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Dredging Operations and Environmental Research

Identification of Sediment Sources to Calumet River through Geochemical Fingerprinting

David W. Perkey, Mark A. Chappell, Jennifer M. Seiter, and Heidi M. Wadman

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Identification of Sediment Sources to Calumet River through Geochemical Fingerprinting

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Final report

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Under Project A1060-449021-56B673, "Geochemical and Other Tracers for Dredged Material Fate"

Abstract

Geochemical sedimentary markers provide a well-established methodology for fingerprinting various sources of sediments accumulating within a region of concern. To help identify possible sources of contaminated sediment infilling a reach of the Calumet River between the Thomas J. O'Brien lock and Lake Michigan, multiple bottom sediment samples were collected in 2014 from six potential sediment sources. Additionally, geochemical data from historic dredging records spanning a 40-year period were examined to develop a historic geochemical fingerprint for sediments within this reach of the river.

Geochemical measurements and advanced multivariate statistics were used to successfully distinguish between all six potential sediment sources. Calumet River samples showed elevated levels of heavy metals and a geochemical signature that was distinct from the potential source sediments, suggesting that heavy metal contamination was occurring locally within the river. Additionally, multivariate analysis was able to show historic reductions in heavy metal and organic contaminants in the dredge records that likely correlates with implementation of the Clean Water Act. Geochemical fingerprints were successfully used to track changes in sediments both spatially and temporally within the project area. These fingerprints could be used further to identify any future changes in sediments within this reach of the Calumet River.

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Preface

This study was conducted for the Dredging Operations and Environmental Research (DOER) Program under Project Number A1060-449021-56B673, "Geochemical and Other Tracers for Dredged Material Fate." The DOER Program Manager was Dr. Todd Bridges.

The work was performed by the Field Data Collection and Analysis Branch (HN-F) of the Navigation Division (CF), U.S. Army Engineer Research and Development Center, Coastal and Hydraulics Laboratory (ERDC-CHL). At the time of publication, Thad C. Pratt was Chief, CEERD-HN-F; Dr. Jackie Pettway was Chief, CEERD-HN; and W. Jeff Lillycrop, CEERD-HV-T, was the Technical Director for Navigation. Jeffrey R. Eckstein was the Deputy Director of ERDC-CHL, and José E. Sánchez was the Director.

COL Bryan S. Green was the Commander of ERDC, and the Acting Director was Dr. David W. Pittman.

Unit Conversion Factors

Multiply	Ву	To Obtain
cubic inches	1.6387064 E-05	cubic meters
cubic yards	0.7645549	cubic meters
degrees Fahrenheit	(F-32)/1.8	degrees Celsius
feet	0.3048	meters
inches	0.0254	meters
microinches	0.0254	micrometers
miles (US statute)	1,609.347	meters
pounds (mass)	0.45359237	kilograms

1 Introduction

The U.S. Army Corps of Engineers (USACE) dredges more than 200 million yd³ of material annually in order to operate and maintain thousands of miles of navigation channels and hundreds of harbors. Managing and monitoring these dredging operations is a key mission of the USACE. The high costs of dredging are only expected to increase over the near future, highlighting the imperative need to quantify the sources of infilling sediment. Identification of these sediment sources could potentially allow USACE to reduce dredging volumes through engineering and/or regulatory action that minimizes or prevents sedimentation when possible. Geochemical fingerprinting is a well-established method for distinguishing among sediment sources (Collins and Walling 2002; Collins et al. 2010; Papanicolaou et al. 2003; Gireeshkumar et al. 2013). This technique relies on the link between the geochemical properties of the sediments transported into an area of interest (AOI) and those of their sources.

Identifying and establishing distinct geochemical markers for the sources of sediments is the first and most critical step when trying to quantify sediment inputs to a given AOI. These distinct markers may be a unique trace metal, an elemental composition or ratio of elements, the presence of radio or stable isotopes, a sediment/soil mineral, or a combination of many of these parameters. Once the likely sources of infilling sediment have been characterized, then samples are collected from areas where sediment mixing has occurred. The origin of these samples can then be analyzed and essentially fingerprinted to determine the source of the sediment in question.

Objective

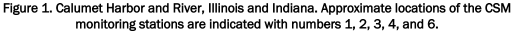
The objective of this report is to showcase how geochemical fingerprinting can be coupled with multivariate statistics as a tool to determine sediment sources impacting an area of the Calumet River between the Thomas J. O'Brien lock and Lake Michigan. This methodology was applied to sediment samples collected from the Calumet River and Harbor, Lake Calumet, and lower Lake Michigan areas as supplementary work to help further identify and constrain sources of sediment and contamination for a conceptual site model being conducted in the region.

Background

The U.S. Army Corps of Engineers, Chicago District (LRC) annually dredges approximately 19,000 m^3 (25,000 yd^3) of contaminated sediment from the Calumet River. An additional 19,000 m³ (25,000 yd³) of less-contaminated sediments are also dredged from Calumet Harbor annually. Currently, both sources of material are placed in a lined confined disposal facility (CDF) located at the mouth of the Calumet River with Lake Michigan. The present CDF is projected to have approximately 5 years of remaining capacity. LRC is developing a dredged material management plan (DMMP) to determine options for the next 20 years of maintenance dredging. The DMMP is exploring opportunities for reducing dredging requirements, beneficial use of the material, and confined placement of contaminated material. LRC is seeking to identify the least-cost DMMP, which could include reduction of sediment loads to the river and harbor. In 2013, at the request of LRC, the U.S. Army Engineer Research and Development Center, Coastal and Hydraulics Laboratory (ERDC-CHL), was asked to develop a conceptual site model (CSM) to identify sediment sources to the Calumet Harbor and River. As part of that CSM, a monitoring plan to identify sediment sources to the Calumet Harbor and River was implemented. The original scope of the plan proposed by ERDC-CHL called for geochemical analysis of potential sediment sources; however, due to funding restrictions, sediment sampling for geochemical analysis was not included in the 2013 study. In 2014, the Dredging Operations and Environmental Research (DOER) Program funded a Research Task (RT) to demonstrate the use of geochemical techniques to identify sediment sources into and out of areas impacted by dredging operations. This DOER Program RT partnered with the LRC on the CSM project to provide geochemical information on sediments infilling the harbor and river channel in an attempt to identify those sediment sources.

Site description

Calumet Harbor is located on the southwest shore of Lake Michigan, west of the Illinois-Indiana state border and approximately 18 km (11 miles) southeast of Chicago. The harbor is protected to the north and northeast by a breakwater with a single gap on the northeast facing wall (Figure 1). Suspended sediment concentrations along the southern Lake Michigan shoreline are typically low; however, during large winter and spring wave events, suspended sediment levels become elevated as a result of the erosion of fine sediments on the southern Lake Michigan coastal shelf (Eadie et al. 1996, 2002; Lou et al. 2000; Beletsky et al. 2003). These storms may produce sediment plumes lasting several days to weeks, which are available for transport into Calumet Harbor.





The Calumet River flows south approximately 11 km (7 miles) from Lake Michigan and Calumet Harbor towards the Thomas J. O'Brien Lock and Dam (Figure 1). Severe rainfall events occasionally result in flow reversal, with river discharge from the lock and dam flowing towards Lake Michigan. This stretch of river is industrially developed with multiple stockpiles of road salt, coal, and other bulk materials located adjacent to the river banks, which are predominantly lined with sheet pile or riprap. A total of nine stormwater and combined sewer outfalls along with two channel outlets were identified as potential sources of sediment to the Calumet River. The two channels were identified as Indian Creek, which enters the river near Turning Basin 3, and Pullman Creek, which enters the system on the western side of Lake Calumet near I-94¹.

Lake Calumet is generally less than 1.2 m (4 ft) deep; however, a deeper legacy channel still exists connecting the southern end of Lake Calumet to the Federal channel. Lake Calumet receives watershed runoff from Pullman Creek, which receives flow from a highly urbanized watershed, composed primarily of freeway. No information exists quantifying the exchange of sediment between Lake Calumet and the Calumet River.

