2-4: Formation of PCDDs from the direct photolysis of TCS and CTDs. (A) TCS \rightarrow 2,8-DCDD, (B) 4-Cl-TCS \rightarrow 2,3,7-TriCDD, (C) 6-Cl-TCS \rightarrow 1,2,8-TriCDD, (D) 4,6-Cl-TCS \rightarrow 1,2,3,8-TCDD.

PCDDs are among a class of 210 chlorinated tricyclic aromatic compounds that are known to be persistent in the environment. Due to their hydrophobicity and resistance to metabolism, PCDDs tend to bioaccumulate in animals and humans [84-87]. Several congeners have been found to be toxic, carcinogenic, mutagenic, and cause adverse effects on animal reproductive systems [86]. These effects are magnified with increased chlorine substitution of PCDDs in the lateral positions, where 2,3,7,8-TCDD is the most toxic congener (Figure 2-5). Although toxicity data for TCS and CTD derived PCDDs is lacking, two of the four congeners (2,3,7-TriCDD, 1,2,3,8-TCDD) have three lateral positions filled. Thus, TCS derived dioxins may pose a risk to aquatic environments.



Figure 2-5: General structure of PCDDs. Toxicity of congeners increase with chlorines substituted in the lateral positions (2, 3, 7, or 8).

Historically, the dominant source of PCDDs has been attributed to two major processes: combustion and chemical production. Municipal and industrial incineration, forest fires, and vehicle exhaust comprise the majority of the combustion source. PCDDs produced in chemical production are generally derived from the manufacturing of chlorinated organic compounds (e.g. pentachlorophenol (PCP) or herbicides). The production of paper products from chlorine bleached wood pulp and metal smelting have also been recognized as important PCDD sources [88].

Since the 1970s, PCDD emissions from incineration and chemical production sources have been reduced, primarily due to regulation by the U.S. EPA [89,90]. Recent studies, however, have illustrated that TCS and CTD derived PCDDs are accounting for an increasing fraction of total dioxin loading to the environment. A pivotal study by Buth et al. [47] recently found that up to 31% of the total PCDD loading in Lake Pepin sediment is attributed to TCS and CTDs. This substantial load of PCDDs from a single antimicrobial is a direct input into the aquatic environment. Thus, further investigation is warranted on the impacts of TCS, CTDs, and their derived PCDDs on wastewater-impacted aquatic systems.

Chapter 3: Quantification of Triclosan, Chlorinated Triclosan Derivatives, and their Dioxin Photoproducts in Lacustrine Sediment Cores.

3.1 Introduction and Objectives

Triclosan (TCS) is a common antimicrobial found in a wide array of personal care products, including liquid handsoaps, bar soaps, and plastics [29-31]. Up to 96% of TCS in these personal care products is washed down the drain during normal use, leading to concentrations on the order of 1-10 μ g/L in wastewater treatment plant (WWTP) influent [11,24,25,33-37]. Although WWTPs with conventional activated sludge treatment typically remove > 90% of TCS through sorption to biosolids or biodegradation, a significant amount persists through treatment and into surface waters [8,11]. As a lipophilic compound, TCS has been shown to bioaccumulate in organisms in natural waters [50,52,54]. Moreover, environmentally relevant TCS concentrations have the potential to induce major changes in the community structure of primary producers, and thus, influence higher trophic levels in an aquatic ecosystem [51,53,57].

During the disinfection of water and wastewater with free-chlorine, TCS can be chemically transformed to a series of chlorinated triclosan derivatives (CTDs) [20-23]. Electrophilic addition of chlorine on the *ortho* and *para* positions of the TCS phenolic moiety has been shown to yield three CTD congeners: 4,5-dichloro-2-(2,4-dichlorophenoxy)phenol (4-Cl-TCS), 5,6-dichloro-2-(2,4-dichlorophenoxy)phenol (6-Cl-TCS), and 4,5,6-trichloro-2-(2,4-dichlorophenoxy)phenol (4,6-Cl-TCS). Upon formation in wastewater, TCS and CTDs are discharged into surface waters, where they are susceptible to photochemical transformation.

Direct photolysis is the primary mechanism by which TCS and CTDs are photochemically transformed in surface waters [26,75,76]. Several studies have observed that TCS and CTDs undergo a cyclization reaction, yielding a suite of polychlorinated dibenzo-*p*-dioxins (PCDDs) in natural waters. Specifically, TCS has been documented to form a specific PCDD congener: 2,8-dichlorodibenzo-p-dioxin (2,8-DCDD) [26,75-78]. Furthermore, 4-Cl-TCS, 6-Cl-TCS, and 4,6-Cl-TCS yield 2,3,7-trichlorodibenzo-*p*dioxin (2,3,7-TriCDD), 1,2,8-trichlordibenzo-*p*-dioxin (1,2,8-TriCDD), and 1,2,3,8tetrachlorodibenzo-*p*-dioxin (1,2,3,8-TCDD), respectively [26]. Because PCDDs are a class of compounds known to be toxic and carcinogenic in the environment, their production from TCS and CTDs is of clear concern [86].

The aim of this study was to investigate the depositional trends of TCS, CTDs, and their derived PCDDs in sediment cores from an array of wastewater-impacted Minnesota (MN) lakes. Due to their hydrophobicity, these compounds tend to sorb to suspended sediment and deposit in the sedimentary record. Thus, sediment cores provide a useful proxy for evaluating the exposure of an aquatic ecosystem to TCS, CTDs, and their PCDDs over time. Cores were collected in large and small scale aquatic systems with varying degrees of wastewater impaction. Furthermore, changes in historical treatment practices at relevant WWTPs were probed to evaluate how they influenced the trends of TCS, CTDs, and their PCDDs in the sediment record. Finally, the contribution of TCS and CTD derived PCDDs to total PCDD loading and toxicity was evaluated for each lake system.
3.2 Experimental

3.2.1 Chemicals and Laboratory Procedures

Chemicals and Materials. TCS (>97%) was purchased from Sigma-Aldrich. $^{13}C_{12}$ -triclosan ($^{13}C_{12}$ -TCS; >99% chemical and isotopic purity) was obtained in methanol from Wellington Laboratories. The three chlorinated triclosan derivatives (CTDs), 4-Cl-TCS, 6-Cl-TCS, and 4,6-Cl-TCS were synthesized and purified by Matthew Grandbois, Ph.D. (see [43]). All polychlorinated dibenzo-p-dioxin (PCDD) standards were purchased by Pace Analytical Services, with the exception of select congeners. Solutions of 2,8-DCDD and 2,3,7-TriCDD (50 µg/mL) in isooctane were purchased from AccuStandard. A mixture of 1,2,3,7/1,2,3,8-TCDD (50 µg/mL) in *n*-nonane was obtained from Cambridge Isotopes, along with ¹³C₁₂-2,3-DCDD (99% chemical and isotopic purity; 50 μ g/mL) and ¹³C₁₂-2,3,7-TriCDD (99% chemical and isotopic purity; 50 μ g/mL). Sulfuric acid (H₂SO₄) and sodium hydroxide (NaOH) were purchased from BDH. Ultrapure water (18.2 M Ω -cm) was obtained from a Millipore Simplicity UV purification system. All solvents used for extractions and cleaning were of HPLC grade, with the exception of methyl-tert-butyl-ether (MTBE). MTBE was purchased from Sigma-Aldrich at >99% purity. Ottawa sand (OS) was acquired from Fisher Scientific. Ultra-high purity and industrial-grade nitrogen were purchased through Matheson.

Laboratory Cleaning Procedures. To prevent analyte cross-contamination and carryover in the laboratory, a specific set of cleaning procedures were required for glassware, tools, and instrumentation. Appendix A in this thesis provides a detailed description of these cleaning procedures for TCS and CTD analysis.

3.2.2 Sediment Core Collection

Coring Locations. To characterize the impact of TCS, CTDs, and their derived PCDDs on regional lake systems, sediment cores were collected from eight sampling sites around Minnesota (MN) (Figure 3-1). Differential global positioning system (dGPS) coordinates are provided for each site in Table 3-1.

From July to August 2010, cores were collected from two fluvial lakes: Lake Pepin and Lake St. Croix. Lake Pepin is 34 km-long wastewater-impacted depositional zone along the Mississippi River. With a watershed of over 122,000 km², Lake Pepin drains approximately two-thirds of the state of MN and integrates many wastewater sources. The Metropolitan WWTP in St. Paul, MN is a major wastewater source in this watershed, serving approximately 1.8 million people and discharging up to 251 million gallons per day (MGD) directly into the Mississippi River. Lake St. Croix is an impacted depositional zone along the St. Croix River that drains approximately 20,000 km². The major wastewater source in this system is from the St. Croix Valley WWTP in Stillwater, MN, a facility serving 30,000 people and discharging up to 4.5 MGD.

Sediment Core	GPS Coordinates			
Lake St. Croix	44° 56.858' N	92°45.325' W		
Lake Pepin	44°29.987' N	92° 17.653' W		
East Lake Gemini	45°35.445' N	94°23.493' W		
Lake Winona	45° 52.425' N	95°24.350' W		
Duluth Harbor	46°43.996' N	92°3.920' W		
Lake Superior	46°49.059' N	91°51.820' W		
Lake Shagawa	47°54.685' N	91°53.382' W		
Lake Little Wilson	47°39.421'N	91°4.227' W		

Table 3-1: Differential global positioning system (dGPS) coordinates for sediment core locations.

To assess the impact of wastewater on smaller-scale aquatic systems, two lakes were cored in central MN during September 2010. One core was collected from East Lake Gemini, a 0.1 km² artificial impoundment that receives wastewater from St. Johns University (SJU) near Collegeville, MN. The SJU WWTP serves a population of 1200 during the summer and 2600 during the academic year. Wastewater discharge correspondingly varies throughout the year, with a maximum discharge of 0.23 MGD. Lake Winona, a natural lake of similar size (0.85 km²), was cored near Alexandria, MN. The Alexandria Lakes Area Sanitation District (ALASD), which serves 23,000 people, discharges up to 3.8 MGD directly into Lake Winona.



Figure 3-1: Sediment coring sites in MN.

In October 2010, sediment cores were collected in Duluth Harbor and Lake Superior. The Duluth Harbor core was obtained in St. Louis Bay, near the outfall of the Western Lake Superior Sanitation District (WLSSD) wastewater facility. WLSSD is a large treatment plant that treats up to 40 MGD, serving approximately 130,000 people in the Duluth, MN area. Approximately half of the wastewater treated by WLSSD is derived from industrial sources. The City of Superior, with a population of 27,000, also discharges approximately 4-5 MGD of wastewater into this aquatic system. The Lake Superior core was collected 22 km northeast of St. Louis Bay on the RV Blue Heron, a research vessel affiliated with the Large Lakes Observatory.

Finally, the last two cores were collected in June 2011 on two lakes near the Boundary Waters Canoe Area (BWCA) in MN. One core was acquired at Lake Shagawa, a 9.4 km² water body that is the receiving water for the Ely, MN WWTP. The Ely WWTP discharges 1.5 MGD from a population of 3500. The last core was collected from Little Lake Wilson, a reference lake with no wastewater input and little anthropogenic contact.

Core Collection Methods. All sediment cores, with the exception of Lake Superior, were collected using a piston corer with a polycarbonate barrel operated from the water surface. The Lake Superior core was collected on the RV Blue Heron using an Ocean Instruments Multicorer. Upon collection, cores were vertically extruded and sectioned at designated intervals. Down-core smearing was removed from the outer circumference of each interval to prevent analyte drawdown in the core. Sections were transferred to baked glass jars, homogenized, and sub-sampled for radiometric dating. Jars were then covered with foil, cooled to 4°C, and frozen within 24 hours of collection.

One core (Lake Pepin) was transported to the Limnological Research Center (LRC) at the University of Minnesota for magnetic susceptibility analysis (see Section 3.2.3 – Radiometric Dating). The core was then sub-sampled using the aforementioned procedures.

Prior to radiometric dating or analyte extraction, loss-on-ignition (LOI) was preformed on each sediment core. Standard LOI techniques were applied to determine sediment dry density, water content, and weight percentage of organics and carbonate [91,92]. Briefly, core intervals were homogenized and sub-sampled into crucibles. Samples were sequentially dried for 12 hours at 105°C, 4 hours at 550°C, and 2 hours at 1000°C to determine sample water content, organic percent, and carbonate percent, respectively.

3.2.3 Radiometric Dating

Lead-210 and Cesium-137 Dating. Sediment cores were dated using lead-210 (²¹⁰Pb) and cesium-137 (¹³⁷Cs) methods, as described in recent literature [93,94]. Briefly, ²¹⁰Pb activity is measured through its granddaughter product, polonium-210 (²¹⁰Po), in 15-20 sections of each core. ²¹⁰Po is quantified using isotope-dilution methodology, with polonium-209 (²⁰⁹Po) as an internal recovery surrogate. ²¹⁰Po/²⁰⁹Po are distilled from dry sediment and analyzed via alpha spectrometry [94]. Total ²¹⁰Pb in the core profile is used to determine radiometric dates and sedimentation rates through the constant rate of supply (CRS) model [93].

To supplement ²¹⁰Pb dating, ¹³⁷Cs activity was measured in 6-10 sections in each core using gamma spectrometry. ¹³⁷Cs is a particularly useful radioactive isotope, as it is

derived from above-ground nuclear testing. A spike of ¹³⁷Cs in the sedimentary record is known to directly correlate with the peak of testing, between 1963 and 1964. Thus, provided that a sediment profile is relatively undisturbed, ¹³⁷Cs provides a useful marker for core dating [94].

Magnetic Susceptibility. Many sediment cores have been collected and radiometrically dated in Lake Pepin. Moreover, several of these cores have been analyzed for magnetic susceptibility, where stratigraphic variations in ferromagnetic mineral concentrations are measured. The current Lake Pepin core was dated by conducting whole-core magnetic susceptibility measurements using a Geotek LTD multisensory core logger in the LRC. The susceptibility profile was matched to those from recently dated cores in the same depositional area of the lake, providing a dating profile for the current core.

3.2.4 Triclosan (TCS) and Chlorinated Triclosan Derivative (CTD) Analysis

Wet samples, corresponding to 10 g dry weight from each core interval, were freeze dried for 3-5 days at the LRC or the St. Croix Watershed Research Station (SCWRS). Dried samples were subsequently transferred to 150 mL beakers. Each sample was spiked with a dilute solution of ${}^{13}C_{12}$ -TCS (500 ng in 1 mL of acetonitrile (ACN)) as an isotope-dilution surrogate for TCS and CTD recovery. For each core, a method blank (freeze dried Ottawa sand) was spiked with ${}^{13}C_{12}$ -TCS and processed to ensure there was no analyte contamination. Spiked samples were covered with foil and allowed to equilibrate for 12 hours prior to extraction.

Accelerated Solvent Extraction. Following equilibration, samples were extracted on a Dionex Model 350 accelerated solvent extractor (ASE 350) (Sunnyvale, CA, USA). Briefly, samples were transferred to 22 or 66 mL stainless steel ASE cells. To provide cleaner extracts, two glass fiber filters were placed at the base of each ASE cell. Sediment was loaded into the cell and mixed with baked Ottawa sand. Any remaining void volume was filled with baked Ottawa sand. Finally, a single glass fiber filter was placed at the top and the cell was sealed for extraction.

Table 3-2 provides optimized parameters for ${}^{13}C_{12}$ -TCS, TCS, and CTD extraction on the ASE 350. HPLC grade dichloromethane (100% DCM) was determined to be the most appropriate extraction solvent.

Optimized Value
100 °C
1500 psi
5 mins
5 mins
100%
100 s
2

Table 3-2: Optimized parameters for ${}^{13}C_{12}$ -TCS, TCS, and CTD extraction from sediment on the ASE 350.

During a typical extraction cycle, sample cells are loaded into an oven, filled with solvent, and subsequently pressurized and heated. After a designated static time, solvent is then purged from the cell with ultra-high purity nitrogen (N_2) into a 250 mL collection vessel. Two extraction cycles were performed to (1) maximize ¹³C₁₂-TCS, TCS, and

CTD recovery in complex sediment matrices, and (2) minimize the presence of common interferences (e.g. NOM) in the final ASE extract.

Due to a wide range of organic content in the sediment cores collected (Table 3-3), two separate clean-up methods were developed for ASE extracts (Figure 3-2). Methods 1 and 2 were designed for cores with high (17-40%) and low (6-16%) organic content, respectively. These methods were modified from Buth [43] to accommodate the range of sediment matrices in this study.

Table 3-3: Mean organic content of sediment cores. Organic content data are derived from LOI results.

Sediment Core	Mean Organic Content (%)
Lake Superior	6
Duluth Harbor	10
Lake Pepin	12
Lake St. Croix	16
Lake Winona	17
Lake Shagawa	26
East Lake Gemini	29
Lake Little Wilson	40

Method 1 – High Organic Content. For sediments with high organic content, extracts were cleaned up on solid-phase extraction (SPE) cartridges and silica columns.

Solid-Phase Extraction. SPE cleanup was performed using Waters Oasis HLB 6 cc/200 mg cartridges. Prior to cleanup, ASE extracts were sub-sampled (1/5 aliquot by mass; ~ 15 mL by volume; ~ 2 g dry sediment weight), transferred to 17 mL glass centrifuge tubes, and blown down to dryness with industrial-grade N₂. Samples were then resuspended in 500 μ L of methanol (MeOH) and dissolved in 30 mL of pH 4 water (H₂O) adjusted with H₂SO₄/NaOH. A pH of 4 was selected to keep \geq 99% of TCS and

CTDs in their neutral, phenol form, maximizing retention on lipophilic moieties in the SPE media.

Before sample loading, SPE cartridges were preconditioned with sequential 5 mL aliquots of MTBE, MeOH, and pH 4 H₂O on a Restek 12-port vacuum manifold. Transfer lines between samples and cartridges were also washed with 10 mL aliquots of MTBE, MeOH, and pH 4 H₂O to remove potential analyte contamination prior to use. Samples were then loaded onto cartridges at 2 mL/min. After loading, cartridges were washed with three sequential 5 mL aliquots of 50:50 MeOH:H₂O (v/v) to remove polar DOM from the SPE media. Cartridges were then dried for 20-25 minutes by vacuum and eluted sequentially with 5 mL of MeOH and 5 mL of 90:10 MTBE:MeOH (v/v) into 15 mL centrifuge tubes. Final SPE extracts were blown down to ~ 500 µL with industrial-grade N₂ and prepped for silica column cleanup.