Previous conceptual site model results

From December 2012 through May 2014, ERDC-CHL collected data to construct a CSM to describe the primary sediment sources and the physical processes that deliver sediment to the Calumet system. The CSM was developed from comparisons of environmental forcing conditions and measured turbidity within the system. Primarily through turbidity monitoring and bottom sediment erosion testing, several potential sediment sources and delivery mechanisms were identified. Large wave events on Lake Michigan that frequently occur from late autumn through early spring were identified as transport mechanisms for delivering fine sediments into Calumet Harbor and River. Turbidity responses to these events were limited to the northern section of the river, closest to the harbor and Lake Michigan, and were not transported throughout the entire river. On the inland side of the system, Pullman Creek, which empties into western Lake Calumet, was identified as a primary source of water and sediment into the system during precipitation events. Sediments from Pullman Creek were found to either transport directly into the river or deposit in the shallow waters of Lake Calumet, where they were available for resuspension and transport during strong (>8 m/s) wind events. Again, however, these observed increases in turbidity levels were limited to the stretches of river closest to Lake Calumet and were not transported through the entire system. In contrast, a heavy rain event in April of 2013 resulted in flow reversal of the Calumet River. A large turbidity plume was observed throughout the system for three days. This extreme event was shown to transport a large amount of sediment to the Calumet River and Harbor, though at lower (decadal scale) frequency than Pullman Creek sediment plumes. In addition to transport of sediments in from outside sources, the bottom sediments within the Calumet River

¹ ERDC CSM-TR in progress.

were found to be mobilized as a result of deep-draft vessel passage. Once resuspended, these bottom sediments are redistributed by currents and waves within the system.

In addition to identifying likely sources and transport mechanisms of sediments in the Calumet system, the CSM also identified two potential sources that did not appear to contribute significant sediment loads to the system. Neither monitoring station 3, located at 106th Street near a stretch of unprotected river bank, nor station 4, positioned just downstream of Indian Creek, showed turbidity responses associated with precipitation events. The lack of response of turbidity in these locations throughout the study indicated that neither potential source was likely to be a significant contributor of sediment to the Calumet River.

2 Methods

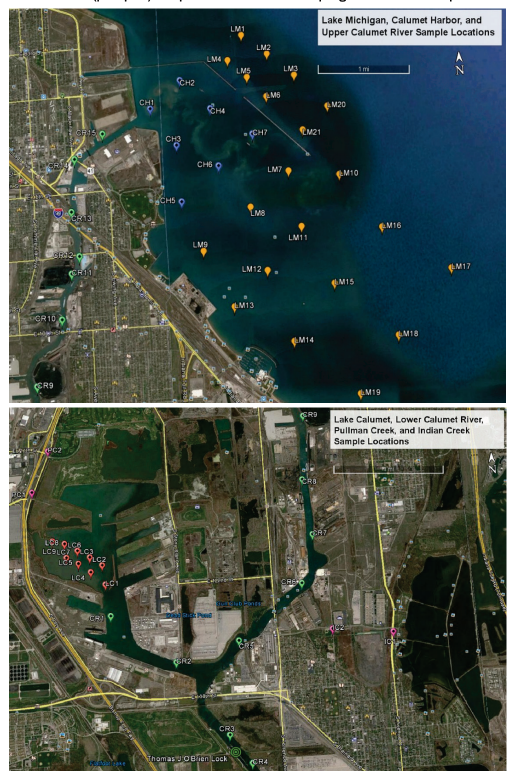
This section describes the field sampling and laboratory methods, along with data analysis methods, used in characterizing the sediments from various sources to the Calumet Harbor and River in the vicinity of Calumet City, IL. Sample collection techniques are presented first, followed by detailed methodology of all the geochemical techniques and multivariate statistics used. Results are presented in the context of the study objective: to identify the sources of sediment to the Calumet system.

Sample collection

To help identify sediment sources to the Calumet system, the project area was divided into six regions composed of Lake Calumet, Calumet River, Calumet Harbor, Lake Michigan, Pullman Creek, and Indian Creek. A total of 55 bottom sediment samples were collected and analyzed to characterize the sediment from these regions. Sample collection occurred 18 through 20 May 2014. Maps of sample locations are found in Figure 2. Table 1 lists their longitude and latitude. No sample was able to be recovered from site LM20.

Samples collected from the Calumet Harbor and River, along with those from Lake Michigan, were obtained with a petite Ponar bottom sampler (Figure 3). Prior to sampling at each location, all equipment contacting sediment samples was decontaminated with Simple Green cleaning solution, distilled water, and site water. Sediment recovered from each Ponar cast was placed in a stainless steel basin and visually examined. Multiple casts were made as needed to collect approximately 1–2 L of sediment. The sediment was then homogenized in the basin and transferred to a sterile whirl-pak bag.

In the shallow water sampling areas of Lake Calumet, Indian Creek, and Pullman Creek, sediments were obtained with a 5 cm (2 in.) diameter push core (Figure 3). Core lengths at these sampling locations typically ranged from 5 to 13 cm (2 to 5 in.). Multiple cores were collected at each location until sufficient material was obtained. As with Ponar sampling, material at each push core site was placed in a stainless steel basin and homogenized prior to being transferred to sterile whirl-pak bags. A new, 2 in. diameter, clear, polyvinyl chloride coring sleeve was utilized at each location and all other equipment contacting the sediment was decontaminated with Simple Green cleaning solution, distilled water, and site water prior to sampling at another location. Figure 2. Locations of samples collected in the northern sample area from Lake Michigan (orange pins), Calumet Harbor (blue pins), and northern Calumet River (green pins) are shown in panel A. Lower Calumet River (green pins), Lake Calumet (red pins), and Pullman and Indian Creek (pink pins) samples in the southern sampling area are found in panel B.



Α

В

Sample Latitude		Longitude	Sample	Latitude	Longitude
LM1*	41°44'38.12"N	87°30'22.33"W	CR1*	41°40'8.60"N	87°35'14.04"W
LM2	41°44'26.66"N	87°30'3.83"W	CR2	41°39'45.54"N	87°34'35.33"W
LM3	41°44'13.83"N	87°29'44.38"W	CR3*	41°39'11.69"N	87°34'5.41"W
LM4	41°44'24.52"N	87°30'33.93"W	CR4	41°38'58.74"N	87°33'52.89"W
LM5	41°44'14.48"N	87°30'20.15"W	CR5*	41°39'53.43"N	87°33'56.55"W
LM6	41°44'3.06"N	87°30'6.50"W	CR6*	41°40'17.88"N	87°33'15.42"W
LM7*	41°43'21.69"N	87°29'53.83"W	CR7	41°40'40.48"N	87°33'6.47"W
LM8*	41°43'3.74"N	87°30'23.90"W	CR8*	41°41'7.20"N	87°33'10.21"W
LM9	41°42'42.20"N	87°31'0.65"W	CR9	41°41'38.46"N	87°33'6.43"W
LM10	41°43'17.82"N	87°29'16.16"W	CR10*	41°42'11.76"N	87°32'46.65"W
LM11	41°42'51.51"N	87°29'47.22"W	CR11	41°42'35.22"N	87°32'38.34"W
LM12	41°42'29.65"N	87°30'14.52"W	CR12*	41°42'44.10"N	87°32'32.18"W
LM13	41°42'12.00"N	87°30'40.48"W	CR13	41°43'7.85"N	87°32'36.80"W
LM14	41°41'51.95"N	87°29'58.59"W	CR14*	41°43'36.05"N	87°32'32.97"W
LM15	41°42'20.42"N	87°29'26.07"W	CR15*	41°43'48.77"N	87°32'11.26"W
LM16	41°42'48.15"N	87°28'47.73"W	LC1*	41°40'23.42"N	87°35'16.54"W
LM17	41°42'23.90"N	87°27'59.39"W	LC2*	41°40'32.52"N	87°35'18.19"W
LM18*	41°41'51.20"N	87°28'42.46"W	LC3	41°40'36.64"N	87°35'25.69"W
LM19	41°41'22.84"N	87°29'13.53"W	LC4*	41°40'29.61"N	87°35'25.34"W
LM20	41°43'55.44"N	87°29'21.03"W	LC5*	41°40'33.99"N	87°35'32.89"W
LM21	41°43'43.37"N	87°29'40.99"W	LC6	41°40'40.10"N	87°35'33.22"W
CH1	41°44'0.72"N	87°31'34.45"W	LC7	41°40'37.12"N	87°35'40.15"W
CH2*	41°44'15.26"N	87°31'10.96"W	LC8	41°40'43.75"N	87°35'41.10"W
CH3*	41°43'39.81"N	87°31'16.13"W	LC9*	41°40'45.38"N	87°35'48.64"W
CH4*	41°43'58.53"N	87°30'49.27"W	IC1	41°39'52.67"N	87°32'21.81"W
CH5	41°43'9.14"N	87°31'14.66"W	IC2	41°39'55.91"N	87°32'58.76"W
CH6*	41°43'27.08"N	87°30'45.65"W	PC1	41°41'9.40"N	87°36'0.78"W
CH7*	41°43'43.06"N	87°30'18.74"W	PC2	41°41'29.09"N	87°35'50.07"W

Table 1. Coordinates of sampling locations in Lake Michigan (LM), Calumet Harbor (CH), Calumet River (CR), Lake Calumet (LC), Indian Creek (IC), and Pullman Creek (PC). Asterisk indicates sample was analyzed for radio isotope activity.