Silica Column Cleanup. Silica columns were prepared in 6 mL disposable syringes for further sample cleanup. Each column was packed with a plug of glass wool, a thin layer of baked Ottawa sand, 2 g of baked silica gel (60 – 200 μ m particle size, 60 Å pore size), and an upper layer of baked sand. Prior to sample loading, each column was flushed with 10 mL of ethyl acetate to precondition the silica gel and remove any potential analyte contamination. SPE extracts were then quantitatively loaded onto the column and eluted by gravity with 12 mL of ethyl acetate. Final extracts were blown down to dryness with industrial-grade N₂ and resuspended in 200 μ L of 50:50 ACN:H₂O (v/v). 150 μ L of the final extract was transferred via gas-tight syringe to amber autosampler vials with 200 μ L inserts. Vials were then wrapped with Teflon tape and stored at 4 °C until analysis.



Figure 3-2: Cleanup procedure for ASE extracts. Methods 1 and 2 were designed for sediments with high and low organic contents, respectively.

Method 2 – Low Organic Content. ASE extracts from sediments with low organic content were cleaned up exclusively with silica gel columns. Namely, subsamples of ASE extracts were transferred to 17 mL glass centrifuge tubes, blown down to dryness with industrial-grade N₂, and suspended in ~500 μ L of 55:45 MeOH:MTBE (v/v) (the final eluent compostion from the SPE step in Method 1). These extracts were the cleaned up on silica columns, blown down to dryness, and resuspended as in Method 1.

To assess the efficacy of Methods 1 and 2 for extracting TCS and CTDs in the given sediment matrices, a series of spike and recovery experiments were executed. Extractions were performed in triplicate on baked Ottawa sand and sediment from the base of the Lake Superior, Lake Pepin, and Lake Shagawa cores. Samples (10 g dry weight) were spiked with 500 ng ${}^{13}C_{12}$ -TCS, 100 ng TCS, and 12 ng of each CTD in a 1

mL dilute solution of ACN. Method blanks, spiked with 500 ng ${}^{13}C_{12}$ -TCS, were extracted concurrently for each matrix. The Ottawa sand, Lake Superior (low organic content), and Lake Pepin (medium organic content) were extracted with Method 2, while Lake Shagawa (high organic content) was extracted with Method 1. Spike and recovery results from each matrix are provided in Table 3-6 in Section 3.3.1.

LC-MS-Q³ Analysis. Sediment core extracts in this study were analyzed on an Agilent 1100 capillary HPLC equipped with a Finnigan TSQ Quantum Discovery MAX MS-Q³ tandem mass spectrometer. Analyte separation was achieved with 8 μ L sample injections on a Phenomenex Synergi MAX-RP reverse-phase column (150 × 0.5 mm, 4 μ m particle size, 80 Å pore size) maintained at 30 °C during sample runs. A flow rate of 10 μ L/min was established with a binary gradient of (A) 15 mM ammonium acetate buffer, and (B) 100% HPLC grade ACN. The gradient began at 50% B, ramped to 100% B over 20 mins, and then ramped down to 50% B from 20 to 23 minutes. 50% B was maintained from 23 to 35 minutes to allow for column re-equilibration. Column effluent was diverted to waste during first and last 10 minutes of each sample run to minimize contamination of the MS-Q³ ion source.

¹³C₁₂-TCS, TCS, and CTDs were quantified using selected reaction monitoring (SRM) in negative electrospray ionization (ESI⁻) mode on the MS-Q³. Specific precursor and product mass-to-charge (m/z) ratios were selected for optimal sensitivity (Table 3-4).

Analyte	Precursor Ion		Product Ion	Role of <i>m/z</i>
	(m/z)		(m/z)	
TCS	287	\rightarrow	35	quantification
	289	\rightarrow	37	confirmation
4-Cl-TCS	321	\rightarrow	35	quantification
	323	\rightarrow	37	confirmation
6-Cl-TCS	321	\rightarrow	35	quantification
	323	\rightarrow	37	confirmation
4,6-Cl-TCS	355	\rightarrow	35	quantification
	357	\rightarrow	37	confirmation
${}^{13}C_{12}$ -TCS	299	\rightarrow	35	quantification

Table 3-4: Selected reaction monitoring (SRM) transitions for ${}^{13}C_{12}$ -TCS, TCS, and CTD quantification and confirmation.

In the MS-Q³, precursor ions are selected by the first quadrupole. These precursor ions are then further fragmented, producing product ions that are detected by the third quadrupole. Buth [43] found that chloride ion (35 Cl⁻; *m/z* 35) was the dominant product ion for TCS and CTDs. Thus, SRM transitions with 35 Cl⁻ as the product ion were selected for analyte quantification. Furthermore, SRM transitions corresponding to 37 Cl⁻, the second most common isotope of Cl⁻, were monitored for analyte confirmation. Prior to running on the MS-Q³, a high concentration of 13 Cl₋2-TCS (20 mg/L) was directly infused into the ion source to tune the instrument for the analytes of interest. Furthermore, to maximize analyte response on the MS-Q³, a clean ion transfer tube was preconditioned with existing sediment extracts. The rational for this procedure is discussed in Appendix B.

Analyte Quantification and Quality Control. Calibration standards for LC-MS- Q^3 analysis were prepared in 50:50 ACN:H₂O (v/v) over a range of TCS and CTD concentrations. A constant concentration of ¹³C₁₂-TCS was maintained in each standard.

Seven to eight point calibration curves were prepared by plotting the ratio of analyte to ${}^{13}C_{12}$ -TCS peak area (y-axis) as a function of analyte concentration (x-axis). From these calibration curves, a response factor was obtained for each analyte. This response factor was then (1) multiplied by the analyte to ${}^{13}C_{12}$ -TCS peak area ratio in the sediment matrix and, (2) the amount of ${}^{13}C_{12}$ -TCS (in g) in the spiked sample. By using isotope-dilution methodology, this provides the amount of analyte (in g) in the sediment sample. Moreover, to assess the efficacy of individual extractions, the absolute ${}^{13}C_{12}$ -TCS recovery was determined for each sample. Detailed information on the calculation of analyte concentrations and ${}^{13}C_{12}$ -TCS absolute recovery is provided in Appendix C.

Several quality assurance/quality control (QA/QC) measures were followed when processing samples and running extracts on the LC-MS-Q³. As indicated in the extraction procedure, method blanks, along with spike and recovery experiments, were prepared to test method performance. Furthermore, instrument blanks of 50:50 ACN:H₂O (v/v) were run every 8 samples during LC-MS-Q³ analysis to monitor for analyte carryover between LC injections. To calculate sediment concentrations in each core, a series of criteria were established to determine the limit of quantification (LOQ) of each analyte. These criteria include: (1) the analyte signal-to-noise ratio must be > 10 within the sediment matrix, (2) the analyte concentration must be 10 times higher than that in the method and instrument blanks, and (3) analyte quantification can only occur above the lowest calibration point in its respective calibration curve.

3.2.5 Polychlorinated dibenzo-p-dioxin (PCDD) and furan (PCDF) Analysis

Between 7 and 12 g (dry weight) of each core interval was analyzed for PCDD/F congeners at Pace Analytical in Minneapolis, MN. East Lake Gemini was an exception, where between 1 and 2 g (dry weight) was processed. For all cores, samples were spiked with nineteen ${}^{13}C_{12}$ -labeled di- through octa-CDD/F isomers as isotope-dilution surrogates and analyzed using an expanded version of U.S. EPA Method 1613B [95]. Although the presence of PCDF isomers is not specifically investigated in this study, they were quantified in the sediment cores as part of Method 1613B standard procedures.

Once spiked with labeled recovery surrogates, each sample was extracted with toluene for at least 18 hours using a Soxhlet/Dean Stark apparatus. Extracts were subsequently spiked with ³⁷Cl₄-2,3,7,8-TCDD to measure the efficiency of sample cleanup. Soxhlet/Dean Stark extracts were concentrated using a Snyder column, backextracted with concentrated H₂SO₄ and NaOH, and eluted through multi-layer silica columns (2 g neutral silica, 4 g acidic silica, and 2 g basic silica) with hexane. Eluates were then added to 4 g activated aluminum oxide (Al_2O_3) columns and eluted with 60:40 DCM:hexane (v/v). After solvent exchange into hexane, Al_2O_3 column eluates were cleaned up via carbon chromatography, where samples were passed through 0.5 g of 18%activated carbon mixed with Celite. These columns were preconditioned with 5 mL of toluene, 2 mL of 75:20:5 DCM:MeOH:toluene (v/v/v), 2 mL of 50:50 DCM:cyclohexane (v/v), and 5 mL of hexane. Sample extracts were added to the column and flushed in the forward direction with 2 mL of 50:50 DCM:cyclohexane (v/v) and 2 mL of 75:20:5 DCM:MeOH:toluene (v/v/v) to remove potential interfering compounds. Finally, analytes were washed off the column in the reverse direction with 10 mL of toluene.

The toluene was then concentrated, spiked with ${}^{13}C_{12}$ -1,2,3,4-TCDD and ${}^{13}C_{12}$ -1,2,3,7,8,9-HxCDD as recovery standards, and concentrated to a final volume of 40 μ L.

HRGC-HRMS Analysis. PCDD/F analysis was performed using high-resolution gas chromatography-high-resolution mass spectrometry (HRGC-HRMS). Aliquots of final extracts (1 μ L) were injected into an HP 5890 gas chromatograph with a split/splitless injector and a 60 meter DB-5MS capillary column (0.25 mm ID × 0.25 μ m film). The gas chromatograph was coupled to a Waters Autospec Ultima high-resolution mass spectrometer operated in selected ion monitoring (SIM) mode (positive electron impact, > 10,000 resolution, 32 eV, 280 °C). Acquisition windows were set to include all tetra- through octa-CDD/F isomers. Windows for di- and tri-CDD/F isomers were centered around the di- and tri-CDD congeners in this study. Therefore, total DCDD and TriCDD values presented should be considered an estimate, as the first and last DCDD and TriCDD eluters may have been outside the established acquisition windows for these isomers.

Analyte Quantification and Quality Control. Standards for HRGC-HRMS analysis were prepared using a U.S. EPA Method 1613B calibration set (tetra- through octa-CDD/F isomers). A secondary calibration set for di- and tri-CDD/Fs was prepared at similar levels to tetra-CDD/F in the Method 1613B. From these calibration sets, five-point calibration curves were constructed for each PCDD/F congener, with constant levels of ${}^{13}C_{12}$ -PCDD/F internal standards.

Analogous to the TCS and CTD analysis method, a series of QA/QC procedures were followed during PCDD/F analysis. Method blanks were prepared to demonstrate freedom of analyte contamination during sample extraction and cleanup. Furthermore, spike and recovery experiments in clean sand and sediment were performed to validate the PCDD/F extraction method for each sediment core. Finally, all reported PCDD/F concentrations were required to be 10 times greater than levels in method blanks.

3.3 Results and Discussion

3.3.1 Analytical Method Performance

TCS and CTD Sediment Extraction. The LC-MS-Q³ method for TCS and CTD quantification provided satisfactory peak shape and effectively separated the analytes of interest in all chromatograms. Figures 3-3 and 3-4 illustrate example chromatograms from a representative standard and Lake Pepin sediment extract, respectively. In all cases, the CTDs eluted later than ¹³C₁₂-TCS and TCS due to their greater hydrophobicity. TCS and CTD calibration curves were linear, with R^2 values consistently \geq 0.99 for each analyte. Two calibration curves for 4,6-Cl-TCS had R^2 values > 0.98.

For each sediment core, trace amounts of TCS were detected in method and instrument blanks. Reported TCS sediment concentrations were at least 10 times greater than those observed in the blanks. In all cases, CTDs were not detected in procedural blanks. All reported CTD concentrations, therefore, were required to be above the lowest point in their respective calibration curves. Finally, the raw analyte signal-to-noise ratio was always > 10 for any reported TCS or CTD concentration. Using these criteria, LOQs ranging from 0.36 to 0.9 ng/g (TCS) and 60 to 70 pg/g (CTDs) were achieved.



Figure 3-3: LC-MS-Q³ chromatograms of ${}^{13}C_{12}$ -TCS, TCS, and CTD SRM transitions in a representative standard. Analyte retention times are displayed above each peak.



Figure 3-4: LC-MS-Q³ chromatograms of ${}^{13}C_{12}$ -TCS, TCS, and CTD SRM transitions in a Lake Pepin sediment extract. Analyte retention times are displayed above each peak.

Due to the wide range of sediment matrices in this study, the absolute recovery of ${}^{13}C_{12}$ -TCS varied significantly (Table 3-5). In cores with high organic content (e.g. Lake Shagawa), the complex matrix and ion-suppression during MS-Q³ analysis induced relatively low ${}^{13}C_{12}$ -TCS recoveries. Higher recoveries were observed in cores with low organic content (e.g. Duluth Harbor), where interferences were less severe.

Absolute Re	e Recovery (%)		
¹³ C ₁₂ -TCS			
27 ± 9^a	(n=12)		
33 ± 7	(n=10)		
36 ± 12	(n=9)		
37 ± 12	(n=16)		
41 ± 20	(n=12)		
61 ± 10	(n=10)		
78 ± 9	(n=16)		
87 ± 17	(n=13)		
	Absolute Re ${}^{13}C_{12}$ - 27 ± 9^a 33 ± 7 36 ± 12 37 ± 12 41 ± 20 61 ± 10 78 ± 9 87 ± 17		

Table 3-5: Absolute recovery of ${}^{13}C_{12}$ -TCS in sediment cores.

^{*a*} Reported values are the mean \pm standard deviation

Spike and recovery results for sediment extraction procedures (Methods 1 and 2) are presented in Table 3-6. Overall, results demonstrate the utility of isotope-dilution methodology for quantifying TCS and CTDs in a range of sediment matrices. Recovery of TCS, relative to the ¹³C₁₂-TCS surrogate, is consistently near 100% for all matrices. Relative CTD recovery, however, generally decreases with increased matrix complexity. This may be attributed to a number of factors, including increased matrix suppression at CTD retention times or a disproportional loss of CTDs relative to ¹³C₁₂-TCS and TCS during sample extraction and cleanup.

Table 3-6: Relative recovery of TCS and CTDs for extraction Method 1 (Lake Shagawa) and 2 (Ottawa sand, Lake Superior, Lake Pepin).

	Absolute Recovery (%)	Relative Recovery (%)			6)
Matrix	¹³ C ₁₂ -TCS	TCS	4-Cl-TCS	6-Cl-TCS	4,6-Cl-TCS
Ottawa sand (n=3)	60 ± 6^a	109 ± 2	81 ± 10	81 ± 12	68 ± 18
Lake Superior (n=3)	60 ± 2	108 ± 3	52 ± 4	52 ± 7	36 ± 3
Lake Pepin (n=3)	52 ± 3	108 ± 7	56 ± 3	59 ± 9	44 ± 4
Lake Shagawa (n=3)	38 ± 5	106 ± 3	35 ± 8	42 ± 3	15 ± 4

^{*a*} Reported values are the mean \pm standard deviation

PCDD/F Sediment Extraction. The sediment extraction and HRGC-HRMS methods employed at Pace Analytical provided consistent PCDD/F recovery in the sediment cores. The absolute recovery of nineteen ${}^{13}C_{12}$ -PCDD/F surrogates ranged from 20 – 116% in sediment extracts, all of which were within the target range specified in Method 1613B (Table 3-7). Moreover, the relative recovery of nineteen native PCDD/F isomers in sand spikes ranged from 74-129%, demonstrating the precision and accuracy of the isotope-dilution methodology.

Table 3-7: Absolute recovery range for ${}^{13}C_{12}$ -PCDD/F isomers in sediment cores.

	Absolute Recovery (%)		
Sediment Core	¹³ C ₁₂ -PCDD/F ^{<i>a</i>}		
Lake Shagawa	27 - 99		
Lake Little Wilson	20 - 112		
East Lake Gemini	44 - 98		
Lake St. Croix	30 - 107		
Lake Winona	28 - 85		
Lake Superior	47 - 116		
Lake Pepin	34 - 101		
Duluth Harbor	29 - 107		

^{*a*} 19 isotopically labeled di- through octa-CDD/F isomers

Spike and recovery experiments were performed in duplicate on all sediment matrices, with the exception of East Lake Gemini. East Lake Gemini spikes were fouled during extraction and are currently being reprocessed. Relative recovery of native PCDD/F isomers ranged from 72 – 126% with relative percent differences ranging from 0 – 14.2% for individual analytes between duplicate matrix spikes (Table 3-8). One recovery outlier was observed in the Little Lake Wilson core, where 2,8-DCDD exhibited a maximum relative recovery of 238%. The same PCDD congener, however, consistently exhibited relative recoveries ranging from 102-117% in sand spikes and 89-124% in other core matrices. Thus, it is likely that interference around the 2,8-DCDD retention time in Little Lake Wilson influenced its recovery during HRGC-HRMS analysis.

Relative Recovery (%)	
PCDD/F ^a	RPD $(\%)^{b}$
85 - 125	0.0 - 6.8
73 - 116 (238) ^{<i>c</i>}	0.0 - 14.2
-	-
94 - 124	0.4 - 12.9
72 - 124	0.4 - 10.2
76 - 120	0.3 - 17.5
86 - 119	0.2 - 7.3
85 - 126	0.2 - 9.2
	Relative Recovery (%) PCDD/F ^a 85 - 125 73 - 116 (238) ^c - 94 - 124 72 - 124 76 - 120 86 - 119 85 - 126

Table 3-8: Relative recovery range for PCDD/F isomers in sediment cores.

^{*a*} 19 native di- through octa-CDD/F isomers

^b RPD = relative percent difference between matrix spike replicates

^c value in parenthesis represents the maximum relative recovery of 1 outlier (2,8-DCDD)

A comprehensive list of isomer-specific ${}^{13}C_{12}$ -PCDD/F recoveries for each sediment core is provided in Appendix D. This appendix also provides isomer-specific PCDD/F recoveries for each matrix spike and recovery experiment.

For all cores, trace amounts of specific di- through octa-CDD/F isomers were detected in method blanks, most of which were well below calibration ranges. All reported PCDD/F concentrations were > 10 times the levels in these blanks.

3.3.2 TCS, CTDs, and their PCDDs in MN Lakes

Lake Pepin and Lake St. Croix. Sediment cores collected in large-scale, wastewater-impacted riverine systems provide important insight into the accumulation of TCS and its byproducts in aquatic environments. Figures 3-5 and 3-6 illustrate the temporal trend of TCS, CTDs, and their derived PCDDs for Lake Pepin and Lake St. Croix, respectively. In both cores, TCS concentrations increase with time, reflecting consumer usage since the mid-1960s. A similar trend in the 2,8-DCDD profile, the known photoproduct of TCS, is observed in these systems. Furthermore, concentrations of 2,3,7-TriCDD, 1,2,8-TriCDD, and 1,2,3,8-TCDD, the photoproducts of CTDs, generally parallel TCS trends in each core.

Wastewater treatment practices can have a profound impact on the presence of TCS, CTDs, and their derived PCDDs in aquatic sediments. The Metropolitan WWTP, for example, is a major wastewater source to the Mississippi River that is approximately 80 km upstream from Lake Pepin. Since its inception in 1938, the plant has disinfected its effluent with various forms of free chlorine (e.g. Cl_2 or NaOCl). The effect of

chlorine disinfection, coupled with increased TCS use over the last 40 years, is directly realized in the CTD and higher chlorinated PCDD profiles in Figure 3-5.



Figure 3-5: Concentration profiles of TCS (A); Σ CTDs (B); 2,8-DCDD (C); and 1,2,8-TriCDD, 2,3,7-TriCDD, and 1,2,3,8-TCDD (D) in the Lake Pepin core. Focus-corrected accumulation rates are presented in (E-H) for the respective analytes. Details on sediment focusing calculations can be found in Appendix E. Focusing factors specific to the Lake Pepin core are provided in Table AE-13.



Figure 3-6: Concentration profiles of TCS (A); Σ CTDs (B); 2,8-DCDD (C); and 1,2,8-TriCDD, 2,3,7-TriCDD, and 1,2,3,8-TCDD (D) in the Lake St. Croix core. Focus-corrected accumulation rates are presented in (E-H) for the respective analytes. Details on sediment focusing calculations can be found in Appendix E. Focusing factors specific to the Lake St. Croix core are provided in Table AE-14.

In the St. Croix sediment core, changes in wastewater disinfection appear to be recorded in the CTD profile. The St. Croix Valley WWTP opened in 1959 approximately 10 km upstream from the selected coring site. The facility treated its effluent with Cl₂ until 1993, when UV disinfection technology was installed during a plant expansion.

The visible inflection in the CTD profile around 1990 may be due to this change in disinfection technology (Figure 3-6). This inflection, however, is not present in the 1,2,8-TriCDD, 2,3,7-TriCDD, and 1,2,3,8-TCDD profiles, which may suggest that photochemical yields of these PCDDs from CTDs were low during this time period.

CTDs and their PCDDs are present in the St. Croix core at appreciable levels, even after UV disinfection was installed at the St. Croix Valley WWTP. This is likely due to other wastewater discharges, including that from the City of Hudson, WI WWTP, a facility 2.5 km upstream from the coring site that disinfects with Cl₂.

East Lake Gemini and Lake Winona. In contrast to large-scale riverine systems that integrate many wastewater sources, sediment cores collected from East Lake Gemini and Lake Winona highlight the impact of TCS, CTDs, and their PCDDs on small-scale lakes with a single wastewater source. Figures 3-7 and 3-8 depict analyte trends in East Lake Gemini and Lake Winona, respectively.

Improvements in wastewater treatment and changes in disinfection have greatly influenced the presence of TCS, CTDs, and their derived PCDDs in East Lake Gemini. After formation in 1966, East Lake Gemini received combined wastewater and stormwater from the SJU campus. Due to the transient nature of the campus population, coupled with extreme flows during storm events, SJU WWTP effluent was of poor quality during the 1960s and 1970s. To mitigate this problem, SJU separated stormwater and wastewater from the mid-1970s to late-1980s. Moreover, to stabilize flow through the WWTP and increase effluent quality, an 80,000 gallon equalization tank was installed upstream from secondary activated sludge treatment in 1980. The same year, the aeration system was upgraded in the activated sludge tank, dramatically improving

oxygen levels. These updates allowed for more consistent hydraulic and solids retention times during treatment and, subsequently, improvements in effluent quality.

Along with changes in wastewater treatment at the SJU WWTP, the mode of effluent disinfection has been modified since 1966. Cl_2 was the primary disinfectant until 1978, when an ozonation system was installed. UV disinfection then replaced ozonation in 1995 and is the current mode of wastewater disinfection at the treatment plant.

When examining analyte profiles in East Lake Gemini, there are clear differences from those observed in Lake Pepin or Lake St. Croix. TCS and 2,8-DCDD concentrations, for example, are much higher in this system, reflecting the direct discharge of wastewater from the SJU WWTP into a small impoundment (0.1 km^2). Moreover, concentrations of TCS, CTDs, and their derived PCDDs consistently decrease from 1966 to 2008 in the sediment core. Prior to 1978, concentrations of TCS, CTDs, and their PCDDs are all high, namely due to poor removal of TCS and disinfection with Cl₂ at the SJU plant. The transition to other forms of disinfection (e.g. ozonation and UV), stabilization of wastewater flow, and improvement activated sludge aeration likely contributed to a reduction in analyte sediment concentrations in this aquatic system.



Figure 3-7: Concentration profiles of TCS (A); Σ CTDs (B); 2,8-DCDD (C); and 1,2,8-TriCDD, 2,3,7-TriCDD, and 1,2,3,8-TCDD (D) in the East Lake Gemini core. Focus-corrected accumulation rates are presented in (E-H) for the respective analytes. Details on sediment focusing calculations can be found in Appendix E. A focusing factor, specific to the East Lake Gemini core, is provided in Table AE-15.



Figure 3-8: Concentration profiles of TCS (A); Σ CTDs (B); 2,8-DCDD (C); and 1,2,8-TriCDD, 2,3,7-TriCDD, and 1,2,3,8-TCDD (D) in the Lake Winona core. Focus-corrected accumulation rates are presented in (E-H) for the respective analytes. Details on sediment focusing calculations can be found in Appendix E. A focusing factor, specific to the Lake Winona core, is provided in Table AE-15.

In recent East Lake Gemini sediment, there is evidence that CTDs are being discharged into the lake, even without chlorinating the SJU wastewater effluent. Although SJU does not chlorinate its potable water, it is likely that CTDs are being formed by the reaction of TCS with chlorinated household products (e.g. bleach) that are washed down the drain. Like TCS, CTDs are typically removed at high rates (>90%) in plants with activated sludge treatment [24,25]. At SJU, however, this may not always be the case. Even with plant improvements, the facility is not equipped to handle such a highly transient population. During large events on the SJU campus (e.g. athletic games, concerts), wastewater flow can exceed plant capacity, leading to poor effluent quality. Conversely, during the summer months, the campus population decreases dramatically. During these time periods, wastewater flow is minimal and biomass levels in secondary activated sludge treatment decrease accordingly. Consequently, when large populations return to SJU, the WWTP is often not operating at fully capacity, which may contribute to TCS and CTD loading into East Lake Gemini.

Lake Winona is a natural water body that is heavily impacted by wastewater. In 1971, the Alexandria Lakes Area Sanitation District (ALASD) was formed, serving the City of Alexandria, MN and surrounding communities. The ALASD WWTP became operational in 1977. Since this date, elevated total phosphorus concentrations have induced hypereutrophic conditions in the lake. In 2002, the Lake Winona was listed as an impaired water body as part of Section 303(d) of the Federal Clean Water Act due excess nutrient levels [96]. From 2000 to 2008, a hydrologic balance on Lake Winona indicated that 63% of total inflow was derived from the ALASD WWTP [96]. Thus, the levels of TCS, CTDs, and their PCDDs in Lake Winona sediments are reflective of a small-scale aquatic environment heavily impacted by wastewater. Although there is a discrepancy between radiometric dating and the onset of TCS, CTDs, and their PCDDs in the sediment core (see Figure 3-8), analyte trends provide important insight into the exposure of Lake Winona to these compounds. TCS concentrations reach as high as 175 ng/g in sections of the core, followed by a steady decrease to 50 ng/g in recent sediment. When accounting for increased sedimentation rates, however, TCS loading is relatively constant since 1970 (Figure 3-8). The ALASD WWTP has consistently disinfected its wastewater with Cl₂, allowing CTD concentrations to rise to over 2000 pg/g in the sediment core.

Even in a hypereutrophic lake, direct photochemical transformation of TCS and CTDs to their respective PCDDs is still a concern. Lake Winona is a relatively shallow aquatic system, with a mean depth of 4.5 ft. Moreover, the current residence time and average Secchi depth for the lake are 70 days and 1.5 ft, respectively [97]. With wastewater accounting for 63% of inflow into Lake Winona, it is not surprising that TCS and CTD derived PCDDs are prevalent in sediments at this site.

Duluth Harbor and Lake Superior. Analyte trends are depicted in Figures 3-9 and 3-10 for the Duluth Harbor and Lake Superior cores, respectively. The Duluth Harbor core was collected approximately 2 km from the Western Lake Superior Sanitation District (WLSSD) wastewater outfall in St. Louis Bay. In the mid-1970s, the WLSSD WWTP began discharging effluent that was disinfected year-round with Cl₂. In June 1994, the treatment plant was granted a variance by the Minnesota Pollution Control

Agency (MPCA) to disinfect wastewater effluent only when fecal coliform levels exceeded 100 MPN/100 mL, where MPN is the most probably number of fecal coliforms in a sample. The WLSSD WWTP continues to operate under this variance today and disinfects on an as-needed basis.



Figure 3-9: Concentration profiles of TCS (A); Σ CTDs (B); 2,8-DCDD (C); and 1,2,8-TriCDD, 2,3,7-TriCDD, and 1,2,3,8-TCDD (D) in the Duluth Harbor core. Focus-corrected accumulation rates are presented in (E-H) for the respective analytes. Details on sediment focusing calculations can be found in Appendix E. A focusing factor, specific to the Duluth Harbor core, is provided in Table AE-15.



Figure 3-10: Concentration profiles of TCS (A); 2,8-DCDD (B); and 1,2,8-TriCDD, 2,3,7-TriCDD, and 1,2,3,8-TCDD (C) in the Lake Superior core. Focus-corrected accumulation rates are presented in (D-F) for the respective analytes. CTDs were below their respective LOQs in each core section. Details on sediment focusing calculations can be found in Appendix E. A focusing factor, specific to the Lake Superior core, is provided in Table AE-15.

TCS, CTD, and PCDD trends in Figure 3-9 clearly demonstrate the impact of WLSSD wastewater effluent on St. Louis Bay. Overall, TCS and 2,8-DCDD increase in the sediment core since the mid-1960s. The CTDs and their PCDDs, however, tend to co-vary with changes in wastewater disinfection at WLSSD. Specifically, total CTD levels drop from approximately 1700 pg/g in 1988 to 650 pg/g in 2002. A corresponding decrease is observed in the CTD derived tri- and tetra-CDDs around the same time period. Similar trends are present when accounting for varying sedimentation rates at the core site (Figure 3-9).

When examining the temporal trends of 2,8-DCDD, 1,2,8-TriCDD, 2,3,7-TriCDD, and 1,2,3,8-TCDD in Duluth Harbor, it is apparent that these PCDD congeners are present at low concentrations in the sediment core, prior to the patenting of TCS in the mid-1960s. 2,8-DCDD, for example, is present at approximately 10 pg/g in core sections dating back to the late-1800s. Furthermore, concentrations of the tri-and tetra-CDDs average around 4 pg/g during the same time period. The same PCDD congeners, however, are at much lower concentrations at the base of the Lake Pepin and Lake St. Croix cores. 2,8-DCDD is present at < 1 pg/g prior to 1930 in these cores. Moreover, 1,2,8-TriCDD, 2,3,7-TriCDD, and 1,2,3,8-TCDD are not detectable in the St. Croix core and are at very low levels (< 0.06 pg/g) in the Lake Pepin core before 1947. Although evidence suggests that TCS and CTDs are the primary source of these PCDDs, their presence at the base of the Duluth Harbor core may suggest a secondary, region-specific source.

The sediment core collected in Lake Superior provides evidence for TCS contamination in this aquatic system, albeit at low levels. TCS is first detected in the

sediment core around 1975, with concentrations increasing to 0.94 ng/g in 2005 (Figure 3-10). CTDs were not detected in the core, but may be present below the quantification limits established for this study. The source of TCS to this site on the open lake may be from distal wastewater sources, such as WLSSD or the City of Superior. Another possible source is the WWTP in Two Harbors, MN, where an average TCS concentration of 572 ng/L in wastewater effluent was recently documented [98].

Similar to Duluth Harbor, background levels of 2,8-DCDD, 1,2,8-TriCDD, 2,3,7-TriCDD, and 1,2,3,8-TCDD are present at the base of the Lake Superior core. Concentrations of 2,8-DCDD increase between 1975 and 2005, reflecting a contribution of this PCDD congener from the phototransformation of TCS in the system. The tri- and tetra-CDD congeners, however, do not follow a particular trend relative to TCS. A fraction of these PCDDs may be derived from CTDs after 1975. This, however, cannot be confirmed with the data in this study.

It is important to note that 2,8-DCDD is present at ~ 10 pg/g in the Lake Superior core, prior to the detection of TCS in the system. Furthermore, the CTD derived tri-and tetra-CDD congeners range from 1 to 6 pg/g throughout the core profile. These levels are similar to those observed in Duluth Harbor sediment before 1960, which may provide additional support for a secondary, region-specific source of these PCDDs.

Lake Shagawa. The City of Ely, MN has been discharging wastewater into Lake Shagawa for over a century. In the late 1880s, a primary wastewater treatment plant was established, where a single clarifier reduced the discharge of total suspended solid (TSS) and biological oxygen demand (BOD) into the lake. The city concurrently used Lake Shagawa as a drinking water source until 1932, when the water was rendered no longer suitable for potable use [99]. In 1954, the facility was upgraded to a secondary treatment plant, where a secondary clarifier, trickling filters, and disinfection with Cl₂ improved effluent quality [100]. This treatment facility, however, did not remove significant amounts of nutrients from wastewater effluent. By 1970, the Ely WWTP was contributing 80 percent of external total phosphorus loading to Lake Shagawa, contributing to eutrophic conditions in the water body [101].

To mitigate the nutrient loading problem to Lake Shagawa, advanced tertiary treatment, funded by the U.S. EPA, was established in 1973. In this tertiary facility, secondary wastewater effluent was treated with lime, alum, and/or polymeric flocculating agents to remove high levels of phosphorus. Following this treatment step, wastewater was passed through dual media filters (e.g., sand, anthracite) and disinfected with Cl₂ before being discharged into the lake.

Figure 3-11 illustrates the depositional trends of TCS, CTDs, and their PCDDs in Lake Shagawa sediment. Similar to Lake Winona, there is a discrepancy between the radiometric dating and the onset of TCS and CTD appearance in this lake system. The dating for this core, however, is currently being re-evaluated. It is anticipated that 1930 in the current dating profile corresponds to ~ 1960, based on the analyte trends.



Figure 3-11: Concentration profiles of TCS (A); Σ CTDs (B); 2,8-DCDD (C); and 1,2,8-TriCDD, 2,3,7-TriCDD, and 1,2,3,8-TCDD (D) in the Lake Shagawa core. Focus-corrected accumulation rates are presented in (E-H) for the respective analytes. Details on sediment focusing calculations can be found in Appendix E. A focusing factor, specific to the Lake Shagawa core, is provided in Table AE-15.
The TCS profile in Lake Shagawa sediments, like that of Lake Pepin and Lake St. Croix, mirrors the increased use of TCS since the 1960s. The CTD profile directly parallels this trend, reflecting the disinfection of the Ely WWTP effluent with Cl₂ since 1954. Moreover, 2,8-DCDD, 1,2,8-TriCDD, 2,3,7-TriCDD, and 1,2,3,8-TCDD trends directly follow with their TCS and CTD precursors.

Lake Little Wilson (Reference Lake). Figure 3-12 illustrates the analytes present in Lake Little Wilson, a remote lake with no wastewater input near the BWCA in MN. TCS and CTDs were not quantifiable in all core segments. Their derived PCDDs, however, are present at low levels throughout the core. 2,8-DCDD, for example, is present from 1872 to 2007 at an average concentration of 25 pg/g. The tri- and tetra-CDD congeners derived from CTDs are only detectable from 1933 to 2007 and at much lower concentrations (~ 0.5 pg/g). Due to the remote nature of Little Lake Wilson, it is likely that these PCDD congeners, among others, entered the lake via atmospheric deposition.

Several studies have investigated the deposition of PCDDs in the Great Lakes region [102-104]. Baker and Hites [103], for example, analyzed the temporal trends of tetra- through octa-CDD isomers in sediment cores collected from Siskiwit Lake on Isle Royale. Isle Royale is an isolated island in Lake Superior that is uninhabited and infrequently visited. Moreover, Siskiwit Lake is approximately 17 m above the elevation of Lake Superior. Thus, the only likely PCDD source to this lake is through atmospheric deposition. Through their study, Hites and Baker documented a rapid increase in total PCDD flux to their core sites from 1930 to 1970, with octa- and hepta-CDD being the most abundant isomers. This increase is attributed to the manufacturing and incineration

of organochlorine-based products during and after World War II. In 1970, emissions control devices aided in the reduction of PCDD loading to the atmosphere from incineration facilities. The Siskiwit Lake sediment cores record this reduction, where a 50% decrease in total PCDD loading from 1970 to 1998 was observed. A study by Pearson et al. [104] recorded similar PCDD depositional trends in a different remote lake near Lake Superior.

The presence of tetra- through octa-CDD isomers in remote lakes confirms that atmospheric deposition is an important source of higher chlorinated PCDDs to these aquatic systems. Although the studies above did not quantify di- or tri-CDD isomers, it is possible that very low concentrations of these PCDDs are present in their sediment cores. Typically, when PCDDs are emitted from a combustion source (e.g., a municipal incinerator), they are associated with particles, mix with the ambient atmosphere, and become diluted. During this process, lower chlorinated PCDDs have a higher potential to partition into the vapor phase. Due to their increased hydrophobicity, higher chlorinated PCDDs, like hepta-CDD and octa-CDD, are more likely to remain sorbed onto particles. Consequently, these particles become enriched with higher chlorinated PCDDs as they move away from the incineration source and eventually deposit in surface waters. This phenomena likely explains why most PCDD profiles derived from atmospheric sources are dominated by higher chlorinated PCDDs and only have traces of lower chlorinated congeners.