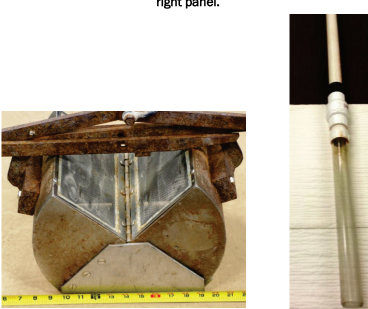


Figure 3. Images of the sampling devices used for sediment collection. A petite Ponar sampler is shown in the left panel, and an image of the 5 cm (2 in.) diameter push corer is shown in the right panel.

After collection, all sample bags were packed into coolers and transported back to ERDC-CHL in Vicksburg, MS. Upon arrival at ERDC-CHL, samples were kept in a walk-in cooler and maintained at a temperature of approximately 4 °C (39 °F).

Grain size analysis

Each sediment sample was homogenized, subsampled (1-2 g), and deflocculated overnight in a solution of sodium metaphosphate (40 g/L) prior to grain size analysis. A Malvern Mastersizer 2000 laser diffraction particle-sizer was used to measure the particle-size distributions of the subsamples. The particle-sizer measures particle size over the range 0.02 to 2000 μ m. Particle size distributions were determined by first removing and sieving (#18 mesh) debris and organic material larger than 1000 μ m. The passing portion of the sample was added to the instrument's reservoir and sonicated for 60 s prior to analysis. The sample was then pumped and recirculated through the optical module, which includes a spatial filter assembly containing a laser diode and laser beam collimator. The diffraction detector assembly contains a custom photodetector array that is used for the measurement of light scattering by the suspended particles. The distribution of grain sizes and median grain sizes was derived from this light scattering measurement. Organic material was not oxidized before grain

size analysis was performed; therefore, grain size distributions include organic material less than 1000 μ m.

Four samples collected from Lake Michigan were composed of coarse material beyond the range of laser diffraction detection limits. Grain size distribution of these samples was obtained through dry sieving according to ASTM D6913-04 (2009) protocols.

Light element analysis

Total inorganic and organic carbon, total nitrogen, and total sulfur, were determined by catalytic combustion using CN628 Carbon/Nitrogen Determinator by LECO Corporation, Inc. Analysis was performed on whole sediment samples and completed in triplicate to evaluate any heterogeneity within samples. Carbon/nitrogen ratios were calculated from the averaged results and determined for both percent and atomic ratios.

Trace metal analysis

Total metal content of all sediment samples were determined through total metal digestion using the USEPA 3050B (USEPA 1986) method.

Bulk mineralogy

Sediment mineralogy was determined through X-ray diffraction (XRD) of the bulk sediment material. Whole sediment samples were dried, sieved, and ground. A subset of 20 of the 55 total surface samples was selected for bulk XRD analysis. Initially, 10 samples were selected from each of the designated areas (Lake Michigan, Calumet Harbor, Calumet River, Lake Calumet, Pullman Creek, and Indian River). Data processing indicated that minimal changes were apparent among all samples. Therefore, an additional 10 samples were prepared, and data were collected. All samples were run on a Panalutical X'Pert Pro Materials Research Diffractometer unit at Co-Ko wavelength, and the data were processed using MDI Jade 2010 software.

Electrical conductivity and pH measurements

Soil pH was conducted using standard methods using 10 g of sediment to 10 mL of deionized (DI) water, being careful to maintain a 1:1, sediment:water ratio. The system was allowed to stand and settle for 10 min before reading the pH with a calibrated Cole-Palmer combination, solid-state pH electrode.

Electrical conductivity (EC) was conducted using standard methods using 10 g of sediment to 10 mL of DI water, being careful to maintain a 1:1, sediment:water ratio. The EC was then measured using a calibrated solidstate conductivity probe.

Radioisotope analysis

A subset of 23 of the 55 total surface samples was identified for radio isotope analysis. Samples (indicated with an asterisk in Table 1) were selected from four of the six project areas. Subsamples were initially selected based on grain size, with preference given to fine-grained (muddy) samples as the higher surface area of fine-grained particles yields substantially better radioisotope results. Accordingly, no samples from Indian or Pullman Creek could be used for isotope analysis due to the coarse grain size of those samples. Gamma activities of 7Be, supported ²³⁴Th, and ¹³⁷Cs were measured by GEL Laboratories LLC. Samples were dried, pulverized, and packed into a 100 cc gamma can (~150-200 g of dried sediment). Samples were then placed into lead-shielded, standard highpurity germanium detectors and counted to achieve a minimum detection count (MDC) above the required detection limit for 137 Cs (0.1 pCi/g; generally approximately 24 hours). If an MDC for activity for 137Cs was not achieved, the gamma spectrum was examined for evidence of activity for the other radioisotopes. If activity was noted, the samples were counted to the MDC, if necessary, for those isotopes. If no activity was noted, counting ceased. All measured activity was calibrated to individual detector efficiency. Radioisotope activity errors were calculated for each sample by GEL Laboratories LLC and are a function of software sensitivity as well as the efficiency of each individual radioisotope detector. Activities were converted from picocuries per gram of sediment (pCi/g) to disintegrations per minute per gram of sediment (dmp/g), as per GEL Laboratories LLC recommended standard, by multiplying the value of pCi by 2.22.

Surface activities of ²¹⁰Pb were determined via alpha spectrometry by measuring the activity of its granddaughter, secular equilibrium isotope ²¹⁰Po. Dried samples were spiked with a known amount of ²⁰⁹Pb and partially digested in 16N HNO₃ and 6N HCl, resulting in the release of ²¹⁰Pb from the fine fraction. Following the methods of Flynn (1968) and Nittrouer et al. (1979), the polonium was leached onto a silver planchette placed in the resulting solution. Alpha activity on the planchette was measured using gas scintillation counting on a standard alpha detector for a minimum of 24 hours.

Historical sediment data

Historic sampling records of dredged material from the Calumet River and Harbor had been maintained in a database by the LRC spanning a 40-year period from 1967 to 2007. The database includes results from various geochemical analyses that had been performed on dredged material from this region. Parameters that have been measured in the past include levels of metals, nitrogen, phosphorous, carbon, and organic contaminants. Prior to the collection of the samples previously described, the LRC database was reviewed to identify possible geochemical parameters that could be helpful in distinguishing Calumet River and Harbor sediments from other regional sediments. To help elucidate this information, principal component analysis was performed on the historical sediment data.

Principal component analysis

When considering methods to geochemically distinguish sediments, it is important to realize that physicochemical characteristics rarely exist independently of each other in soils/sediments but are instead spatially and temporally interrelated. The presence of these covariate relationships among sediment geophysical and geochemical properties complicates distinguishing sources with a single defining property, as is commonly attempted. Therefore, the *formulated* nature of sediment's geochemical characteristics makes them better suited to be distinguished by multivariate approaches. This allows consideration of the entire geochemical characterization matrix instead of arbitrarily selecting from one or a handful of seemingly important properties, which could potentially bias the study. A well-established multivariate technique is principal component analysis or PCA. This method quantifies inherent (and often unseen) relationships existing in samples based on the existing covariance among variables (Esbensen 2010). The combination of PCA with a Partial Least Squares Discriminant Analysis (PLS-DA) and a Soft Independent Models for Classification Analogy (SIMCA) was used to determine if sediments from different geographic areas (e.g., Lakes Calumet and Michigan, Calumet River and Harbor) were geochemically distinguishable.