Figure 3-12: Concentration profiles of 2,8-DCDD (A); and 1,2,8-TriCDD, 2,3,7-TriCDD, and 1,2,3,8-TCDD (B) in the Lake Little Wilson core. Focus-corrected accumulation rates are presented in (C-D) for the respective analytes. TCS and CTDs are below their respective LOQs in all core segments. Details on sediment focusing calculations can be found in Appendix E. A focusing factor, specific to the Lake Little Wilson core, is provided in Table AE-15.

While it is possible that incineration contributes to the 2,8-DCDD, 1,2,8-TriCDD, 2,3,7-TriCDD, and 1,2,3,8-TCDD deposition in aquatic environments, it is clear that TCS and CTDs are typically the dominant source of these PCDD congeners after 1965 in systems heavily impacted by wastewater. This is exemplified when comparing analyte accumulation rates at the eight coring sites (Figures 3-5 to 3-12).

The majority of atmospheric loading of PCDDs to aquatic environments in recent history has been from the combustion of chlorinated synthetic compounds. Before 1930, however, low levels of PCDDs were produced from a variety of sources. A number of studies have suggested that the burning of coal and wood were important sources in the past [105-109]. Furthermore, forest fires have been cited as natural PCDD sources [108,110-111]. Gullet and Touati [110], in particular, recently evaluated PCDD emissions from different types of forest biomass. In general, Gullet and Touati found that the type of biomass can have a significant effect on the PCDD isomer patterns emitted during forest fires. Due to the distinct differences in forest biomes in MN (e.g., coniferous in the northeast, deciduous in the central and southeast), it is feasible that PCDD emissions and isomer abundances from these environments are unique. This may explain why TCS and CTD derived PCDDs are observed at low levels during the late 1800s in the Duluth Harbor, Lake Superior, and Lake Little Wilson cores, but are virtually non-detectable in the Lake Pepin and Lake St. Croix cores during the same time period. It is also important to note that sedimentation rates in Lake Pepin and Lake St. Croix are generally higher than in the northern lakes during this time period. Thus, higher sedimentation rates may have diluted a background level of TCS and CTD derived PCDDs below detection limits in these cores.

1.3.3 TCS and CTDs – An Emerging PCDD Source

The temporal trends of 2,8-DCDD, 1,2,8-TriCDD, 2,3,7-TriCDD, and 1,2,3,8-TCDD provide compelling evidence that TCS and CTDs are their dominant source in wastewater-impacted aquatic systems. Further evidence is granted by examining the ratio of these PCDDs to their TCS and CTD precursors in sediment core intervals. Tables 3-9 and 3-10 present these ratios, expressed as a percentage, for the sediment cores in this study. This allows for direct comparison to PCDD yields from TCS and CTDs presented in the literature.

Table 3-9: Ratio of 2,8-DCDD to TCS in sediment core intervals. Values are represented as a percentage for a comparison to PCDD yields from TCS in natural waters.

Sediment Core	[TCS derived PCDD]	^a]/[TCS](%)
Lake Pepin	1.0 ± 0.5^{b}	(n = 12)
Lake St. Croix	1.2 ± 0.4	(n = 9)
East Lake Gemini	1.6 ± 0.5	(n = 8)
Lake Winona	6.4 ± 5.0	(n = 11)
Duluth Harbor	1.5 ± 0.6	(n = 11)
Lake Superior	6.0 ± 2.2	(n = 4)
Lake Shagawa	1.8 ± 1.6	(n = 12)

^a [2,8-DCDD]

^b Reported values are the mean ± standard deviation

As outlined in Chapter 2, the experimental photochemical yields of TCS and CTD derived PCDDs range from 0.5-3% in natural waters [26,75]. The ratios in Tables 3-9 and 3-10 are within or near this range, with a few exceptions. The mean ratio for Lake Winona in Table 3-10, for example, is quite high. When accounting for the relative recovery of CTDs in the sediment core extractions, however, this ratio is close to

photochemical yields presented in the literature. The fact that the mean values presented in Tables 3-9 and 3-10 are close to experimental yields suggests that TCS and CTDs, when present, are the primary sources of their known photoproducts in the sediment cores.

Table 3-10: Ratio of 1,2,8-TriCDD, 2,3,7-TriCDD, and 1,2,3,8-TCDD to CTDs in sediment core intervals. Values are represented as a percentage for a comparison to PCDD yields from CTDs in natural waters. The presented ratios should be considered overestimates, as CTD concentrations are not corrected for relative recovery.

Sediment Core	Σ ([CTD derived PCDDs ^{<i>a</i>}]) / Σ[CTDs] (%)
Lake Pepin	2.1 ± 1.0^{b}	(n = 10)
Lake St. Croix	0.6 ± 0.2	(n = 8)
East Lake Gemini	2.7 ± 1.5	(n = 6)
Lake Winona	13.8 ± 3.8	(n = 8)
Duluth Harbor	4.9 ± 5.5	(n = 12)
Lake Superior	-	-
Lake Shagawa	1.2 ± 0.7	(n = 12)

^{*a*} [1,2,8-TriCDD], [2,3,7-TriCDD], [1,2,3,8-TCDD]

^{*b*} Reported values are the mean \pm standard deviation

PCDD Trends – Lake Pepin. Evaluating the isomer-specific trends of PCDDs in a sediment core can help fingerprint the dominant dioxin sources to an aquatic environment over time. Figure 3-13 illustrates the focus-corrected accumulation rates of PCDD isomers to the Lake Pepin core site from 1940 to 2008. Since 1965, 2,8-DCDD has comprised at least 90% of the total DCDD loading to the system, and in many cases, was the only DCDD congener detected. Furthermore, 1,2,8-TriCDD and 2,3,7-TriCDD constituted 99% of detected TriCDD isomers. 1,2,3,8-TCDD was not detected before 1958, but constituted up to 40% of total TCDD isomers in recent sediment. Buth et al. [47] collected two sediment cores in Lake Pepin and recorded similar contributions of these TCS and CTD derived PCDD congeners. Moreover, acquisition windows for diand tri-CDD isomers were expanded during HRGC-HRMS analysis in Buth et al. [47] to evaluate the contribution of other DCDD and TriCDD congeners to PCDD loading. Expansion of these acquisition windows demonstrated that di- and tri-CDD congeners, other than those derived from TCS and CTDs, were not present at significant levels. Thus, the di- and tri-CDD contributions from TCS and CTDs to the total DCDD and TriCDD pool are considered representative for the current Lake Pepin core.

The temporal trends of the di- through octa-CDD isomers in Figure 3-13 illustrate distinct sources of PCDDs since 1940. The increase in di- and tri-CDD accumulation rates since 1965 is reflective of 2,8-DCDD, 1,2,8-TriCDD, and 2,3,7-TriCDD loading from TCS and CTDs. Similarly, the increase in TCDD accumulation rate from 1970 to 2008 is largely due to 1,2,3,8-TCDD, the primary photoproduct of 4,6-Cl-TCS. The higher chlorinated PCDD isomers, however, follow a different trend. OCDD accumulation in the Lake Pepin core, for example, increases in the 1940s and 1950s, peaks around 1970, and subsequently decreases after 1980. Similar to the higher chlorinated isomer profiles observed in Lake Siskiwit sediment, this trend directly mirrors the atmospheric input of PCDDs from incineration sources in the 20th century.

To illustrate the differences between PCDD sources in the Lake Pepin core, specific congeners and isomers profiles were correlated against 2,8-DCDD, the known photoproduct of TCS, and OCDD, the most abundant PCDD derived from incineration sources. Figure 3-14 depicts the results of these correlations.



profiles include the contribution of the PCDD congeners derived from TCS and CTDs. Details on sediment focusing calculations can be referenced in Appendix E. Figure 3-13: Focus-corrected PCDD accumulation rates at the Lake Pepin core site. DCDD, TriCDD, and TCDD



Figure 3-14: Correlation of PCDD congener and isomer accumulation profiles with 2,8-DCDD (A) and OCDD (B) in the Lake Pepin core. Grey bars represent the TCS and CTD derived PCDD congeners.

Results from the correlation analysis are clear for Lake Pepin. CTD derived PCDDs positively correlate with 2,8-DCDD, with R² values of 0.93, 0.94, and 0.73 for 1,2,8-TriCDD, 2,3,7-TriCDD, and 1,2,3,8-TCDD, respectively. Overall, 2,8-DCDD also correlates well with the DCDD, TriCDD, and TCDD isomer groups. Higher chlorinated isomers, however, do not correlate well with 2,8-DCDD, suggesting a different source. OCDD, for example, correlates well with other higher chlorinated PCDDs, but poorly with lower chlorinated isomers and the congeners derived from TCS and CTDs. The contrasting PCDD isomer profiles in Figure 3-13, coupled with the correlations in Figure 3-14, suggest that TCS and CTDs are a distinct and emerging PCDD source in Lake Pepin.

PCDD Trends – Lake Superior. As a contrast to Lake Pepin, PCDD trends were examined in Lake Superior, a large lake with a less pronounced wastewater impact. Figure 3-15 illustrates the temporal loading of di- through octa-CDD isomers to the Lake Superior core site from 1880 to 2006. In this system, 2,8-DCDD contributes > 91% of total DCDD loading after 1978, a time period corresponding to the detection of TCS in the sediment profile. Before this date, however, the 2,8-DCDD contribution averages 80% and is as low as 61%. This may reflect that, prior to the presence of TCS in the lake, 2,8-DCDD was part of a broader suite of lower chlorinated PCDDs derived from a secondary source. The contribution of 1,2,8-TriCDD and 2,3,7-TriCDD to the TriCDD isomer pool is consistently > 98% throughout the core profile. This contribution may be unrepresentative, as the first and last TriCDD eluters during HRGC-HRMS analysis were not quantified. Thus, similar to 2,8-DCDD, 1,2,8-TriCDD and 2,3,7-TriCDD may be part of a larger array of lower chlorinated PCDD congeners in the core.



PCDD Accumulation (pg/cm² yr)

focusing calculations can be referenced in Appendix E. TCDD profiles include the contribution of the PCDD congeners derived from TCS and CTDs. Details on sediment Figure 3-15: Focus-corrected PCDD accumulation rates at the Lake Superior core site. DCDD, TriCDD, and



Figure 3-16: Correlation of PCDD congener and isomer accumulation profiles with 2,8-DCDD (A) and OCDD (B) in the Lake Superior core. Grey bars represent the TCS and CTD derived PCDD congeners.

Unlike the di- and tri-CDD analysis, all tetra-CDD congeners were quantified during HRGC-HRMS analysis for the Lake Superior core. Thus, the contribution of 1,2,3,8-TCDD to the TCDD pool should be considered representative. As indicated above, 1,2,3,8-TCDD became a prominent TCDD congener in the Lake Pepin core after 1958, contributing up to 40% of TCDD in the presence of CTDs. In the Lake Superior core, however, 1,2,3,8-TCDD contributes an average of 8% and a maximum of 17% throughout the core profile. A portion of this contribution may be derived from CTDs, below their limit of quantification in the core. However, because CTDs are not quantifiable, it is feasible that 1,2,3,8-TCDD is introduced to Lake Superior from another source.

Figure 3-16 presents the results of a correlation analysis for Lake Superior. Analogous to the Lake Pepin analysis, 2,8-DCDD and OCDD are correlated against specific PCDD congeners and isomers. Interestingly, 2,8-DCDD does not correlate well with 1,2,8-TriCDD or 2,3,7-TriCDD in the core. OCDD, however, does correlate relatively well with these congeners, which may support the theory of a secondary source in the region. 2,8-DCDD does not correlate with OCDD. This is likely due to the contribution of this congener from TCS in more recent sections of the core.

The PCDD trends and correlations in Figures 3-15 and 3-16 clearly illustrate that there may be a series of PCDD sources to Lake Superior, of which TCS is only a small component. Low levels of di-through-octaCDD at the base of the core may suggest that natural sources, such as forest fires, have contributed to a background level of PCDDs in this environment. Moreover, because low levels of higher chlorinated PCDD congeners are present at the base of the core, one cannot discount the potential for anaerobic reductive dechlorination as potential source of lower chlorinated PCDDs in old sediments. Finally, it is possible that traces of TCS and CTD derived PCDDs were introduced from nearby incineration sources, such as steel or paper mills in the Duluth, MN area throughout the 20th century. With the data in this study, however, it is difficult to implicate a specific secondary source of these PCDDs.

3.4 Environmental Implications

The depositional trends of TCS and CTDs in lake sediments provide important insight into the historical exposure of wastewater-impacted aquatic ecosystems to these compounds. This exposure is dynamically linked to the size of the receiving water body and degree of wastewater impact. For example, in small-scale wastewater impacted lakes, such as East Lake Gemini and Lake Winona, high concentrations of TCS and CTDs in the sediment record likely reflect the long-term exposure of aquatic organisms to elevated levels of these contaminants. Consequently, this exposure may be adversely impacting algal and macrophyte community structure, enhancing bioaccumulation in higher trophic levels, and inducing a range of endocrine disrupting effects on aquatic biota. Moreover, because wastewater typically contains a diverse array of PPCPs, it is probable that TCS and CTDs are part of a complex mixture of bioactive and persistent compounds, leading to unknown consequences for the receiving aquatic ecosystem.

Due to the harmful effects of PCDDs in the environment, the loading of TCS and CTD derived PCDDs to aquatic ecosystems is of clear concern. Therefore, the contribution of 2,8-DCDD, 1,2,8-TriCDD, 2,3,7-TriCDD, and 1,2,3,8-TriCDD to the total PCDD pool, in terms of mass and toxicity, was determined for the lakes in this

study. The contribution to total toxicity was determined by using toxic equivalency factors (TEFs), where toxicity of specific PCDD congeners and isomers are normalized to the most toxic PCDD congeners: 2,3,7,8-TCDD and 1,2,3,7,8-PeCDD (Table 3-11). TEFs are multiplied by the concentration of a given PCDD congener or isomer to provide their toxic equivalency (TEQ) in an environmental matrix. These TEQs are then added to determine the total TEQ attributed to PCDDs in a given sample (Eq. 6):

$$Total TEQ = \Sigma \left[PCDD_i \times TEF_i \right]$$
(6)

Table 3-11: Toxic equivalency factors (TEFs) for PCDD congeners and isomers, relative to 2,3,7,8-TCDD.

PCDD Congener/Isomer	Toxic Equivalency Factor (TEF)
DCDD	0.001^{a}
TriCDD	0.01^{a}
2,3,7,8-TCDD	1^b
other TCDDs	0.01^{a}
1,2,3,7,8-PeCDD	1^b
other PeCDDs	0.1^b
HxCDD	0.1^b
HpCDD	0.01^{b}
OCDD	0.0003^{b}

^a Ontario Ministry of the Environment, 1985

^b Van den Berg et al., 2006

The TEF values presented in Table 3-11 were obtained from two sources [112,113]. Due to an abundance of toxicity data on tetra- through octa-CDDs in the literature, TEF values for these isomer groups have been well refined in recent years. The values in Table 3-11, for example, were determined by the World Health Organization (WHO) in 2005 as part of a re-evaluation of mammalian TEFs for dioxins

and dioxin-like compounds. Lower chlorinated congeners (e.g., di- and tri-CDDs), however, have been poorly studied. Values for these isomers in Table 3-11 are derived from a 1985 study by the Ontario Ministry of the Environment. More recent work has proposed that di- and tri-CDDs may be less toxic, based on a series of bioassays on specific chlorinated and brominated dioxins [114]. Therefore, di- and tri-CDD TEFs presented in Table 3-11 in should be considered a worst-case scenario when calculating the TEQ for these isomers.

Figures 3-17 and 3-18 illustrate the contribution of 2,8-DCDD, 1,2,8-TriCDD, 2,3,7-TriCDD, and 1,2,3,8-TCDD, in terms of mass and toxicity, to the PCDD pool in each sediment core. In the Lake Pepin core, the contribution of these PCDD congeners has been increasing since the patent of TCS in 1964, reaching up to 30% of total PCDD mass and 5% of total PCDD toxicity in recent sediment. This trend is reflective of the photochemical transformation of TCS and CTDs in the Mississippi River. A similar trend is observed in the Lake St. Croix sediment core. Due to a less pronounced wastewater impact to this system, however, these PCDD congeners only constitute up to 6.5% of total PCDD mass and 1.6% of PCDD toxicity in recent years.



Figure 3-17: Percent contribution of 2,8-DCDD, 1,2,8-TriCDD, 2,3,7-TriCDD, and 1,2,3,8-TCDD by mass and toxic equivalents (TEQ) to the total PCDD pool for Lake Pepin (A), Lake St. Croix (B), East Lake Gemini (C), and Lake Winona (D). Mass and TEQ contributions for all dated core sections are provided in Appendix F.

The contribution of 2,8-DCDD, 1,2,8-TriCDD, 2,3,7-TriCDD, and 1,2,3,8-TCDD to the East Lake Gemini and Lake Winona PCDD pools is reflective of the heavy wastewater impact in each system. In East Lake Gemini, these PCDD congeners consistently account for greater than 46% of total PCDD loading to the system and up to 23% of PCDD toxicity. Although the formation of these PCDDs in East Lake Gemini has decreased since the 1960s, atmospheric loading of higher chlorinated PCDDs to the lake has been reduced over a similar timeline. Thus, the overall contribution of TCS and CTD derived PCDDs to this system remains significant. The photochemical transformation of TCS and CTDs has had a major impact on the PCDD loading and toxicity in Lake Winona. The di-, tri-, and tetra-CDDs derived from TCS and CTDs contribute up to 57% of PCDD mass loading and 32% of PCDD toxicity to the system.

The contribution of TCS and CTD derived PCDDs to total PCDD mass and toxicity in Duluth Harbor, Lake Superior, Lake Shagawa, and Little Lake Wilson is presented in Figure 3-18. Based on the estimation that a secondary, region-specific source has contributed to the loading of TCS and CTD derived PCDDs near Duluth, MN, it is possible that a fraction of each profile is attributed to this source. In the Duluth Harbor core, TCS and CTDs have clearly been the dominant source of 2,8-DCDD, 1,2,8-TriCDD, 2,3,7-TriCDD, and 1,2,3,8-TCDD since the 1960s. The contribution of these congeners to total PCDD mass and toxicity in harbor sediment, however, is relatively low. This is due to heavy contamination of the sediments with higher chlorinated congeners from a variety of industrial operations in near Duluth, including paper and steel mills.