In essence, PCA clusters samples (e.g., sediments) are based on their statistical *similarity*. These statistical similarities are the calculated scores for each reduced dimension, defined as a principle component (PC). To formally assess the statistical validity of the assigned class memberships for the sediment samples, or groupings, as defined by PCA, the

unsupervised classification method called SIMCA and PLS-DA was utilized. SIMCA allows for the classification of samples based on the relative *distance* among the different PCA models built for each group of sediments. This classification thus allows the sediment samples or groups to be sorted with geochemically similar sediments (Wold 1977). PLS-DA is subsequently performed in order to sharpen the separation between groups of observations by analyzing and rotating PCA components such that a maximum separation among classes (in this case, sediment groupings) is obtained. From this analysis, the variables (e.g., geochemical parameters) that carry the class-separating information are defined. Altogether, the results are then used to infer information about geochemically indistinguishable sediments (e.g., likely sediment sources). The following paragraphs provide a detailed description of the PCA, PLS-DA, and SIMCA analyses applied to the Calumet River system data.

Loadings in PCA are defined as the correlation (r) between two variables (vectors), x and y as

$$r(x,y) = \frac{Cov(x,y)}{S_x S_y} \tag{1}$$

where Cov = covariance between x and y and $S_i =$ standard deviation of the variables. Here, the variables represent the geochemical characterization data used to describe each sample type. If projected in reduced space, based on the direction of the principal axis, then the relationship between the variable covariance and angle of the vectors separating x and y is

$$r(x,y) = \cos\theta = \frac{x^T y}{\|x\| \|y\|}$$
(2)

where x^T represents the transpose of the variable x and ||i|| represents the mean-centered variables. Thus, the correlation of variables x and y to the PCs as well as to each other is indicated graphically by their respective angles to the principal axes. For example, if the angle between the vectors (plotted in reduced space) describing variable x and the principal axis (PC-1) is close to zero, then variable x is well described by PC-1 (because $\cos 0 = 1$). Similarly, if the angle between variable x and PC-1 is close to 180° , then variable x is considered negatively correlated to PC-1 (because $\cos(180^\circ) = -1$). Finally, if the angle between two variables is close to 180° ,

the two variables *x* and *y* are considered anti-correlated. In summary, if the variables are close together geometrically, then they are correlated.

By quantifying the correlations among all the variables in a data matrix, PCA can essentially reduce the dimensionality of large, multivariable datasets to small subsets of orthogonal principal components. The advantage of this technique is that it elucidates latent structure in multivariate data matrices that may not be readily apparent from a univariate comparison of the absolute values within the characterization data. Furthermore, these relationships extend to the samples by which the variables are described through the decomposition of the raw data matrix (*X*) as indicated by the general centered PCA model (Esbensen 2010):

$$X = TP^T + E \tag{3}$$

where T = score matrix, $P^T = \text{accompanying loading matrix (transposed)}$, and $E = \text{the error (nonstructural) or residual matrix in the data isolated from the analysis. Thus, the PC model represents the matrix product of TP^T.$

These relationships are important when comparing the score and loading plots. The two plots are considered complementary and are most revealing when the positions of the objects and variables are studied together. By mentally superimposing the score and loading plots, the same rules described above apply for matching correlated variables with objects. Thus, PCA is a powerful approach that has proven particularly effective in finding relationships among complex soil and sediment data that may not have been apparent from simply qualitatively looking at populations of geochemical data (Mathangwane et al. 2008; Chappell et al. 2011; Steevens et al. 2011; Chappell et al. 2014).

For this project, prior to commencing PCA, the data were preprocessed by mean-centering (around zero) and normalizing by the standard deviation to remove any parametric bias. Characterization data reported as below the reported limit or nondetects were removed from the matrix and treated as missing data. Dimension reduction was carried out via the Nonlinear Iterative Partial Least Squares algorithm (which is appropriate for missing data) with the model allowed to select the optimal number of PCs. In the PCA, soil characterization data were treated as variables while the samples were treated as objects. Variables containing > 10% missing data and poorly contributing to the PCs were eliminated to minimize model uncertainty. PCA models were statistically cross validated using a bootstrapping approach and a Leave-One-Out Method for samples. Note that the radioisotope data were not included in the PCA analysis for this project. Cost restrictions as well as grain size limitations resulted in radioisotope analysis of only 23 of the 55 collected samples. This limited sample population risks potentially introducing spatial bias into the population that would likely impact the PCA results. The radioisotope data were thus analyzed on a stand-alone basis, in the context of the other data, post-PCA analysis.

3 Results and Discussion

Bottom sediments collected from the Calumet River and surrounding area were analyzed to determine likely sources of sediment infilling the river. The results of the data analyses are presented in the following sections. This section will also include general observations as well as interesting contrasts in the data. The reader will be referred to technical appendices for full presentation of the analyzed dataset.

Grain size analysis

Weight percentages of sand, silt, and clay fractions based on the Wentworth scale for the 55 bottom samples collected throughout the study can be found in Appendix A. Table 2 presents a summary table that lists the average sand, silt, and clay content for each region. Overall, samples collected from Lake Michigan were generally found to have the highest percentage of sand with an average sand content of 79% (n=20). The bottom samples collected from Pullman Creek and Indian Creek were similarly sand dominated with average sand contents of 78% (n=2) and 57% (n=2), respectively. Sediments within the protected breakwater of Calumet Harbor had a lower average sand content (41%, n=7) than those sediments located just outside the harbor. The fine portion ($<63 \mu m$) of these sediments was composed mostly of silt, which on average accounted for 50% of the total sample. By comparison, samples collected from within the Calumet River and Lake Calumet were found to be silt dominated, with average silt contents of 58% (n=15) and 61% (n=9), respectively. There was very little textural difference observed between samples collected from these two sampling regions. Average sand content for the river was 28% compared to 24% for Lake Calumet, and average clay contents were within 1% of each other.

Region	Sand	Silt	Clay
Pullman Creek (PC)	78	19	3
Indian Creek (IC)	57	41	2
Lake Calumet (LC)	24	61	15
Calumet River (CR)	28	58	14
Calumet Harbor (CH)	41	50	9
Lake Michigan (LM)	79	19	2

Table 2. Average weight percentage sand, silt, and clay of sediments collected from the six sampling regions.

Light element analysis

Weight percentages of total organic carbon (TOC), total nitrogen (TN), and the atomic organic carbon to nitrogen ratios (OC/N) for the 55 samples collected are presented in Appendix B. Table 3 provides a summary of the results via averages of TOC, TN, and OC/N for each region. Overall, the measured TOC of sediments ranged from 7.96% to below detection limits. Samples collected from the Calumet River (CR) tended to have the highest TOC values with a mean of 3.39% (n=15). These levels of TOC were similar to some of the higher TOC percentages previously reported in the historical dredge material records for this area. Calumet Harbor samples showed the lowest TOC values with a mean of 1.32% (n=7). Note that while Table 3 shows Lake Michigan samples (LM) as having the second highest mean TOC (2.17%, n=17), there were three samples from this group (LM 2-4) that had TOC values below detection and were therefore not included in the reported average. Had values <0.1% been utilized for these three samples, then the resulting mean TOC for the LM samples would have been <2.0%.

Region	TOC	TN	OC/N
Pullman Creek (PC)	1.87	BD*	N/A
Indian Creek (IC)	1.90	0.18	12.31
Lake Calumet (LC)	1.48	0.12	14.39
Calumet River (CR)	3.39	0.20	19.78
Calumet Harbor (CH)	1.32	0.12	12.83
Lake Michigan (LM)	2.17	0.13**	19.51**

Table 3. Average weight percentage TOC, TN, and OC/N for sediments collected from the six sampling regions.

*Indicates measurements below detection thresholds of 0.1%.

**LM mean calculated from only 2 of 20 samples that had TN values above detection threshold.

As with TOC, CR samples tended to have the highest %TN with a mean of 0.2% (n=15). In many instances, the other sediment groupings contained samples with TN values below the detection point of 0.1%. For instance, only 2 LM samples (LM9 & LM13) and none of the Pullman Creek (PC) samples had TN percentages above detection limits. In contrast, all 15 samples from the CR group had measurable levels of TN.

Overall, the atomic ratios calculated for this dataset ranged from 4.33 to 54.63. In instances where TN values were below the detection limit, no OC/N could be calculated. CR samples once again showed the highest mean

OC/N (19.78, n=15) for the study area. As previously described, only two of the twenty LM samples had TN values high enough to allow for the calculation of OC/N. Therefore, the reported mean of 19.51 in Table 3 is not representative of the majority of LM samples collected. The other sediment groupings had mean OC/N ratios that ranged from 12.31 to 14.39.

Trace element analysis

Trace element analysis of the 55 samples showed that there is variability across the system. Given the large amounts of data yielded by this methodology, this report summarizes the data only for common metals of concern. Table 4 presents mean values reported in mg/kg of sediment for each of the six sampling regions for the selected metals arsenic, chromium, copper, lead, and zinc. Complete data of all trace element analysis can be found in Appendix C.

Region	As	Cr	Cu	Pb	Zn
Pullman Creek (PC)	13.85	31.05	32.50	136.50	204.00
Indian Creek (IC)	20.96	97.65	37.75	268.40	287.00
Lake Calumet (LC)	22.51	17.41	20.89	46.82	133.44
Calumet River (CR)	13.85	37.26	60.62	150.05	457.87
Calumet Harbor (CH)	9.91	13.28	18.41	38.90	78.70
Lake Michigan (LM)	7.03	7.22	7.57	17.57	80.44

Table 4. Average As, Cr, Cu, Pb, and Zn concentrations (mg/kg) for the six sampling regions.

Arsenic values ranged from 2.75 to 35.70 mg/kg with higher concentrations typically being found in LC samples, which had a mean value of 22.51 mg/kg (n=9) and lower concentrations associated with LM samples with a mean of 7.03 mg/kg (n=20). Chromium levels ranged from 2.42 to 172.0 mg/kg. The highest (172.0 mg/kg) concentration was associated with a sample from Indian Creek (IC1) which was found to be anomalously high when compared to the rest of the data set. Excluding this sample, the highest concentrations of chromium were typically associated with sediments from the CR region. A similar trend was apparent in the lead (Pb) data. Again, the highest single Pb concentration (441.0 mg/kg) was found to be anomalously high and associated with Indian Creek sediment (IC2). Outside of this sample, elevated Pb concentrations were commonly found to be associated with sediments from the CR region. Calumet River sediments also typically showed the highest concentrations of both copper (Cu) and zinc (Zn) and had the highest mean levels of both at 60.62 mg/kg and 457.87 mg/kg, respectively (n=15). By comparison, Lake Michigan samples showed the lowest mean concentrations of As, Cr, Cu, and Pb (Table 4). The lower trace metal concentrations associated with these samples are likely linked to their sand dominated texture. Overall, the samples with the highest trace metal values were associated with areas of active industry and higher anthropogenic activity (i.e., the Calumet River).

Mineralogy

XRD data did not show significant mineralogical differences between the Calumet River and the potential sediment sources. All samples contained common soil-forming minerals including quartz, iron oxides (hematite), clay minerals (chlorite, kaolinite, and muscovite), feldspars (albite, orthoclase), dolomite, and calcite. However, a mineralogical anomaly was discovered in the Calumet River at sampling site 9. This sampling location is located among industrial areas and slag piles. At this site, XRD results showed the formation of an iron/chromium oxide mineral: chromite (FeCr₂O₄). Chromite is an oxide mineral belonging to the spinel group and is an industrially important mineral for the production of metallic chromium, used as an alloying ingredient in stainless steels.

Below are two examples of the Calumet River system mineralogy as determined by XRD. Figure 4 shows representative patterns from Calumet Harbor, Lake Michigan, Pullman Creek, Lake Calumet, and the Calumet River. Although peak intensity may change from sample to sample, the overall mineralogy does not differ significantly. Arrows indicate major peaks for some of the dominant mineral forms in the samples.

By comparison, Figure 5 shows three XRD spectra from the Calumet River (CR5, CR7, and CR9). Overall, the mineralogy is fairly consistent as demonstrated by the similar XRD spectra. However, there are some subtle changes in the 38-42 2θ region of the spectra, due to the presence of hematite and chromite. Chromite in particular was only found in portions of the Calumet River and not in any of the potential sediment sources. These data further support the idea that anthropogenic activity greatly influences the geochemical nature of the sediments in the river.

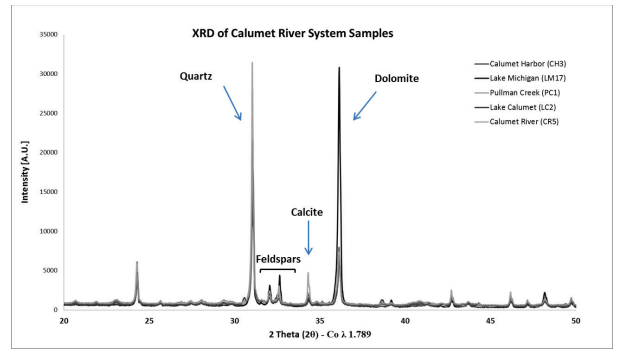
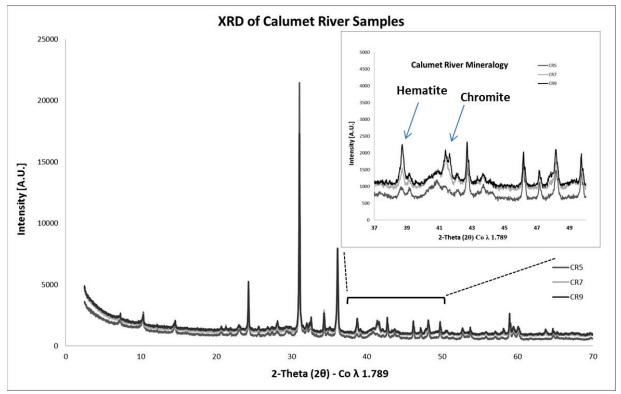


Figure 4. XRD patterns of representative samples from Calumet Harbor, Lake Michigan, Pullman Creek, Lake Calumet, and the Calumet River.

Figure 5. XRD patterns of selected Calumet River samples showing the presence of chromite.



Electrical conductivity (EC) and pH measurements

Sediment pH and EC were measured for all 55 sediment bottom samples collected in May 2014. Individual sediment pH measurements ranged from 6.82 in the Calumet Harbor (CH4) to 8.84 in Lake Michigan (LM21). The average pH for each of the sampling regions was within 0.4 of each other and ranged from 7.46–7.86 (Table 5).

Region	pН	EC
Pullman Creek (PC)	7.65	1.68E+03
Indian Creek (IC)	7.49	2.78E+03
Lake Calumet (LC)	7.82	686.46
Calumet River (CR)	7.46	579.11
Calumet Harbor (CH)	7.63	449.43
Lake Michigan (LM)	7.86	258.86

Table 5. Average pH and electrical conductivity values $(\mu S/cm)$ for the bottom sediments collected from the six sampling regions.

EC ranged from 112.9 μ S/cm in Lake Michigan (LM4) to 4198 μ S/cm in the Indian Creek (IC1). This wide range indicates that there is a high degree of variability throughout the system. The Indian Creek and Pullman Creek samples exhibited higher mean EC values when compared to the rest of the system: 2784 and 1686μ S/cm, respectively (Table 5). These values are $\sim 2.5-4$ times greater than the mean EC value of Lake Calumet sediments, which had the next highest mean EC. EC in sediment is commonly linked to texture (sands exhibit lower EC values, and clays exhibit higher EC values), nutrients and salinity (cations/anions), organic matter, and cation exchange capacity (Grisso el al. 2009). With the exception of the IC and PC samples, mean EC values increased from Lake Michigan inland towards Lake Calumet (Table 5). This trend followed the textural pattern commonly associated with conductivity; sandy LM samples displayed low EC while the fine-dominated CR and LC samples showed higher EC. While the reasons for why the sand-dominated PC and IC samples had such elevated EC values is not known for sure, one possible explanation could be due to higher salt content. Due to their urban environment, both of these shallow water creeks would receive a large amount of road runoff, which could transport a significant amount of road salt into their systems. While salinity was not directly measured for any of the sediments, it was observed that both PC and IC samples had the highest mean concentrations of sodium compared to the other regions

(Appendix C). Sodium is a major component of road salt, and while there are many different forms of salt, the elevated level of sodium in the PC and IC samples offers some support to the idea that they may have had a higher salt content than the other sediments. Complete data of all the pH and EC analyses can be found in Appendix D.