Figure 3-18: Percent contribution of 2,8-DCDD, 1,2,8-TriCDD, 2,3,7-TriCDD, and 1,2,3,8-TCDD by mass and toxic equivalents (TEQ) to the total PCDD pool for Duluth Harbor (A), Lake Superior (B), Lake Shagawa (C), and Lake Little Wilson (D). Mass and TEQ contributions for all dated core sections are provided in Appendix F.

As illustrated in Section 3.3.3, there is likely a diverse array of PCDD sources to the Lake Superior coring site, some of which may have contributed to the di-,tri-, and tetra-CDD loading to the sediment. After 1975, however, the PCDD contribution by mass and toxicity from 2,8-DCDD increased by a factor of 5. This is likely attributed to the photochemical transformation of TCS in the lake and suggests that TCS is a dominant source of this PCDD to Lake Superior today.

TCS and CTD derived PCDDs contribute appreciably to the total PCDD loading in Lake Shagawa. Since 1930, which likely corresponds to ~1960 in the core profile, these PCDDs have contributed at least 5% and up to 36% of total PCDD mass in Lake Shagawa sediment. Toxicity contributions reach as high as 6% in recent sediment. The visible inflection in the profiles presented in Figure 3-18 are attributed to changes in atmospheric PCDD loading throughout the 20th century.

The presence of 2,8-DCDD, 1,2,8-TriCDD, 2,3,7-TriCDD, and 1,2,3,8-TCDD in Lake Little Wilson suggest that traces of these PCDDs can be derived from atmospheric sources. The mass and toxicity contributions of these PCDD congeners are depicted in Figure 3-18 from 1950 to 2010, a time period relevant to TCS and CTDs in the environment. During this time interval, the di-, tri-, and tetra-CDDs consistently account for 5% of total PCDD mass and ~ 0.4% of total PCDD toxicity. However, if you account for the high relative recovery of 2,8-DCDD in Lake Little Wilson sediment (up to 238% - see Section 3.3.1), then the mass and toxicity contributions are on the order of 1.5% and 0.2%, respectively.

Chapter 4: Conclusions

Results presented in this study document the presence of TCS, CTDs, and their derived PCDDs in an array of wastewater-impacted MN lakes. Dated sediment cores provide a unique perspective on the historical exposure of regional aquatic ecosystems to these deleterious compounds. Furthermore, the sediment cores provide valuable insight into how wastewater treatment practices and lake system scale influence the depositional trends of TCS, CTDs, and their PCDDs in the sedimentary record. Finally, evaluation of the TCS and CTD derived PCDD contribution to the total PCDD pool, in terms of mass and toxicity, aids in assessing the potential risk that TCS poses to aquatic environments.

When comparing large and small-scale lake systems, there are clear differences in the concentrations and depositional trends of TCS, CTDs, and their PCDDs. Lake Pepin, for example, integrates many wastewater sources in the Mississippi River watershed, including the Metropolitan WWTP. Increasing analyte concentrations in this core are reflective of the large-scale integration of TCS and CTDs from various wastewater sources since the mid-1960s. Conversely, in small-scale systems like East Lake Gemini and Lake Winona, analyte trends are attributed to a single wastewater source. In East Lake Gemini, improvements in wastewater treatment and changes in disinfection at the SJU WWTP are directly recorded in TCS, CTD, and derived PCDD profiles. Furthermore, the high analyte concentrations found in Lake Winona are attributed to chlorinated ALASD WWTP effluent accounting for 63% of total inflow into the small, shallow water body.

Changes in wastewater disinfection can have a dramatic impact on the discharge of CTDs and formation of tri- and tetra-CDDs into aquatic environments. The Duluth Harbor core, for example, records a 50% decrease in CTD and 1,2,8-TriCDD, 2,3,7-TriCDD, and 1,2,3,8-TCDD loading from 1988 to 2002. This decrease is directly attributed to the disinfection variance granted to the WLSDD WWTP in 1994.

Several observations throughout this study suggest that TCS and CTDs are the primary sources of 2,8-DCDD, 1,2,8-TriCDD, 2,3,7-TriCDD, and 1,2,3,8-TCDD in aquatic systems heavily impacted by wastewater. The ratios of these PCDDs to their TCS and CTD precursors in the sediment cores are similar to those observed in photolysis experiments. Furthermore, the analysis of di- through octa-CDD trends in the Lake Pepin core suggest that TCS and CTDs have been a distinct and emerging source of these PCDDs since the 1960s in wastewater-impacted systems. Low levels of TCS and CTD derived PCDDs, however, are present prior to the patent of TCS in the Duluth Harbor core. Moreover, similar levels of these PCDDs are present throughout the Lake Superior core in the absence of detectable CTDs and in Lake Little Wilson, an aquatic system with no wastewater impact. This suggests that a secondary, region-specific atmospheric source may be contributing to low background levels of these PCDDs in northern MN. To delineate this secondary source, future work may analyze sediment extracts with expanded di- and tri-CDD acquisition windows during HRGC-HRMS analysis. If a broader array of DCDD and TriCDD congeners are present, this may help fingerprint a specific atmospheric source.

An integral component of this research was to determine the potential risk that TCS and its byproducts pose to aquatic ecosystems. In general, the loading of TCS and CTD derived PCDDs, in terms of mass and toxicity, has been increasing in MN lakes since the 1960s. The Lake Pepin core is a prime example of this trend, with TCS and CTD derived PCDDs accounting for up to 30% of total PCDD mass and 5% of total PCDD toxicity in recent sediment. In small-scale lakes, like East Lake Gemini and Lake Winona, the TCS and CTD derived PCDD burden is much greater. In Lake Winona sediment, for example, these PCDDs account for up to 52% of total PCDD mass and 32% of total PCDD toxicity. This PCDD contribution, coupled with the presence of high concentrations of TCS and CTDs, likely poses a substantial risk to this aquatic ecosystem. Moreover, it is likely that similar risks are present in other small-scale aquatic systems that receive municipal and industrial wastewater around the U.S. and the world.

To better quantify the risks associated with TCS and CTD derived PCDDs, future work should focus on refining the estimated TEF values for di- and tri-CDD congeners. With improved estimates of 2,8-DCDD, 1,2,8-TriCDD, 2,3,7-TriCDD, and 1,2,3,8-TCDD toxicity to a variety of aquatic organisms, more informed risk assessments could be made on the harmful effects of TCS byproducts in surface waters. Furthermore, if TCS use continues to increase in personal care products, strict regulations may be needed for reducing levels of TCS and CTDs in wastewater effluents. Advanced oxidation processes, including ozonation [115,116] and photocatalysis [117], have been shown to rapidly degrade TCS in water with low dissolved organic carbon (DOC). Furthermore, membrane filtration may be a promising treatment option for TCS removal in wastewater effluent [118]. Consequently, integration of these processes in WWTPs may reduce TCS in wastewater effluent, minimize the formation of CTDs, and prevent the formation of their PCDDs in the environment.

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Appendix A

General Laboratory Preparation and Practice

To prevent analyte contamination during sample preparation and extraction, a series of detailed procedures were followed in the laboratory. Freeze drying tins, plastic syringes for silica columns, and all glassware, with the exception of disposable glass pipettes, were triple washed consecutively with a dilute solution of alconox and deionized (DI) water. The glassware, including the pipettes, were then covered with foil and fired at 550 °C for 4 hours. Freeze drying tins, plastic syringes, and any glassware that could not be fired were triple rinsed with methanol and ethyl acetate. All cleaned materials were stored in a designated cabinet to prevent contamination. Silica gel, glass wool, and Ottawa sand were all fired at 550 °C for 4 hours before use in ASE cells or silica columns.

Prior to working in the laboratory, bench-tops areas and fumehoods to be used were wiped down with alconox. Surfaces were then covered with foil to help prevent contamination.

Gas-tight, glass syringes were used to prepare standards, spike samples, and transfer final extracts to autosampler vials. Three separate syringes were designated for $^{13}C_{12}$ -TCS, TCS, and CTD solutions to prevent analyte cross-contamination. Moreover, one syringe was always kept clean for sample resuspension. Finally, a fifth syringe was used to transfer final extracts to autosampler vials. Regardless of use, all syringes were rinsed 5 times with acetone, acetonitrile, and methanol prior to each transfer.

Accelerated Solvent Extraction (ASE) 350 - Cleaning Procedures

Several cleaning procedures are required when using the ASE 350 to minimize analyte contamination. The stainless steel ASE cell end caps, in particular, must be carefully cleaned, as they tend to accumulate analytes over time. For this project, ASE end caps were triple rinsed with tap water, DI water, methanol, and ethyl acetate. Detergents, including alconox, cannot be used on the endcaps because they degrade the seals. After this washing procedure, the cells were disassembled, placed in a clean jar, and sonicated in acetone for 10 minutes. The end caps were then reassembled and subsequently rinsed three times with methanol and ethyl acetate to remove any remaining contamination.

The stainless steel bodies of the ASE cells were cleaned by triple rinsing with alconox, DI water, methanol and ethyl acetate. All specialty tools were cleaned using the same procedure.

After a sequence of ASE extractions, the solvent lines were rinsed 5 times consecutively with acetone as a preventative maintenance step. Dichloromethane (DCM), the mobile phase used for extraction in this study, is a very strong solvent that will corrode the ASE pump if left in the lines for long periods of time (e.g., for a month). To help maintain the ASE, lines should always be flushed with acetone prior to shutting the instrument down.
Appendix B

LC-MS-Q³ – Operational Guidance for TCS and CTD Analysis

Prior to analyzing sediment extracts for TCS and CTDs on the LC-MS-Q³ at the Masonic Cancer Research Center, a specific instrument preparation procedure must be followed. Specifically, a clean ion-transfer tube should be installed in the MS-Q³ and preconditioned with 3-5 existing sediment extracts, before running any instrument blanks, standards, or samples. This process has been shown to provide consistent and improved analyte response in sediment matrices. For an unknown reason, the analytes of interest ($^{13}C_{12}$ -TCS, TCS, and CTDs) take a period of time to equilibrate in the MS-Q³ (e.g., to provide stable and consistent analytical response). Preconditioning of a clean ion-transfer tube with existing sediment extracts containing these analytes reduces this equilibration time.

Appendix C

Calculation of TCS and CTD Concentrations in Sediment Matrices

The following material outlines the calculations used to determine TCS and CTD concentrations in sediment matrices using isotope-dilution methodology. Concentrations are determined by multiplying (1) a response factor from the analyte calibration curve, (2) the analyte to ${}^{13}C_{12}$ -TCS peak area ratio in the sediment matrix, and (3) the amount of ${}^{13}C_{12}$ -TCS spiked into the sample. Figure AC-1 provides an example calibration curve for the calculations.





Figure AC-1: Example TCS calibration curve. Notation: PA = peak area, std = standard.

Equation 1 outlines the calculation of TCS mass in a sample, using isotope dilution methodology:

$$TCS_{\text{sample}}(g) = [RF] \times \frac{[TCS PA_{\text{sample}}]}{[^{13}C_{12} - TCS PA_{\text{sample}}]} \times [^{13}C_{12} - TCS_{\text{sample}}(g)]$$
(1)

where:

$$RF = \frac{1}{[{}^{13}C_{12} - TCS_{std} (g/L)][calibration slope]}$$

and:

calibration slope =
$$\frac{\text{TCS PA}_{,\text{std}}}{[^{13}\text{C}_{12} - \text{TCS PA}_{,\text{std}}][\text{TCS}_{,\text{std}} (g/L)]}$$

Equation 1 can be expanded as follows:

$$TCS_{,sample} (g) = \frac{[{}^{13}C_{12} - TCS PA_{,std}][TCS_{,std} (g/L)]}{[{}^{13}C_{12} - TCS_{,std} (g/L)][TCS PA_{,std}]} \times \frac{[TCS PA_{,sample}]}{[{}^{13}C_{12} - TCS PA_{,sample}]} \times [{}^{13}C_{12} - TCS PA_{,sample}]$$
(2)

$$\times [{}^{13}C_{12} - TCS_{,sample} (g)]$$

When normalized to the amount of sediment analyzed in an extract, Equation 2 provides the concentration of TCS in a given sample.

Absolute Recovery of ¹³C₁₂-TCS

The absolute recovery of ${}^{13}C_{12}$ -TCS in sediment extracts was determined using Equation 3:

$${}^{13}C_{12}\text{-}TCS \text{ Recovery} = \frac{[{}^{13}C_{12} - TCS \text{ PA},_{\text{sample}}]}{[{}^{13}C_{12} - TCS \text{ PA},_{\text{std}}(\text{avg})]} \times \frac{[{}^{13}C_{12} - TCS,_{\text{std}}(g/L)]}{[{}^{13}C_{12} - TCS,_{\text{sample}}(g/L)]} \times 100$$
(3)

where: ${}^{13}C_{12}$ -TCS PA, $_{std}$ (avg) = the average ${}^{13}C_{12}$ -TCS PA in the calibration curve.

Appendix D

¹³C₁₂-PCDD/F and PCDD/F Recoveries – Pace Analytical

Table AD-1: Absolute ${}^{13}C_{12}$ -PCDD/F recovery in the Lake Pepin sediment core.

¹³ C ₁₂ -PCDD/F Congener	Absolute Recovery (%)	
¹³ C ₁₂ -2,3-DCDD	^a 47 ± 7.4	(n = 16)
¹³ C ₁₂ -2,3,7-TriCDD	77 ± 9.1	(n = 16)
¹³ C ₁₂ -2,3,7,8-TCDF	73 ± 7.4	(n = 16)
¹³ C ₁₂ -2,3,7,8-TCDD	87 ± 7.2	(n = 16)
¹³ C ₁₂ -1,2,3,7,8-PeCDF	75 ± 7.2	(n = 16)
¹³ C ₁₂ -2,3,4,7,8-PeCDF	75 ± 8.6	(n = 16)
¹³ C ₁₂ -1,2,3,7,8-PeCDD	85 ± 9.8	(n = 16)
¹³ C ₁₂ -1,2,3,4,7,8-HxCDF	79 ± 10.5	(n = 16)
¹³ C ₁₂ -1,2,3,6,7,8-HxCDF	74 ± 4.0	(n = 16)
¹³ C ₁₂ -2,3,4,6,7,8-HxCDF	66 ± 4.2	(n = 16)
¹³ C ₁₂ -1,2,3,7,8,9-HxCDF	75 ± 7.7	(n = 16)
¹³ C ₁₂ -1,2,3,4,7,8-HxCDD	83 ± 13.4	(n = 16)
¹³ C ₁₂ -1,2,3,6,7,8-HxCDD	69 ± 6.0	(n = 16)
¹³ C ₁₂ -1,2,3,4,6,7,8-HpCDF	61 ± 5.7	(n = 16)
¹³ C ₁₂ -1,2,3,4,7,8,9-HpCDF	60 ± 6.4	(n = 16)
¹³ C ₁₂ -1,2,3,4,6,7,8-HpCDD	62 ± 6.1	(n = 16)
¹³ C ₁₂ -OCDD	55 ± 6.8	(n = 16)

^aReported values are the mean ± standard deviation

Table AD-2: Relative PCDD/F recovery in Lake Pepin sediment (spin)	ke and recovery).
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PCDD/F Congener	Relative Recovery (%)	^b RPD
2,8-DCDD	^a 92	7.3
2,3,7-TriCDD	108	4.6
2,3,7,8-TCDF	119	1.4
2,3,7,8-TCDD	87	1.4
1,2,3,7,8-PeCDF	110	2.6
2,3,4,7,8-PeCDF	108	0.2
1,2,3,7,8-PeCDD	95	3.8
1,2,3,4,7,8-HxCDF	107	2.2
1,2,3,6,7,8-HxCDF	107	1.5
2,3,4,6,7,8-HxCDF	106	0.5
1,2,3,7,8,9-HxCDF	106	0.2
1,2,3,4,7,8-HxCDD	108	1.2
1,2,3,6,7,8-HxCDD	117	0.4
1,2,3,7,8,9-HxCDD	109	0.1
1,2,3,4,6,7,8-HpCDF	108	3.7
1,2,3,4,7,8,9-HpCDF	98	1.7
1,2,3,4,6,7,8-HpCDD	98	0.3
OCDF	97	1.7
OCDD	98	1.8

^a Reported values are the mean relative recovery (n = 2) ^b RPD = relative percent difference between matrix spike replicates

¹³ C ₁₂ -PCDD/F Congener	Absolute Recovery (%)	
¹³ C ₁₂ -2,3-DCDD	^a 48 ± 9.4	(n = 16)
¹³ C ₁₂ -2,3,7-TriCDD	76 ± 5.8	(n = 16)
¹³ C ₁₂ -2,3,7,8-TCDF	82 ± 5.0	(n = 16)
¹³ C ₁₂ -2,3,7,8-TCDD	88 ± 7.1	(n = 16)
¹³ C ₁₂ -1,2,3,7,8-PeCDF	84 ± 4.4	(n = 16)
¹³ C ₁₂ -2,3,4,7,8-PeCDF	82 ± 5.4	(n = 16)
¹³ C ₁₂ -1,2,3,7,8-PeCDD	88 ± 6.4	(n = 16)
¹³ C ₁₂ -1,2,3,4,7,8-HxCDF	81 ± 11.5	(n = 16)
¹³ C ₁₂ -1,2,3,6,7,8-HxCDF	79 ± 7.5	(n = 16)
¹³ C ₁₂ -2,3,4,6,7,8-HxCDF	76 ± 6.5	(n = 16)
¹³ C ₁₂ -1,2,3,7,8,9-HxCDF	71 ± 12.7	(n = 16)
¹³ C ₁₂ -1,2,3,4,7,8-HxCDD	78 ± 5.7	(n = 16)
¹³ C ₁₂ -1,2,3,6,7,8-HxCDD	67 ± 4.3	(n = 16)
¹³ C ₁₂ -1,2,3,4,6,7,8-HpCDF	50 ± 9.7	(n = 16)
¹³ C ₁₂ -1,2,3,4,7,8,9-HpCDF	45 ± 12.1	(n = 16)
¹³ C ₁₂ -1,2,3,4,6,7,8-HpCDD	45 ± 8.6	(n = 16)
¹³ C ₁₂ -OCDD	39 ± 8.3	(n = 16)

Table AD-3: Absolute ${}^{13}C_{12}$ -PCDD/F recovery in the Lake St. Croix sediment core.