Radioisotope analysis

Significant activity for 7Be, ²³⁴Th, ¹³⁷Cs, and ²¹⁰Pb is generally accepted to be measurable activities reported as disintegrations per minute per gram (dpm/g) of +2 or higher for 7Be,²³⁴Th, and ²¹⁰Pb, or 0.1 or higher for ¹³⁷Cs, after counting and detector efficiency errors are accounted for. Of the 23 samples selected for radioisotope analysis, none showed any significant 7Be activity. One sample (LC1) showed detectable levels of ²³⁴Th, and three sites (CR8, CR15, and CH7) had detectable ¹³⁷Cs levels. In contrast, 10 samples showed significant measurable gamma (γ) and alpha (α) ²¹⁰Pb activity, and a further six samples had significant measurable α , but no measurable y, ²¹⁰Pb activity. Only one sample had measurable y, but not α , ²¹⁰Pb activity: LCO2. This is likely due to a laboratory error. The α ²¹⁰Pb activity measured for this sample was less than 1 dpm/g, which usually indicates that the ²⁰⁹Po isotope used to extract ²¹⁰Pb from the sediment sample did not properly leach onto the silver planchette during standard alpha preparation, resulting in a lack of measurable α radioisotope activity. Overall, of the 23 samples selected for radioisotope analysis, 11 did not have significant gamma activity of any of the selected radioisotopes. Further, of those 11 samples, 5 did not have measurable α ²¹⁰Pb activity. All of the samples without detectable α or y activity were distributed fairly evenly throughout the study site (Figure 6), with no obvious spatial bias. Appendix E provides a table with the complete, processed results for all of the radioisotope analyses.

In terrestrial systems, significant measurable activities of 7Be and ¹³⁷Cs usually indicate runoff from the land surface into a body of water, as well as discharge from a stream or river into a larger aqueous system (Feng et al. 1999; Mullenbach and Nittrouer 2000; Mullenbach et al. 2004; Wilson et al. 2005). Complicating the use of 7Be is that the isotope's activity is strongly dependent on latitude and season. Peak 7Be deposition occurs at latitudes of 30° or lower, and during the early spring and/or late summer (Olsen et al. 1985; Baskaran 1995; Kim et al. 2000). In essence, the higher the latitude, and the further in time from early spring in which the sampling occurs, the lower overall 7Be activities that will be expected in the samples.



Figure 6. Location of radioisotope samples classified by (1) measurable gamma activity of ⁷Be, ¹³⁷Cs, ²³⁴Th, and/or ²¹⁰Pb (white pins); (2) measurable alpha activity only of ²¹⁰Pb (yellow pins); or (3) no measurable radioisotope activity at all (red pins).

Given the timing of the bottom sampling event (late May 2014), as well as the latitude of the Calumet region (41°), it is not surprising that no measurable activities of 7Be were seen at the site. In contrast, sedimentary ¹³⁷Cs is concentrated highest in regions with significant soil runoff. The Calumet River, as well as the adjacent Lake Calumet and surrounding environment, is highly urbanized, with very little exposed riverbank or shoreline. In addition, the previous CSM developed by ERDC in 2014 indicated that an exposed stretch of erodible shoreline on Calumet River in the vicinity of CR10 was not a significant source of turbidity to the main Calumet system. Accordingly, the lack of input of both isotopes resulted in the low (¹³⁷Cs) to nonexistent (7Be) activities measured in this study.

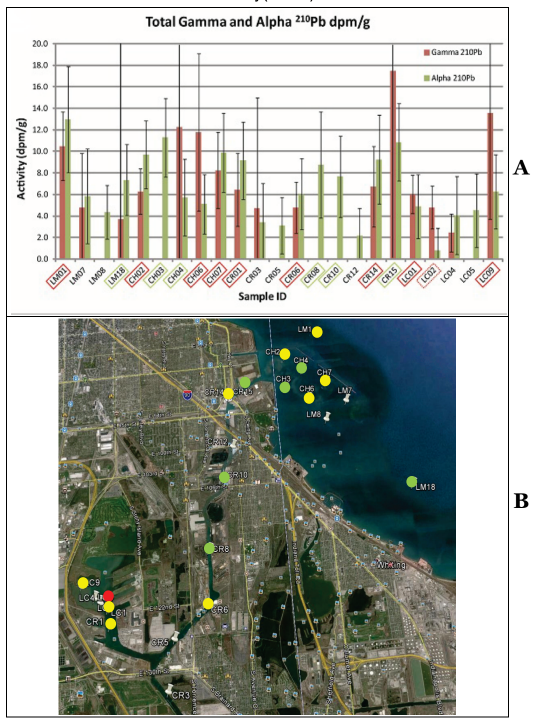
Calumet Harbor consistently showed significant levels of ²¹⁰Pb activities, either in γ , α , or both (Figure 7 [A]). Of the two methods, α -counting of ²¹⁰Pb allows for a higher resolution of small radioisotope activity levels, and generally yields improved detection of lower activities than traditional gamma counting. Two of the four Lake Michigan samples (LM1 & LM18) and both of the upper Calumet River samples (CR15 and CR14) also showed significant ²¹⁰Pb activities that might reflect some resuspension and transport of nearshore Lake Michigan and Calumet Harbor sediments into the upper river. This transport, however, is not extensive as no detectable ²¹⁰Pb activity was seen at the next downstream station, CR12.

In Lake Calumet, the highest activities were measured at LC9, the site closest to Pullman Creek (γ =13.54±9.8 dpm/g; Figure 7 [B]). The remaining activities were confined to the deeper legacy channel running south through the lake towards the Calumet River (Figure 7 [B]). The closest Calumet River sample to Lake Calumet, CR1, also had measurable ²¹⁰Pb activity, but the next sample upriver, CR05, did not. These data indicate that Lake Calumet sediment is not likely transported much farther into the Calumet River than site CR1. Most of the remaining samples in the southern portion of the Calumet River showed lower or no significant ²¹⁰Pb activities, often measurable by alpha spectroscopy only, and likely reflect the limited local runoff from the adjacent banks (Figure 7, [A]).

Given that both ²¹⁰Pb and ²³⁴Th are daughter isotopes of ²³⁸U, which is generally considered to be ubiquitous in terrestrial sediment, one might normally expect to see at least supported levels of ²¹⁰Pb and/or ²³⁴Th in every sediment sample in the Calumet system (Borole et al. 1982; Appleby and Oldfield 1983; Feng et al. 1998; Savoye et al. 2006; Sommerfield 2006; Waples et al. 2006; Jweda et al. 2008; Hancock and Caitcheon 2010). The lack of significant ²³⁴Th and relatively patchy distribution of ²¹⁰Pb activity seen in this region instead suggests two things: (1) sedimentary sources to the system (whether from fluvial, runoff, or wave resuspension) were minimal and isolated spatially; and (2) sediment, once in the system, is not transported in significant enough concentrations to result in an even distribution of ²¹⁰Pb activity throughout the study site. This first point supports the CSM findings of most turbidity responses following either precipitation or wave events being spatially limited. Additionally, while a large precipitation event occurred in April 2014 that resulted in a large turbidity event that was observed throughout the entire system, as the second point above stated, the concentration of sediment in suspension during this event may not have been sufficient enough to result in an even distribution of sediment throughout the region.

Figure 7. (A) Gamma activity of ²¹⁰Pb in all radioisotope samples via gamma (red bars) and alpha (green bars) counting. Error bars are generated via laboratory-reported counting error and detector

efficiency for each individual sample. Red boxes outlining the Sample IDs (*X*-axis) indicate significant gamma and alpha activity, green boxes indicate significant alpha activity, only, and the dashed red box indicates significant gamma activity, only. Negative values, either in activity or in error, are not shown for clarity. (B) Spatial distribution of samples with (1) significant measurable gamma and alpha 210Pb activity (yellow ovals), only alpha activity (green ovals), or gamma-only activity (red oval).



Principal component analysis (PCA)

Historic Calumet dredging data

The historical data set includes geochemical data (e.g., metals, organic compounds), referred to here as variables, on sediments from the Calumet River for over 40 years. The type of data and methods used were inconsistent through the years. Therefore, most of the historical variables contained extensive missing data to exceed the 10% limit. These variables were removed from the analysis. For the entire dataset, spanning from 1967 to 2007, only six characterization variables were usable: ammoniumnitrogen (NH₄-N), Pb, Cr, total phosphorous (TP), Cu, and Zn concentrations. A model was developed containing five PCs with 91% of the variance in the data explained by PC 1-2. Calculated F-residuals found that two of the samples, Cal 75-2 (Mar 1975) and CalR 1002-003 (2002) acted as leverage outliers, statistically *far* from the model center, yet still reside in the same hyperplane described by the PC model.