Table AD-4: Relative PCDD/F recovery in Lake St. Croix sediment (spike and recovery).

PCDD/F Congener	Relative Recovery (%)	^b RPD
2,8-DCDD	^a 100	6.8
2,3,7-TriCDD	108	2.6
2,3,7,8-TCDF	119	4.4
2,3,7,8-TCDD	96	0.4
1,2,3,7,8-PeCDF	111	1.8
2,3,4,7,8-PeCDF	109	2.9
1,2,3,7,8-PeCDD	97	6.2
1,2,3,4,7,8-HxCDF	110	8.3
1,2,3,6,7,8-HxCDF	111	4.1
2,3,4,6,7,8-HxCDF	106	3.6
1,2,3,7,8,9-HxCDF	108	2.3
1,2,3,4,7,8-HxCDD	117	12.9
1,2,3,6,7,8-HxCDD	115	8.1
1,2,3,7,8,9-HxCDD	109	5.6
1,2,3,4,6,7,8-HpCDF	109	5.2
1,2,3,4,7,8,9-HpCDF	103	10.9
1,2,3,4,6,7,8-HpCDD	102	6.7
OCDF	100	5.3
OCDD	107	9.0

^a Reported values are the mean relative recovery (n = 2)

¹³ C ₁₂ -PCDD/F Congener	Absolute Recovery (%)	
¹³ C ₁₂ -2,3-DCDD	^{<i>a</i>} 48 ± 3.3	(n = 8)
¹³ C ₁₂ -2,3,7-TriCDD	80 ± 3.7	(n = 8)
¹³ C ₁₂ -2,3,7,8-TCDF	76 ± 3.9	(n = 8)
¹³ C ₁₂ -2,3,7,8-TCDD	89 ± 6.0	(n = 8)
¹³ C ₁₂ -1,2,3,7,8-PeCDF	80 ± 5.4	(n = 8)
¹³ C ₁₂ -2,3,4,7,8-PeCDF	81 ± 5.2	(n = 8)
¹³ C ₁₂ -1,2,3,7,8-PeCDD	91 ± 5.2	(n = 8)
¹³ C ₁₂ -1,2,3,4,7,8-HxCDF	85 ± 5.3	(n = 8)
¹³ C ₁₂ -1,2,3,6,7,8-HxCDF	76 ± 3.0	(n = 8)
¹³ C ₁₂ -2,3,4,6,7,8-HxCDF	69 ± 4.8	(n = 8)
¹³ C ₁₂ -1,2,3,7,8,9-HxCDF	79 ± 3.9	(n = 8)
¹³ C ₁₂ -1,2,3,4,7,8-HxCDD	90 ± 4.9	(n = 8)
¹³ C ₁₂ -1,2,3,6,7,8-HxCDD	73 ± 3.5	(n = 8)
¹³ C ₁₂ -1,2,3,4,6,7,8-HpCDF	63 ± 4.0	(n = 8)
¹³ C ₁₂ -1,2,3,4,7,8,9-HpCDF	61 ± 4.7	(n = 8)
¹³ C ₁₂ -1,2,3,4,6,7,8-HpCDD	62 ± 4.0	(n = 8)
¹³ C ₁₂ -OCDD	56 ± 3.8	(n = 8)

 Table AD-5:
 Absolute ¹³C₁₂-PCDD/F recovery in the East Lake Gemini sediment core.

¹³ C ₁₂ -PCDD/F Congener	Absolute Recovery (%)	
¹³ C ₁₂ -2,3-DCDD	^a 42 ± 6.6	(n = 11)
¹³ C ₁₂ -2,3,7-TriCDD	63 ± 5.8	(n = 11)
¹³ C ₁₂ -2,3,7,8-TCDF	63 ± 6.4	(n = 11)
¹³ C ₁₂ -2,3,7,8-TCDD	78 ± 6.9	(n = 11)
¹³ C ₁₂ -1,2,3,7,8-PeCDF	70 ± 6.7	(n = 11)
¹³ C ₁₂ -2,3,4,7,8-PeCDF	64 ± 5.8	(n = 11)
¹³ C ₁₂ -1,2,3,7,8-PeCDD	70 ± 6.1	(n = 11)
¹³ C ₁₂ -1,2,3,4,7,8-HxCDF	65 ± 5.4	(n = 11)
¹³ C ₁₂ -1,2,3,6,7,8-HxCDF	67 ± 5.5	(n = 11)
¹³ C ₁₂ -2,3,4,6,7,8-HxCDF	60 ± 5.2	(n = 11)
¹³ C ₁₂ -1,2,3,7,8,9-HxCDF	61 ± 5.4	(n = 11)
¹³ C ₁₂ -1,2,3,4,7,8-HxCDD	62 ± 4.9	(n = 11)
¹³ C ₁₂ -1,2,3,6,7,8-HxCDD	53 ± 5.6	(n = 11)
¹³ C ₁₂ -1,2,3,4,6,7,8-HpCDF	48 ± 4.5	(n = 11)
¹³ C ₁₂ -1,2,3,4,7,8,9-HpCDF	44 ± 6.5	(n = 11)
¹³ C ₁₂ -1,2,3,4,6,7,8-HpCDD	45 ± 5.4	(n = 11)
¹³ C ₁₂ -OCDD	38 ± 6.8	(n = 11)

 Table AD-6:
 Absolute ¹³C₁₂-PCDD/F recovery in the Lake Winona sediment core.

Table AD-7: Relative PCDD/F recovery in Lake Winona sediment (spike and recovery).

PCDD/F Congener	Relative Recovery (%)	^b RPD
2,8-DCDD	^a 121	5.8
2,3,7-TriCDD	105	0.5
2,3,7,8-TCDF	119	1.9
2,3,7,8-TCDD	87	3.3
1,2,3,7,8-PeCDF	108	0.6
2,3,4,7,8-PeCDF	107	1.1
1,2,3,7,8-PeCDD	96	1.5
1,2,3,4,7,8-HxCDF	109	3.6
1,2,3,6,7,8-HxCDF	105	0.4
2,3,4,6,7,8-HxCDF	105	2.6
1,2,3,7,8,9-HxCDF	106	2.8
1,2,3,4,7,8-HxCDD	114	6.1
1,2,3,6,7,8-HxCDD	115	3.7
1,2,3,7,8,9-HxCDD	111	0.6
1,2,3,4,6,7,8-HpCDF	111	0.4
1,2,3,4,7,8,9-HpCDF	100	0.9
1,2,3,4,6,7,8-HpCDD	96	4.4
OCDF	101	10.2
OCDD	75	5.9

^a Reported values are the mean relative recovery (n = 2)

¹³ C ₁₂ -PCDD/F Congener	Absolute Recovery (%)	
¹³ C ₁₂ -2,3-DCDD	^a 75 ± 6.5	(n = 13)
¹³ C ₁₂ -2,3,7-TriCDD	80 ± 9.8	(n = 13)
¹³ C ₁₂ -2,3,7,8-TCDF	85 ± 4.1	(n = 13)
¹³ C ₁₂ -2,3,7,8-TCDD	98 ± 4.7	(n = 13)
¹³ C ₁₂ -1,2,3,7,8-PeCDF	85 ± 2.5	(n = 13)
¹³ C ₁₂ -2,3,4,7,8-PeCDF	80 ± 3.5	(n = 13)
¹³ C ₁₂ -1,2,3,7,8-PeCDD	84 ± 6.2	(n = 13)
¹³ C ₁₂ -1,2,3,4,7,8-HxCDF	87 ± 5.6	(n = 13)
¹³ C ₁₂ -1,2,3,6,7,8-HxCDF	86 ± 6.5	(n = 13)
¹³ C ₁₂ -2,3,4,6,7,8-HxCDF	77 ± 3.0	(n = 13)
¹³ C ₁₂ -1,2,3,7,8,9-HxCDF	72 ± 4.2	(n = 13)
¹³ C ₁₂ -1,2,3,4,7,8-HxCDD	74 ± 5.6	(n = 13)
¹³ C ₁₂ -1,2,3,6,7,8-HxCDD	68 ± 5.9	(n = 13)
¹³ C ₁₂ -1,2,3,4,6,7,8-HpCDF	51 ± 4.3	(n = 13)
¹³ C ₁₂ -1,2,3,4,7,8,9-HpCDF	50 ± 5.4	(n = 13)
¹³ C ₁₂ -1,2,3,4,6,7,8-HpCDD	51 ± 6.1	(n = 13)
¹³ C ₁₂ -OCDD	37 ± 5.5	(n = 13)

Table AD-8: Absolute ${}^{13}C_{12}$ -PCDD/F recovery in the Duluth Harbor sediment core.

Table AD-9: Relative PCDD/F recovery in Duluth Harbor sediment (spike and recovery). _

PCDD/F Congener	Relative Recovery (%)	^b RPD
2,8-DCDD	^a 97	9.2
2,3,7-TriCDD	104	5.5
2,3,7,8-TCDF	111	3.4
2,3,7,8-TCDD	86	2.3
1,2,3,7,8-PeCDF	112	2.3
2,3,4,7,8-PeCDF	111	1.9
1,2,3,7,8-PeCDD	94	4.5
1,2,3,4,7,8-HxCDF	116	4.5
1,2,3,6,7,8-HxCDF	111	2.6
2,3,4,6,7,8-HxCDF	109	1.9
1,2,3,7,8,9-HxCDF	110	3.1
1,2,3,4,7,8-HxCDD	106	0.9
1,2,3,6,7,8-HxCDD	120	0.2
1,2,3,7,8,9-HxCDD	124	3.1
1,2,3,4,6,7,8-HpCDF	116	6.0
1,2,3,4,7,8,9-HpCDF	102	5.7
1,2,3,4,6,7,8-HpCDD	99	3.4
OCDF	115	2.9
OCDD	111	0.9

^a Reported values are the mean relative recovery (n = 2) ^b RPD = relative percent difference between matrix spike replicates

¹³ C ₁₂ -PCDD/F Congener	Absolute Recovery (%)	
¹³ C ₁₂ -2,3-DCDD	^a 62 ± 7.3	(n = 10)
¹³ C ₁₂ -2,3,7-TriCDD	78 ± 3.8	(n = 10)
¹³ C ₁₂ -2,3,7,8-TCDF	84 ± 3.6	(n = 10)
¹³ C ₁₂ -2,3,7,8-TCDD	104 ± 3.7	(n = 10)
¹³ C ₁₂ -1,2,3,7,8-PeCDF	92 ± 3.1	(n = 10)
¹³ C ₁₂ -2,3,4,7,8-PeCDF	98 ± 3.0	(n = 10)
¹³ C ₁₂ -1,2,3,7,8-PeCDD	109 ± 3.8	(n = 10)
¹³ C ₁₂ -1,2,3,4,7,8-HxCDF	81 ± 3.2	(n = 10)
¹³ C ₁₂ -1,2,3,6,7,8-HxCDF	87 ± 3.0	(n = 10)
¹³ C ₁₂ -2,3,4,6,7,8-HxCDF	87 ± 2.4	(n = 10)
¹³ C ₁₂ -1,2,3,7,8,9-HxCDF	95 ± 4.5	(n = 10)
¹³ C ₁₂ -1,2,3,4,7,8-HxCDD	85 ± 5.6	(n = 10)
¹³ C ₁₂ -1,2,3,6,7,8-HxCDD	78 ± 2.8	(n = 10)
¹³ C ₁₂ -1,2,3,4,6,7,8-HpCDF	80 ± 2.9	(n = 10)
¹³ C ₁₂ -1,2,3,4,7,8,9-HpCDF	92 ± 4.3	(n = 10)
¹³ C ₁₂ -1,2,3,4,6,7,8-HpCDD	92 ± 3.8	(n = 10)
¹³ C ₁₂ -OCDD	85 ± 3.7	(n = 10)

Table AD-10: Absolute ${}^{13}C_{12}$ -PCDD/F recovery in the Lake Superior sediment core.

Table AD-11: Relative PCDD/F recovery in Lake Superior sediment (spike and recovery).

PCDD/F Congener	Relative Recovery (%)	^b RPD
2,8-DCDD	^a 101	8.0
2,3,7-TriCDD	111	17.5
2,3,7,8-TCDF	114	1.3
2,3,7,8-TCDD	78	4.8
1,2,3,7,8-PeCDF	107	4.2
2,3,4,7,8-PeCDF	107	4.6
1,2,3,7,8-PeCDD	90	5.6
1,2,3,4,7,8-HxCDF	106	0.3
1,2,3,6,7,8-HxCDF	101	5.6
2,3,4,6,7,8-HxCDF	99	1.7
1,2,3,7,8,9-HxCDF	100	4.4
1,2,3,4,7,8-HxCDD	100	2.9
1,2,3,6,7,8-HxCDD	110	1.4
1,2,3,7,8,9-HxCDD	107	1.2
1,2,3,4,6,7,8-HpCDF	101	2.4
1,2,3,4,7,8,9-HpCDF	94	4.2
1,2,3,4,6,7,8-HpCDD	90	2.0
OCDF	103	4.1
OCDD	99	1.4

^{*a*} Reported values are the mean relative recovery (n = 2)

¹³ C ₁₂ -PCDD/F Congener	Absolute Recovery (%)	
¹³ C ₁₂ -2,3-DCDD	^{<i>a</i>} 52 ± 14.7	(n = 12)
¹³ C ₁₂ -2,3,7-TriCDD	75 ± 12.7	(n = 12)
¹³ C ₁₂ -2,3,7,8-TCDF	69 ± 11.7	(n = 12)
¹³ C ₁₂ -2,3,7,8-TCDD	88 ± 14.6	(n = 12)
¹³ C ₁₂ -1,2,3,7,8-PeCDF	74 ± 12.3	(n = 12)
¹³ C ₁₂ -2,3,4,7,8-PeCDF	73 ± 12.6	(n = 12)
¹³ C ₁₂ -1,2,3,7,8-PeCDD	82 ± 14.3	(n = 12)
¹³ C ₁₂ -1,2,3,4,7,8-HxCDF	69 ± 13.1	(n = 12)
¹³ C ₁₂ -1,2,3,6,7,8-HxCDF	74 ± 13.5	(n = 12)
¹³ C ₁₂ -2,3,4,6,7,8-HxCDF	71 ± 12.1	(n = 12)
¹³ C ₁₂ -1,2,3,7,8,9-HxCDF	76 ± 12.7	(n = 12)
¹³ C ₁₂ -1,2,3,4,7,8-HxCDD	73 ± 12.4	(n = 12)
¹³ C ₁₂ -1,2,3,6,7,8-HxCDD	65 ± 11.6	(n = 12)
¹³ C ₁₂ -1,2,3,4,6,7,8-HpCDF	65 ± 11.5	(n = 12)
¹³ C ₁₂ -1,2,3,4,7,8,9-HpCDF	70 ± 13.9	(n = 12)
¹³ C ₁₂ -1,2,3,4,6,7,8-HpCDD	70 ± 14.5	(n = 12)
¹³ C ₁₂ -OCDD	64 ± 12.8	(n = 12)

 Table AD-12:
 Absolute ¹³C₁₂-PCDD/F recovery in the Lake Shagawa sediment core.

Table AD-13: Relative PCDD/F recovery in Lake Shagawa sediment (spike and recovery). _

PCDD/F Congener	Relative Recovery (%)	^b RPD
2,8-DCDD	^a 116	6.8
2,3,7-TriCDD	112	1.4
2,3,7,8-TCDF	124	1.9
2,3,7,8-TCDD	86	1.7
1,2,3,7,8-PeCDF	115	0.4
2,3,4,7,8-PeCDF	113	1.3
1,2,3,7,8-PeCDD	100	1.1
1,2,3,4,7,8-HxCDF	116	0.0
1,2,3,6,7,8-HxCDF	113	2.3
2,3,4,6,7,8-HxCDF	110	0.5
1,2,3,7,8,9-HxCDF	112	0.8
1,2,3,4,7,8-HxCDD	108	5.2
1,2,3,6,7,8-HxCDD	122	4.4
1,2,3,7,8,9-HxCDD	116	2.0
1,2,3,4,6,7,8-HpCDF	113	0.2
1,2,3,4,7,8,9-HpCDF	101	1.6
1,2,3,4,6,7,8-HpCDD	100	6.0
OCDF	109	6.4
OCDD	109	2.9

^a Reported values are the mean relative recovery (n = 2) ^b RPD = relative percent difference between matrix spike replicates

Table AD-14: Absolute ${}^{13}C_{12}$ -PCDD/F recovery in the Lake Little Wilson sedimentcore.

¹³ C ₁₂ -PCDD/F Congener	Absolute Recovery (%)	
¹³ C ₁₂ -2,3-DCDD	^a 76 ± 7.4	(n = 10)
¹³ C ₁₂ -2,3,7-TriCDD	79 ± 5.6	(n = 10)
¹³ C ₁₂ -2,3,7,8-TCDF	84 ± 9.3	(n = 10)
¹³ C ₁₂ -2,3,7,8-TCDD	102 ± 6.6	(n = 10)
¹³ C ₁₂ -1,2,3,7,8-PeCDF	86 ± 4.7	(n = 10)
¹³ C ₁₂ -2,3,4,7,8-PeCDF	75 ± 11.7	(n = 10)
¹³ C ₁₂ -1,2,3,7,8-PeCDD	93 ± 7.2	(n = 10)
¹³ C ₁₂ -1,2,3,4,7,8-HxCDF	91 ± 8.0	(n = 10)
¹³ C ₁₂ -1,2,3,6,7,8-HxCDF	101 ± 8.8	(n = 10)
¹³ C ₁₂ -2,3,4,6,7,8-HxCDF	48 ± 21.5	(n = 10)
¹³ C ₁₂ -1,2,3,7,8,9-HxCDF	83 ± 4.9	(n = 10)
¹³ C ₁₂ -1,2,3,4,7,8-HxCDD	69 ± 7.9	(n = 10)
¹³ C ₁₂ -1,2,3,6,7,8-HxCDD	90 ± 8.6	(n = 10)
¹³ C ₁₂ -1,2,3,4,6,7,8-HpCDF	62 ± 6.0	(n = 10)
¹³ C ₁₂ -1,2,3,4,7,8,9-HpCDF	59 ± 7.0	(n = 10)
¹³ C ₁₂ -1,2,3,4,6,7,8-HpCDD	50 ± 8.6	(n = 10)
¹³ C ₁₂ -OCDD	47 ± 6.8	(n = 10)

Table AD-15:	Relative PCDD/F re	ecovery in Lake	Little Wilson	sediment (sp	ike and
recovery).					

PCDD/F Congener	Relative Recovery (%)	^b RPD
2,8-DCDD	^a 233	4.0
2,3,7-TriCDD	104	1.3
2,3,7,8-TCDF	115	2.4
2,3,7,8-TCDD	74	2.2
1,2,3,7,8-PeCDF	105	0.4
2,3,4,7,8-PeCDF	104	0.8
1,2,3,7,8-PeCDD	87	0.3
1,2,3,4,7,8-HxCDF	108	0.5
1,2,3,6,7,8-HxCDF	96	0.4
2,3,4,6,7,8-HxCDF	110	11.2
1,2,3,7,8,9-HxCDF	103	2.3
1,2,3,4,7,8-HxCDD	98	14.2
1,2,3,6,7,8-HxCDD	108	10.7
1,2,3,7,8,9-HxCDD	103	6.6
1,2,3,4,6,7,8-HpCDF	104	0.9
1,2,3,4,7,8,9-HpCDF	91	0.0
1,2,3,4,6,7,8-HpCDD	93	0.0
OCDF	103	4.5
OCDD	103	0.5

^a Reported values are the mean relative recovery (n = 2)

Appendix E

Core Radiometric Dating: Lead-210 (²¹⁰Pb) and Cesium - 137 (¹³⁷Cs)

 Table AE-1:
 ²¹⁰Pb dating for the Lake St. Croix sediment core.

Top of	Base of	Cum.	Unsup.	Error of	Cum. Act.	Age: Base	Error of	Date	Sediment	Error of
Interval	Interval	Dry Mass	Activity	Unsup. Act.	below Int.	of Int.	Age	A.D.	Accum.	Sed. Accum.
(cm)	(cm)	(g/cm²)	(pCi/g)	(±s.d.)	(pCi/cm ²)	(yr)	(±s.d.)		(g/cm ² yr)	(±s.d.)
0	4	0.3449	10.7482	0.4304	63.7538	1.81	2.83	2008.7	0.1900	0.0140
4	8	0.8063	10.0442	0.4880	59.1196	4.24	2.95	2006.3	0.1904	0.0155
12	16	1.9456	7.4293	0.3203	49.9999	9.62	3.23	2000.9	0.2189	0.0191
20	24	3.4456	7.5128	0.3692	38.7598	17.80	3.84	1992.8	0.1728	0.0184
32	36	6.1345	3.7362	0.2460	25.8279	30.83	3.61	1979.7	0.2297	0.0249
44	48	9.2211	2.0437	0.2126	17.9582	42.50	3.64	1968.1	0.2900	0.0388
56	60	12.7143	1.2498	0.1838	12.7341	53.54	4.05	1957.0	0.3358	0.0584
68	72	16.5234	0.8977	0.1611	8.8850	65.10	4.97	1945.5	0.3280	0.0707
80	84	20.3662	0.8728	0.1741	5.4991	80.51	7.21	1930.0	0.2154	0.0579
92	96	24.3010	0.6530	0.1568	2.6491	103.96	13.13	1906.6	0.1461	0.0598
104	108	28.6548	0.3370	0.1341	0.7461	144.65	37.85	1865.9	0.0904	0.0876
116	120	33.0477	0.0484	0.1492	0.1208	203.12	165.94	1807.4	0.0985	0.4763
				_						
Supported	Pb-210:	2.1407 ± 0.12	223 pCi/g]			Cum. Unsu	p. Pb-210:	67.4607 p	oCi/cm ²
Number of	Supported	Samples:	3]			Unsup. Pb-	210 Flux:	2.1638 p	Ci/cm ² yr

 Table AE-2:
 ²¹⁰Pb dating for the East Lake Gemini sediment core.

Top of	Base of	Cum.	Unsup.	Error of	Cum. Act.	Age: Base	Error of	Date	Sediment	Error of
Interval	Interval	Dry Mass	Activity	Unsup. Act.	below Int.	of Int.	Age	A.D.	Accum.	Sed. Accum.
(cm)	(cm)	(g/cm ²)	(pCi/g)	(±s.d.)	(pCi/cm ²)	(yr)	(±s.d.)		(g/cm ² yr)	(±s.d.)
0	4	0.1550	15.3132	0.5917	53.7821	1.39	0.94	2009.3	0.1117	0.0048
8	12	0.5727	13.4634	0.5371	47.9863	5.05	0.99	2005.6	0.1145	0.0051
12	16	0.8259	9.0967	0.3374	45.6826	6.63	1.01	2004.1	0.1603	0.0069
18	20	1.0656	7.2508	0.3411	43.8598	7.94	1.03	2002.8	0.1902	0.0100
20	22	1.2143	6.5013	0.3261	42.8933	8.65	1.05	2002.0	0.2078	0.0116
30	32	2.0967	6.2348	0.2721	37.3007	13.14	1.13	1997.6	0.1893	0.0098
40	42	3.1501	4.9285	0.2651	31.5929	18.47	1.13	1992.2	0.2030	0.0122
50	52	4.3007	5.2137	0.2607	25.7255	25.07	1.28	1985.6	0.1573	0.0093
60	62	5.5392	3.9961	0.2198	20.2116	32.82	1.18	1977.9	0.1614	0.0099
70	74	6.9945	4.4321	0.2469	14.0000	44.61	1.40	1966.1	0.1055	0.0067

Supported Pb-210:	0.8053 ± 0.1542 pCi/g
Number of Supported	d Samples: 3

Cum. Unsup. Pb-210:	56.1564 pCi/cm ²	
Unsup. Pb-210 Flux:	1.7934 pCi/cm ² yr	

Top of	Base of	Cum.	Unsup.	Error of	Cum. Act.	Age: Base	Error of	Date	Sediment	Error of
Interval	Interval	Dry Mass	Activity	Unsup. Act.	below Int.	of Int.	Age	A.D.	Accum.	Sed. Accum.
(cm)	(cm)	(g/cm ²)	(pCi/g)	(±s.d.)	(pCi/cm ²)	(yr)	(±s.d.)		(g/cm ² yr)	(±s.d.)
0	4	0.5716	3.4376	0.1563	19.2571	3.12	1.23	2007.6	0.1832	0.0093
4	8	1.2719	3.1088	0.1237	17.0799	6.97	1.30	2003.7	0.1818	0.0087
8	12	2.0417	3.4899	0.1862	14.3935	12.47	1.42	1998.2	0.1401	0.0083
12	14	2.4322	3.0827	0.1288	13.1898	15.27	1.50	1995.4	0.1392	0.0076
14	16	2.8204	3.1380	0.1614	11.9717	18.38	1.59	1992.3	0.1248	0.0079
20	22	4.0042	2.9427	0.1630	8.4126	29.71	2.04	1981.0	0.0951	0.0072
24	26	4.8196	2.7515	0.1438	6.1311	39.87	2.66	1970.8	0.0756	0.0066
26	28	5.2445	2.8396	0.1271	4.9244	46.91	3.23	1963.8	0.0604	0.0058
30	32	6.0921	2.0997	0.1121	2.9986	62.84	4.88	1947.9	0.0508	0.0071
32	34	6.5615	1.4514	0.0641	2.3173	71.12	6.27	1939.6	0.0567	0.0099
34	36	7.0602	1.1574	0.0466	1.7401	80.32	8.30	1930.4	0.0542	0.0123
36	38	7.5843	0.8124	0.0387	1.3143	89.33	10.96	1921.4	0.0582	0.0174
38	40	8.1047	0.4820	0.0304	1.0635	96.13	13.52	1914.6	0.0765	0.0293
40	42	8.6647	0.2749	0.0263	0.9095	101.15	15.80	1909.5	0.1115	0.0516
44	46	9.8029	0.2646	0.0265	0.6054	114.22	23.69	1896.5	0.0798	0.0532
48	50	11.0632	0.3046	0.0268	0.2340	144.75	61.16	1865.9	0.0330	0.0471
Supported	Pb-210:	0.1978 ± 0.0)18 pCi/g				Cum. Unsu	p. Pb-210:	21.2221 p	Ci/cm ²
Number of	Supported	Samples:	3				Unsup. Pb-	210 Flux:	0.6773 pC	Ci/cm² yr

 Table AE-3:
 ²¹⁰Pb dating for the Lake Winona sediment core.

Table AE-4:	²¹⁰ Pb dating for the Duluth Harbor sediment core.	
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Top of	Base of	Cum.	Unsup.	Error of	Cum. Act.	Age: Base	Error of	Date	Sediment	Error of
Interval	Interval	Dry Mass	Activity	Unsup. Act.	below Int.	of Int.	Age	A.D.	Accum.	Sed. Accum.
(cm)	(cm)	(g/cm ²)	(pCi/g)	(±s.d.)	(pCi/cm ²)	(yr)	(±s.d.)		(g/cm ² yr)	(±s.d.)
0	2	0.5809	3.3972	0.1766	28.8504	2.12	1.57	2008.6	0.2734	0.0167
2	4	1.2771	3.3137	0.1625	26.5435	4.80	1.64	2005.9	0.2601	0.0156
4	6	2.1365	3.0028	0.1630	23.9629	8.09	1.73	2002.7	0.2617	0.0171
8	10	4.1734	2.3712	0.1378	18.8419	15.81	1.94	1994.9	0.2638	0.0191
10	12	5.1894	1.9331	0.1100	16.8780	19.34	2.09	1991.4	0.2874	0.0217
12	14	6.1535	1.6272	0.1218	15.3092	22.47	2.24	1988.3	0.3077	0.0283
16	18	8.1187	1.5510	0.1002	12.2243	29.70	2.66	1981.0	0.2605	0.0245
18	20	9.0667	1.4454	0.1076	10.8541	33.52	2.93	1977.2	0.2483	0.0264
22	24	10.9643	1.1798	0.0969	8.4934	41.39	3.58	1969.4	0.2387	0.0302
24	26	11.9326	1.0367	0.0923	7.4896	45.43	4.01	1965.3	0.2397	0.0337
28	30	13.9559	1.0520	0.0926	5.3687	56.12	5.44	1954.6	0.1744	0.0300
30	32	14.9859	0.9970	0.0815	4.3418	62.94	6.65	1947.8	0.1511	0.0300
34	36	17.1446	0.6514	0.0797	2.7620	77.47	10.07	1933.3	0.1484	0.0444
36	38	18.2038	0.7364	0.0638	1.9820	88.12	13.95	1922.6	0.0994	0.0374
40	42	20.3218	0.5127	0.0600	0.7824	117.97	34.73	1892.8	0.0626	0.0529
42	44	21.5748	0.2482	0.0573	0.4714	134.24	57.41	1876.5	0.0770	0.1089
				-						
Supported	Pb-210:	0.7726 ± 0.05	509 pCi/g	1			Cum. Unsu	p. Pb-210:	30.824 pC	Ci/cm ²
Number of	Supported	Samples: 5		1			Unsup. Pb-	210 Flux:	0.9878 p	Ci/cm ² yr

Supported Pb-210:	0.7726 ± 0	.0509	pCi/g
Number of Supported	d Samples:	5	

Cum. Unsup. Pb-210:	30.824 pCi/cm ²
Jnsup. Pb-210 Flux:	0.9878 pCi/cm ² yr

Top of	Base of	Cum.	Unsup.	Error of	Cum. Act.	Age: Base	Error of	Date	Sediment	Error of
Interval	Interval	Dry Mass	Activity	Unsup. Act.	below Int.	of Int.	Age	A.D.	Accum.	Sed. Accum.
(cm)	(cm)	(g/cm ²)	(pCi/g)	(±s.d.)	(pCi/cm ²)	(yr)	(±s.d.)		(g/cm ² yr)	(±s.d.)
0	1	0.2670	11.1537	0.4206	29.4076	3.10	2.31	2007.6	0.0862	0.0054
2	3	0.8768	10.1771	0.3825	23.0623	10.90	2.66	1999.8	0.0754	0.0054
4	5	1.6224	6.8575	0.2829	17.4123	19.93	3.07	1990.8	0.0850	0.0073
6	7	2.4250	5.0320	0.2468	13.0340	29.23	3.74	1981.5	0.0868	0.0094
9	10	3.7590	2.2372	0.1538	9.0168	41.06	3.21	1969.7	0.1326	0.0140
11	12	4.6989	2.1241	0.1535	6.9940	49.22	3.91	1961.5	0.1097	0.0137
14	15	6.0382	1.5597	0.1405	4.6616	62.25	5.25	1948.5	0.0997	0.0168
16	17	6.9793	1.3668	0.1354	3.3320	73.03	7.13	1937.7	0.0832	0.0181
18	19	7.9482	0.8562	0.1174	2.3859	83.76	9.65	1927.0	0.0941	0.0284
19	20	8.4457	0.5467	0.1168	2.1139	87.64	10.82	1923.1	0.1280	0.0477
22	23	9.9900	0.2671	0.1074	1.5652	97.29	14.05	1913.5	0.1905	0.1081
24	25	11.0012	0.3761	0.1179	1.2133	105.47	17.94	1905.3	0.1081	0.0643
27	28	12.4561	0.4046	0.1131	0.6387	126.08	33.49	1884.7	0.0563	0.0534
29	30	13.4291	0.2486	0.1076	0.3601	144.48	58.92	1866.3	0.0524	0.0857
30	32	14.4053	0.1426	0.1012	0.2209	160.17	94.95	1850.6	0.0622	0.1496
Supported	Pb-210:	1.131 ± 0.09	304 pCi/g			,	Cum. Unsu	.p. Pb-210	32.3862 p	Ci/cm ²
Number of	Supported	Samples:	3			P	Unsup. Pb	-210 Flux:	1.031 pC	Ji/cm ² yr

 Table AE-5:
 ²¹⁰Pb dating for the Lake Superior sediment core.

 Table AE-6:
 ²¹⁰Pb dating for the Lake Shagawa sediment core.

Top of	Base of	Cum.	Unsup.	Error of	Cum. Act.	Age: Base	Error of	Date	Sediment	Error of
Interval	Interval	Dry Mass	Activity	Unsup. Act.	below Int.	of Int.	Age	A.D.	Accum.	Sed. Accum.
(cm)	(cm)	(g/cm ²)	(pCi/g)	(±s.d.)	(pCi/cm ²)	(yr)	(±s.d.)		(g/cm ² yr)	(±s.d.)
0	2	0.1268	15.9131	0.5688	38.3117	1.65	1.16	2009.8	0.0769	0.0033
4	6	0.4370	13.4765	0.2741	33.9610	5.52	1.23	2005.9	0.0810	0.0028
8	10	0.8125	11.6901	0.2458	29.4168	10.13	1.33	2001.3	0.0814	0.0031
12	14	1.2176	10.4672	0.2209	25.0567	15.28	1.47	1996.2	0.0777	0.0033
16	18	1.6301	9.5049	0.3403	21.0383	20.90	1.66	1990.6	0.0721	0.0040
20	22	2.0497	8.2273	0.3641	17.4577	26.89	1.91	1984.6	0.0693	0.0046
24	26	2.4630	6.6243	0.3027	14.5603	32.72	2.19	1978.7	0.0716	0.0054
30	32	3.1014	5.8806	0.2676	10.6532	42.75	2.84	1968.7	0.0597	0.0054
32	34	3.3130	5.1465	0.1910	9.5644	46.21	3.13	1965.3	0.0611	0.0059
36	38	3.7621	3.1404	0.1794	7.9540	52.13	3.61	1959.3	0.0824	0.0098
42	44	4.5074	2.0919	0.0949	6.1570	60.36	4.44	1951.1	0.0956	0.0131
46	48	5.0426	1.3995	0.0811	5.3219	65.04	5.08	1946.4	0.1226	0.0197
50	52	5.5861	0.9879	0.0881	4.7315	68.81	5.68	1942.7	0.1533	0.0293
52	54	5.8625	1.0100	0.0879	4.4523	70.77	6.03	1940.7	0.1415	0.0282
58	60	6.7377	1.0956	0.0822	3.5183	78.33	7.59	1933.1	0.1046	0.0247
60	64	7.3849	0.8744	0.0856	2.9524	83.96	9.01	1927.5	0.1149	0.0312
64	68	8.0402	0.5663	0.0736	2.5813	88.27	10.28	1923.2	0.1519	0.0490
72	76	9.3706	0.5344	0.0708	1.8597	98.80	14.20	1912.7	0.1185	0.0501
84	88	11.3420	0.3678	0.0713	1.0274	117.86	25.28	1893.6	0.0968	0.0708
Supported	Pb-210:	1.2097 ± 0.04	83 pCi/g				Cum. Unsu	p. Pb-210:	40.3297	oCi/cm ²
Number of	Supported	Samples: 3					Unsup. Pb-	210 Flux:	1.2808 p	Ci/cm ² yr

Top of	Base of	Cum.	Unsup.	Error of	Cum. Act.	Age: Base	Error of	Date	Sediment	Error of
Interval	Interval	Dry Mass	Activity	Unsup. Act.	below Int.	of Int.	Age	A.D.	Accum.	Sed. Accum.
(cm)	(cm)	(g/cm ²)	(pCi/g)	(±s.d.)	(pCi/cm ²)	(yr)	(±s.d.)		(g/cm ² yr)	(±s.d.)
0	2	0.0896	24.7811	0.5491	34.8219	1.99	0.57	2009.5	0.0451	0.0011
4	6	0.3038	21.1130	0.4671	30.1203	6.64	0.58	2004.8	0.0461	0.0012
8	10	0.5559	23.1291	0.5204	24.4140	13.39	0.63	1998.1	0.0348	0.0009
12	14	0.8263	20.5690	0.4664	18.6855	21.98	0.69	1989.5	0.0304	0.0008
14	16	0.9664	19.8735	0.4064	15.9013	27.16	0.76	1984.3	0.0270	0.0007
18	20	1.2692	17.9593	0.5867	10.3258	41.02	0.93	1970.4	0.0202	0.0008
20	22	1.4331	11.9421	0.2293	8.3689	47.77	1.10	1963.7	0.0243	0.0008
22	24	1.6010	9.3088	0.2170	6.8059	54.41	1.31	1957.1	0.0253	0.0010
24	26	1.7720	6.0378	0.2884	5.7735	59.69	1.50	1951.8	0.0324	0.0020
28	30	2.1210	4.1336	0.2167	4.1784	70.08	1.57	1941.4	0.0341	0.0022
32	34	2.4676	2.1717	0.1252	3.2752	77.90	1.15	1933.6	0.0496	0.0031
36	38	2.8205	1.7410	0.1256	2.6243	85.01	1.29	1926.5	0.0496	0.0038
40	42	3.1823	1.4688	0.1104	2.0691	92.65	1.51	1918.8	0.0466	0.0039
42	44	3.3590	1.1524	0.0975	1.8655	95.97	1.64	1915.5	0.0531	0.0050
46	48	3.7053	1.1773	0.1029	1.4600	103.84	2.01	1907.6	0.0413	0.0041
52	54	4.2353	1.0908	0.0991	0.8668	120.59	3.09	1890.9	0.0274	0.0033
56	58	4.5905	0.6612	0.0613	0.5958	132.62	3.92	1878.8	0.0307	0.0043
60	64	5.1150	0.5806	0.0749	0.2828	156.55	7.56	1854.9	0.0201	0.0042
64	68	5.4631	0.3893	0.0510	0.1473	177.50	13.97	1834.0	0.0166	0.0055
68	72	5.8230	0.1587	0.0480	0.0902	193.25	21.96	1818.2	0.0229	0.0135
72	76	6.1939	0.0695	0.0446	0.0644	204.07	29.63	1807.4	0.0343	0.0327
										0
Supported	Pb-210:	1.064 ± 0.03	329 pCi/g	4		I	Cum. Unsu	ıp. Pb-210:	37.0433 p	Ci/cm ²
Number of Supported Samples: 1					I	Unsup. Pb-	-210 Flux:	1.1793 p	Ci/cm² yr	

 Table AE-7:
 ²¹⁰Pb dating for the Lake Little Wilson sediment core.