The bivariate plot (Figure 8) shows that all of these variables were positively loaded on PC-1, with the nutrient-related variables, NH₄-N and TP positively loaded on PC-2 while three of the heavy metals (Cu, Pb, and Zn) were negatively loaded on PC-2. All the variables were significantly loaded on both PC 1-2, but Cr was especially important for PC-1 based on the small angle between this variable and the principal axis. In particular, the explained variance was dominated by PC-1 (78%). Extrapolating from the positions of the variable loadings in Figure 9, it was expected that the scores for the sediment samples containing the highest concentrations of these heavy metal and nutrient solutes (representing the *dirtiest* samples) would be similarly loaded on PC 1-2 while the scores representing the cleanest samples would be anti-correlated to the dirtiest samples. These conclusions are further demonstrated by grouping the samples with respect to their ranges for each variable. For example, Figure 9 shows that the highest sediment concentrations for NH₄-N, Cu, and Zn occur among samples positively loaded in PC-1.

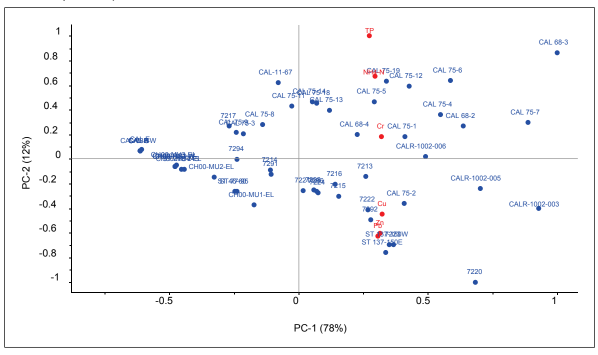


Figure 8. Bivariate plot showing the variable loadings and sample scores (P and T, respectively in Equation 1) for the first two PCs of the historical Calumet sediment characterization data.

Replotting the scores with respect to their sampling time (Figure 10) showed trends with respect to the temporal geochemistry of the sediments. In general, older sediment samples were positively loaded on PC-1 while newer samples were negatively loaded on PC-1, with again, agricultural (NH₄-N, TP) and heavy metal contaminants (Pb, Zn, Cu) generally collapsing into a single cluster highly and negatively loaded on PC-1 (as indicated by the dotted lines). Notable exceptions to these trends occurred with Cal A-E, Cal N, and Cal W samples (collected in 1967–1968) and the three CalR-1002-3,5-6 samples collected in 2002. It is hypothesized that these exceptions arose from particular conditions at the sampling sites. For example, a close visual inspection of the Calumet River maps and historic sampling points indicates that the 2002 samples were collected at positions along the river in proximity to spoil piles located at the river's edge. Thus, it was expected that the sediment geochemistry was impacted by mixture of the spoil material with the sediment such as during erosional (e.g., high rainfall) events.

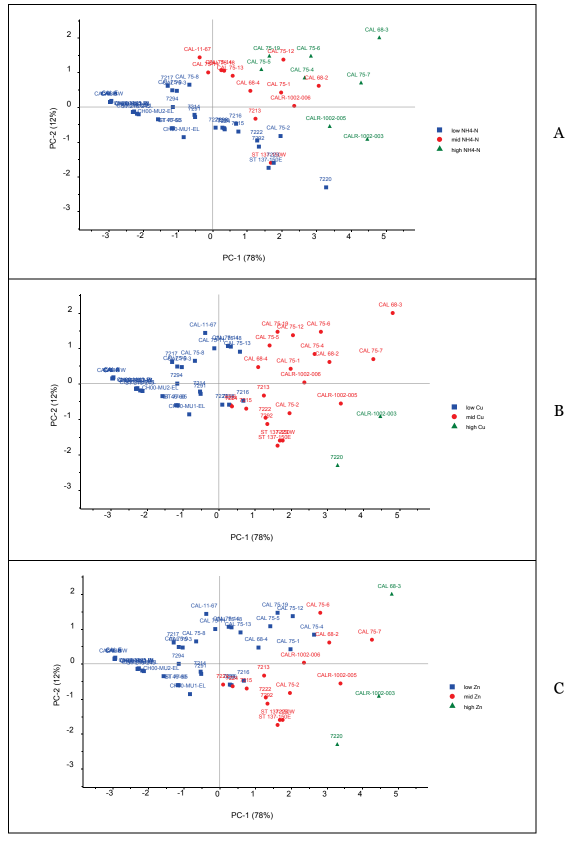


Figure 9. Plots showing the relative concentration ranges for (A) NH₄-N, (B) Cu, and (C) Zn as reported in the historic sediment Calumet data.

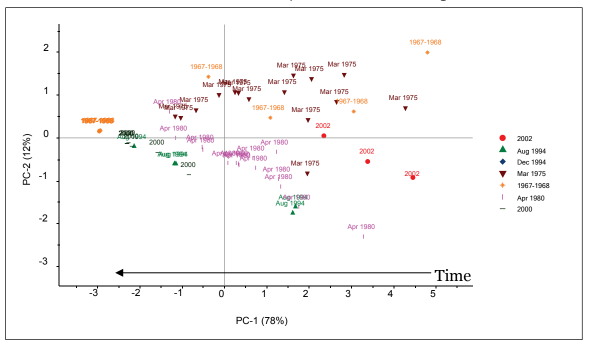
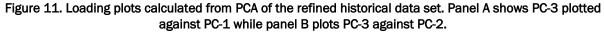


Figure 10. Scores for the historic Calumet samples for PC 1-2 plotted with respect to their sampling time. The dashed lines demonstrate the collapse of the data set to a single cluster with time.

Such clear temporal trends are diagnostic of shifts in the surrounding environment, such as those associated with urban development or unsuspected long-term contaminant leaks. For example, an important temporal shift in sediment geochemical characteristics should be evident in sediments following the passage of substantial environmental legislation, such as U.S. Environmental Protection Act's (USEPA's) Clean Water Act. This hypothesis is borne out by re-examining the data and loosening the constrictions on the potential variable sets that can be used to elucidate these relationships. Narrowing the sample collection times to 1975–2002 allowed for expansion of the analysis to a larger characterization matrix consisting of 13 variables (including Chemical Oxygen Demand [COD], oil and grease, Fe, Mn, Ba, and total potassium and nitrogen [TKN]).

Inclusion of these additional variables allowed for the creation of a 5-PC model explaining 95% of the variance in the data. The model was readily optimized by the removal of two substantial leverage outliers from the December 1994 collections: samples 94-1228 and 94-1219. Figure 11 shows that the variables all load positively in PC-1 (explaining 58% of the variance) similar to the loadings in Figure 8. With the addition of a third PC, Figure 11 (B) shows that PC-2 (explaining 22% of variance) distinguishes the macronutrient and organic-related contaminants/characteristics (negatively loaded on PC-2) from the mineral-relevant characteristics (percent fines)

and heavy metal contaminants (with the exception of Ni). Thus, the Mn, As, Cu, and Zn variables were highly correlated, forming a distinct cluster among these contaminants. Among the macronutrients and organic soluterelated variables, COD, TKN, and TP were closely correlated. Interestingly, oil and grease represented the only variable that could be considered *anticorrelated* with the heavy metal/clay fines samples. This suggests that these variables represent the two extremes of the sample characteristics. In other words, samples high in oil and grease exhibited very different *properties* than sediments only high in heavy metals.



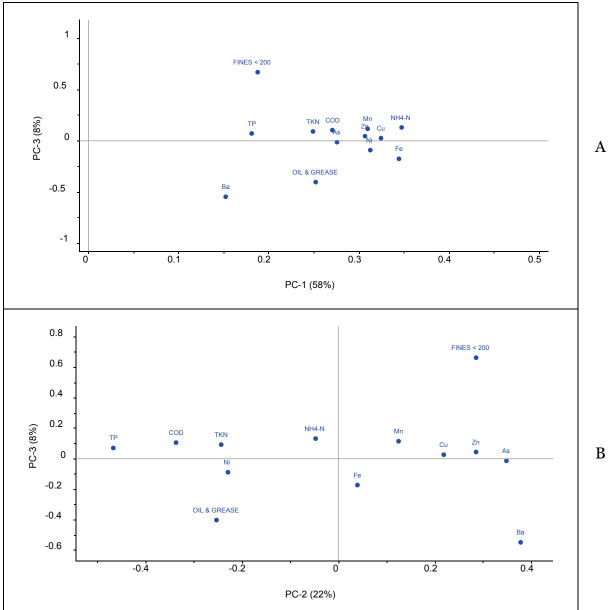
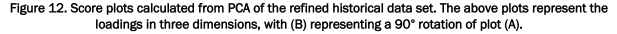
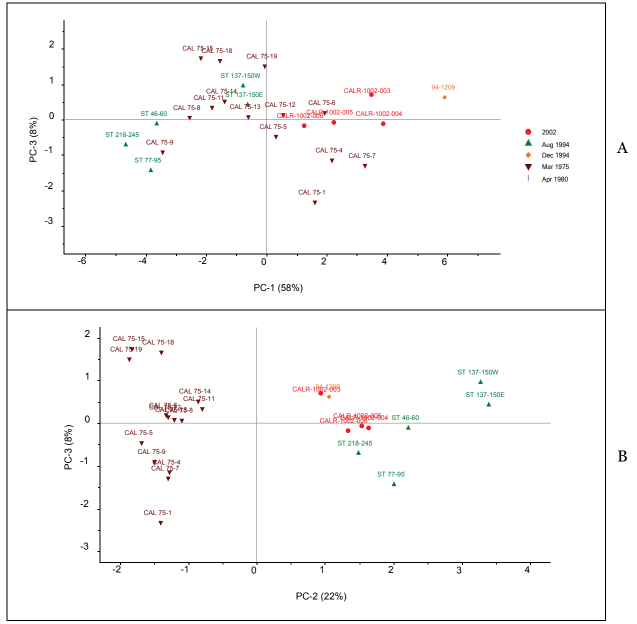