 Table AE-8:
 ¹³⁷Cs dating for the East Lake Gemini sediment core.

Top of Interval	Bottom of Interval	Detector	Total Pb-210	Total Supported Pb-214	Excess Pb-210 (using Pb-214)	Cs-137
(cm)	(cm)		(pCi/g)	(pCi/g)	(pCi/g)	(pCi/g)
20	22	2	7.90	0.63	7.27	0.78
40	42	2	5.22	1.08	4.14	1.46
60	62	1	3.75	1.02	2.73	2.77

 Table AE-9:
 ¹³⁷Cs dating for the Lake Winona sediment core.

Top of Interval	Bottom of Interval	Detector	Total Pb-210	Total Supported Pb-214	Excess Pb-210 (using Pb-214)	Cs-137
(cm)	(cm)		(pCi/g)	(pCi/g)	(pCi/g)	(pCi/g)
20	22	1				0.77
22	24	1				0.77
24	26	1				0.81
26	28	1				1.17
28	30	1				1.04
30	32	1				1.07
32	34	1				0.81
34	36	1				0.83

Top of Interval	Bottom of Interval	Detector	Total Pb-210	Total Supported Pb-214	Excess Pb-210 (using Pb-214)	Cs-137
(cm)	(cm)		(pCi/g)	(pCi/g)	(pCi/g)	(pCi/g)
14 18 22 24 26 30	16 20 24 26 28 32	2 2 2 2 2 2 2	<u> </u>			0.64 0.62 0.92 1.00 1.63 1.06
34 38	36 40	2 2				0.52 ND

 Table AE-10:
 ¹³⁷Cs dating for the Duluth Harbor sediment core.

 Table AE-11:
 ¹³⁷Cs dating for the Lake Shagawa sediment core.

Top of Interval	Bottom of Interval	Detector	Total Pb-210	Total Supported Pb-214	Excess Pb-210 (using Pb-214)	Cs-137
(cm)	(cm)		(pCi/g)	(pCi/g)	(pCi/g)	(pCi/g)
26	28	1				2.76
28	30	1				3.18
30	32	1				3.41
32	34	1				3.45
34	36	1				3.41
38	40	1				1.37
42	44	1				0.67
46	48	1				0.5
50	52	1				0.29
60	64	1	1.190	1.260		0.30
72	76	1	small peak	1.150		ND
84	88	1	small peak	1.230		ND

 Table AE-12:
 ¹³⁷Cs dating for the Lake Little Wilson sediment core.

Top of Interval	Bottom of Interval	Detector	Total Pb-210	Total Supported Pb-214	Excess Pb-210 (using Pb-214)	Cs-137
(cm)	(cm)		(pCi/g)	(pCi/g)	(pCi/g)	(pCi/g)
60	64	2		0.950		ND
68	72	2		0.670		ND
76	80	2		0.250		ND



Figure AE-1: Total ²¹⁰Pb activity in the Lake St. Croix (A), East Lake Gemini (B), Lake Winona (C), and Duluth Harbor (D) cores. Error bars represent one standard deviation in ²¹⁰Pb activity.



Figure AE-2: Total ²¹⁰Pb activity in the Lake Superior (A), Lake Shagawa (B), and Lake Little Wilson (C) cores. Error bars represent one standard deviation in ²¹⁰Pb activity.



Figure AE-3: Total ¹³⁷Cs activity in the East Lake Gemini (A), Lake Winona (B), Duluth Harbor (C), and Lake Shagawa (D) cores.

Radiometric Dating Interval	Focusing Factor
1996-2008	0.25
1990-1996	0.59
1980-1990	0.57
1970-1980	0.66
1960-1970	0.66
1950-1960	0.85
1940-1950	0.93
1930-1940	1.39
1910-1930	1.26
1890-1910	1.19
1860-1890	1.29
1830-1860	1.61
<1830	1.32

 Table AE-13: Sediment focusing factors for the Lake Pepin sediment core.

Table AE-14: Sediment focusing factors for the Lake St. Croix sediment core.

Radiometric Dating Interval	Focusing Factor
1990-2001	0.74
1980-1990	0.77
1970-1980	0.68
1960-1970	0.59
1950-1960	0.41
1940-1950	0.51
1930-1940	0.56
1910-1930	0.77
1890-1910	0.80
1870-1890	0.99
1850-1870	0.83
<1850	0.53

 Table AE-15: Sediment focusing factors for East Lake Gemini, Lake Winona,

 Duluth Harbor, Lake Superior, Lake Shagawa, and Lake Little Wilson.

Sediment Core	Focusing Factor
East Lake Gemini	3.59
Lake Winona	1.35
Duluth Harbor	1.98
Lake Superior	2.06
Lake Shagawa	2.56
Lake Little Wilson	2.36

Notes on Focusing Factor Calculations:

For non-riverine lake systems in this study (East Lake Gemini, Lake Winona, Duluth Harbor, Lake Superior, Lake Shagawa, Lake Little Wilson), a sediment focusing factor (FF) was determined by taking the ratio of the observed, unsupported ²¹⁰Pb flux in each core to the ²¹⁰Pb flux expected from atmospheric deposition to the lake (Equation 4).

$$FF = \frac{\text{unsupported}^{210} Pb \text{ flux (pCi cm}^{-2} \text{ yr}^{-1})}{0.5 \text{ pCi cm}^{-2} \text{ yr}^{-1}}$$
(4)

where:

0.5 pCi cm⁻² yr⁻¹ = an approximate atmospheric ²¹⁰Pb flux to lakes in MN [119].

FF values greater that 1 signify areas of deposition, or sediment focusing to deeper parts of a lake. FF values less than 1 suggests that the core site may be erosional on a transient basis. Because TCS, CTDs, and PCDDs are hydrophobic and rapidly partition to sediment, one should expect that sediment focusing factors will be representative of analyte focusing in a given lake.

Lake Pepin and Lake St. Croix

In riverine systems, like Lake Pepin and Lake St. Croix, a significant amount of ²¹⁰Pb is deposited upstream of these depositional environments. Therefore, the unsupported ²¹⁰Pb flux recorded in these sediment cores cannot be directly compared to the atmospheric deposition of ²¹⁰Pb at the surface of these lakes.

To mitigate this issue, multiple sediment cores were collected in each lake in previous studies [94,120]. The sedimentation rates for these cores were integrated over the lake area to provide whole-lake sedimentation rates over time. The FF values for Lake Pepin (Table AE-13) and Lake St. Croix (Table AE-14) were determined by taking the ratio of the core-specific sediment flux to the whole-lake sediment flux at different time intervals (Equation 5).

$$FF = \frac{\text{core - specific sediment flux } (g \text{ cm}^{-2} \text{ yr}^{-1})}{\text{whole - lake sediment flux } (g \text{ cm}^{-2} \text{ yr}^{-1})}$$
(5)

Correcting Analyte Accumulation Rates for Sediment Focusing

Analyte accumulation rates presented in Figures 3-5 to 3-12, 3-13, and 3-15 are all normalized to their core-specific focusing factors. This normalization accounts for sediment focusing at each core site and provides insight into whole-lake analyte accumulation rates.

Appendix F

TCS and CTD Derived PCDDs – % By Mass and TEQ

Table AF-1: Percent of TCS and CTD derived PCDDs, by mass and TEQ, of the Lake Pepin PCDD pool.

Approximate	TCS Derived PCDDs	Total PCDDs	%	TCS Derived PCDDs	Total PCDDs	%
Date	(pg/g)	(pg/g)	By Mass	(pg TEQ/g)	(pg TEQ/g)	By TEQ
2009	112.10	652.50	17.18	0.221	5.912	3.74
2005	133.20	534.10	24.94	0.252	5.048	4.99
2002	145.30	621.40	23.38	0.283	6.084	4.65
1997	64.40	352.20	18.29	0.131	3.609	3.63
1992	87.10	295.90	29.44	0.142	2.611	5.44
1989	62.70	428.50	14.63	0.105	4.217	2.49
1983	127.10	1019.70	12.46	0.191	7.681	2.49
1980	105.80	1158.20	9.13	0.158	8.167	1.93
1974	61.00	831.60	7.34	0.088	6.460	1.36
1971	77.10	3130.40	2.46	0.123	24.206	0.51
1964	3.61	700.12	0.52	0.013	5.072	0.25
1961	4.19	1054.90	0.40	0.015	8.066	0.18
1958	4.12	1076.48	0.38	0.012	8.283	0.14
1953	1.77	195.57	0.91	0.004	1.383	0.30
1944	0.83	58.08	1.43	0.001	0.677	0.12
1891	0.79	44.01	1.80	0.001	0.198	0.40

Table AF-2: Percent of TCS and CTD derived PCDDs, by mass and TEQ, of the Lake St. Croix PCDD pool.

Approximate	TCS Derived PCDDs	Total PCDDs	%	TCS Derived PCDDs	Total PCDDs	%
Date	(pg/g)	(pg/g)	By Mass	(pg TEQ/g)	(pg TEQ/g)	By TEQ
2007.5	36.50	567.70	6.43	0.059	3.608	1.64
2002.3	40.21	864.01	4.65	0.069	5.447	1.27
1996.9	33.24	914.04	3.64	0.053	5.406	0.99
1992.8	18.26	781.64	2.34	0.030	4.541	0.65
1989.5	28.92	1008.92	2.87	0.046	6.499	0.71
1983.0	18.49	1881.39	0.98	0.032	4.478	0.71
1979.7	15.62	744.02	2.10	0.021	4.906	0.43
1976.8	14.61	1275.21	1.15	0.020	8.205	0.24
1971.0	13.65	1172.20	1.16	0.020	7.974	0.24
1968.1	18.26	1474.52	1.24	0.030	10.582	0.28
1965.3	13.23	1586.50	0.83	0.024	11.974	0.20
1961.2	3.36	1447.26	0.23	0.008	11.819	0.07
1957.0	1.79	1358.09	0.13	0.004	12.819	0.03
1947.0	1.93	615.43	0.31	0.009	7.218	0.13
1930.0	1.30	82.00	1.59	0.001	1.078	0.12
1865.0	0.96	12.50	7.68	0.001	0.334	0.29

Approximate	TCS Derived PCDDs	Total PCDDs	%	TCS Derived PCDDs	Total PCDDs	%
Date	(pg/g)	(pg/g)	By Mass	(pg TEQ/g)	(pg TEQ/g)	By TEQ
2008.4	282.69	419.59	67.37	0.307	1.904	16.12
2004.9	424.90	715.40	59.39	0.469	3.128	14.99
2002.8	374.40	800.50	46.77	0.414	4.523	9.15
1996.3	911.00	1480.40	61.54	1.010	5.737	17.61
1992.2	413.60	601.00	68.82	0.446	2.375	18.78
1985.6	1192.72	1770.09	67.38	1.298	5.533	23.46
1977.9	960.00	1479.00	64.91	1.050	4.754	22.09
1966.1	1419.00	2405.70	58.98	1.590	8.521	18.66

Table AF-3: Percent of TCS and CTD derived PCDDs, by mass and TEQ, of the East Lake Gemini PCDD pool.

Table AF-4: Percent of TCS and CTD derived PCDDs, by mass and TEQ, of the Lake Winona PCDD pool.

Approximate	TCS Derived PCDDs	Total PCDDs	%	TCS Derived PCDDs	Total PCDDs	%
Date	(pg/g)	(pg/g)	By Mass	(pg TEQ/g)	(pg TEQ/g)	By TEQ
2008	4581.00	8268.80	55.40	6.210	19.381	32.04
2004	6314.00	10968.00	57.57	8.240	25.120	32.80
1998	5932.00	11177.00	53.07	8.020	27.087	29.61
1995	6028.00	11151.00	54.06	8.080	26.776	30.18
1992	5932.00	11338.00	52.32	8.020	27.610	29.05
1981	5197.00	11117.00	46.75	6.970	28.720	24.27
1971	4390.00	10892.00	40.30	6.100	28.707	21.25
1964	3570.00	9728.00	36.70	5.100	27.205	18.75
1948	2077.00	11206.00	18.53	2.770	31.786	8.71
1930	324.60	4855.90	6.68	0.456	16.226	2.81
1915	303.90	1849.80	16.43	0.429	23.774	1.80

Table AF-5: Percent of TCS and CTD derived PCDDs, by mass and TEQ, of the Duluth Harbor PCDD pool.

Approximate	TCS Derived PCDDs	Total PCDDs	%	TCS Derived PCDDs	Total PCDDs	%
Date	(pg/g)	(pg/g)	By Mass	(pg TEQ/g)	(pg TEQ/g)	By TEQ
2009	381.10	4115.50	9.26	0.571	33.149	1.72
2006	413.80	4256.00	9.72	0.628	37.362	1.68
2003	432.70	3299.00	13.12	0.637	31.475	2.02
1995	313.30	2727.00	11.49	0.523	25.155	2.08
1991	187.40	3877.00	4.83	0.434	34.569	1.26
1988	186.00	5545.00	3.35	0.510	48.683	1.05
1981	173.60	4812.00	3.61	0.476	44.275	1.08
1977	209.00	6176.00	3.38	0.470	54.191	0.87
1969	228.00	7637.00	2.99	0.660	71.136	0.93
1965	157.30	7392.00	2.13	0.493	72.714	0.68
1955	22.50	8419.80	0.27	0.117	72.313	0.16
1923	33.60	12585.00	0.27	0.192	140.875	0.14
1893	23.10	17807.20	0.13	0.157	363.404	0.04

Approximate	TCS Derived PCDDs	Total PCDDs	%	TCS Derived PCDDs	Total PCDDs	%
Date	(pg/g)	(pg/g)	By Mass	(pg TEQ/g)	(pg TEQ/g)	By TEQ
2006	37.35	223.70	16.70	0.077	3.062	2.50
1998	66.40	503.30	13.19	0.151	7.602	1.99
1988	26.40	314.20	8.40	0.075	4.501	1.67
1979	56.40	1284.70	4.39	0.177	14.728	1.20
1968	18.90	1652.80	1.14	0.081	21.986	0.37
1958	16.00	1362.90	1.17	0.075	23.661	0.32
1946	22.20	1679.90	1.32	0.105	51.044	0.21
1935	6.75	178.05	3.79	0.031	6.046	0.51
1911	2.63	7.63	34.47	0.008	0.211	3.94
1880	14.10	59.09	23.86	0.058	2.480	2.35

Table AF-6: Percent of TCS and CTD derived PCDDs, by mass and TEQ, of the Lake Superior PCDD pool.

Table AF-7: Percent of TCS and CTD derived PCDDs, by mass and TEQ, of the Lake Shagawa PCDD pool.

Approximate	TCS Derived PCDDs	Total PCDDs	%	TCS Derived PCDDs	Total PCDDs	%
Date	(pg/g)	(pg/g)	By Mass	(pg TEQ/g)	(pg TEQ/g)	By TEQ
2009	517.80	1444.90	35.84	0.678	10.179	6.66
2005	509.70	1550.00	32.88	0.687	12.555	5.47
2000	396.80	1260.00	31.49	0.548	10.173	5.39
1995	136.20	404.00	33.71	0.192	3.385	5.67
1989	148.30	880.40	16.84	0.223	7.022	3.18
1983	171.50	933.20	18.38	0.275	8.290	3.32
1976	85.60	708.10	12.09	0.154	6.471	2.38
1971	96.20	755.60	12.73	0.170	6.929	2.45
1964	59.10	690.40	8.56	0.105	7.313	1.44
1957	36.60	677.60	5.40	0.069	9.184	0.75
1945	22.40	146.40	15.30	0.044	2.814	1.56
1933	24.60	111.50	22.06	0.048	1.842	2.61

Table AF-8: Percent of TCS and CTD derived PCDDs, by mass and TEQ, of the Lake Little Wilson PCDD pool.

Approximate	TCS Derived PCDDs	Total PCDDs	%	TCS Derived PCDDs	Total PCDDs	%
Date	(pg/g)	(pg/g)	By Mass	(pg TEQ/g)	(pg TEQ/g)	By TEQ
2007	18.60	455.21	4.09	0.024	7.429	0.32
1998	35.90	1171.51	3.06	0.053	16.352	0.32
1984	38.03	975.10	3.90	0.056	17.087	0.33
1967	19.94	568.81	3.51	0.028	9.120	0.31
1954	24.30	484.78	5.01	0.036	8.849	0.41
1944	15.00	207.60	7.23	0.015	3.369	0.45
1934	24.34	180.40	13.49	0.027	3.588	0.76
1925	24.00	58.80	40.82	0.024	0.813	2.95
1900	19.00	37.50	50.67	0.019	0.260	7.31
1873	49.00	92.40	53.03	0.049	0.295	16.63