Figure 12 shows the refined sample set for the distinct temporal separation of collections in PC-2. Mentally rotating Figure 12 (A) 90° shows strong clustering of the older March 1975 samples from the remaining collections. These results suggest that the first priority in reducing pollution inputs to the Calumet system involved the mitigation of agricultural and petroleum-based contaminants. With time, this effort was followed up with more detailed efforts to mitigate heavy metal contamination, although some lingering heavy metal contamination seems to exist (as of 2002) at discrete locations along the river system.



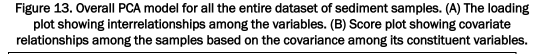


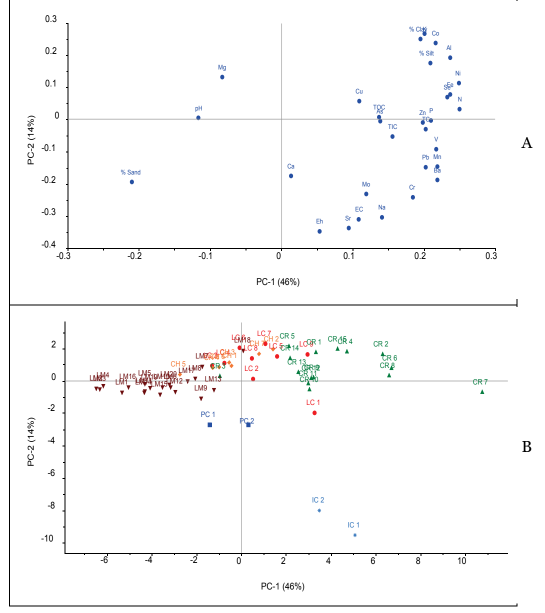
2014 Sediment data

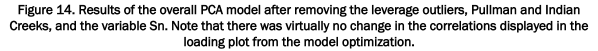
For the sediment samples collected across the Calumet River system, an overall PCA model was found containing six PCs explaining 83% of the variance in the data. This overall PCA model demonstrated strong patterns that there were sensible relationships among the different soils. This suggested that the original groupings (based on sampling location) may serve well to statistically distinguish the sediments. The analysis showed that the Calumet River samples were more correlated with the Lake Calumet samples, while anti-correlated with the Lake Michigan samples. Also, Lake Michigan samples were highly correlated with the Lake Calumet and Calumet Harbor samples, pointing toward their expected geochemical similarities (Figure 13). The loading plot (Figure 13 [A]), showing the total covariance among the characterization data, pointed toward the relationships of the variables to each other, as well as the samples in the score plot (Figure 13 [B]). For example, %sand was correlated with the Lake Michigan samples, while anti-correlated with %clay and %silt, which shared the strongest correlation to Calumet River samples.

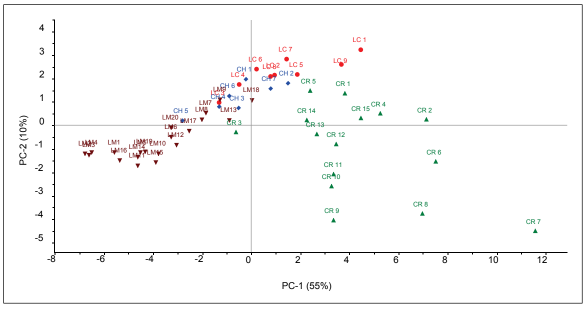
While sensible patterns exist among the different objects and variables, there was substantial overlap in the samples, making it difficult to unambiguously assign them to any single class membership. Part of the reason for this ambiguity was attributed to the two substantial leverage outlier sets, Pullman Creek and Indian Creek. As leverage outliers, Pullman Creek and Indian Creek samples are statistically far from the model center, yet still reside in the same hyperplane described by the PC model. This has the result of stretching out the ellipsoid and thus contributing to the ambiguity in discriminating the appropriate classes for most of the samples. With the PCA model clearly discriminating out the Pullman Creek and Indian Creek samples as distinct classes, the removal of these samples from the overall PCA model provided more clarity for the remaining samples.

The overall PCA model, after the removal of leverage outliers, was further optimized by removing the Sn concentration variable (which made no contribution to any of the PCs). This resulted in a 4-PC model explaining 79% of the variance in the data. The optimized score plot (Figure 14) articulates clearer clustering of samples, yet substantial ambiguity remained as indicated by the overlap of the clusters defining the different classes of samples. Using PLS-DA (Figure 15), it was determined that some of the important variables for overall discriminating samples among the various classes included TOC, Total Carbon (TC), Pb, P, V, Ba, %N, Se, Al, %silt, %clay, Mg, and %sand. The influence of three of these variables (%sand, TC, and Ni) are demonstrated graphically in Figure 16. Here, coarse-textured sediment samples were positively loaded in Factors 1 and 2 while samples with higher TC (including both TOC and Total Inorganic Carbon (TIC)) and heavy metal concentrations were generally negatively loaded on Factor 1.









To formally assess the statistical validity of the assigned class memberships for the sediment samples, the unsupervised classification method called SIMCA was utilized. SIMCA allows for the classification of samples based on the relative *distance* among the different PCA models built for each group of sediments. A sample's distance (D_i) between two classes from the PCA model is calculated as the orthogonal distance from the sample to the different classes previously defined by their principal components as

$$D_{i} = \frac{\frac{1}{K_{q}} \sum_{k=1}^{K_{q}} s_{q}(m,k)^{2} + \frac{1}{K_{m}} \sum_{k=1}^{K_{m}} s_{m}(q,k)^{2}}{\frac{1}{K_{m}} \sum_{k=1}^{K_{m}} s_{m}(m,k)^{2} + \frac{1}{K_{q}} \sum_{k=1}^{K_{q}} s_{q}(q,k)^{2}}$$
(4)

where s_i = the standard deviation for variable k when fitting samples from two different PCA models (e.g., fitting model q onto model m). D_i values close to zero indicate that the two classes (m,q) are virtually identical while values > 1 indicate the two classes are statistically distinguishable (Svante and Michael 1977).

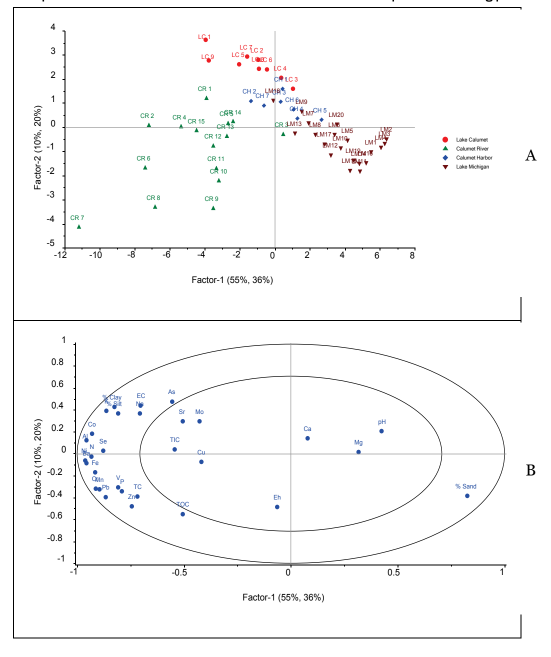
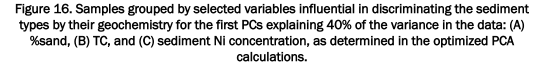
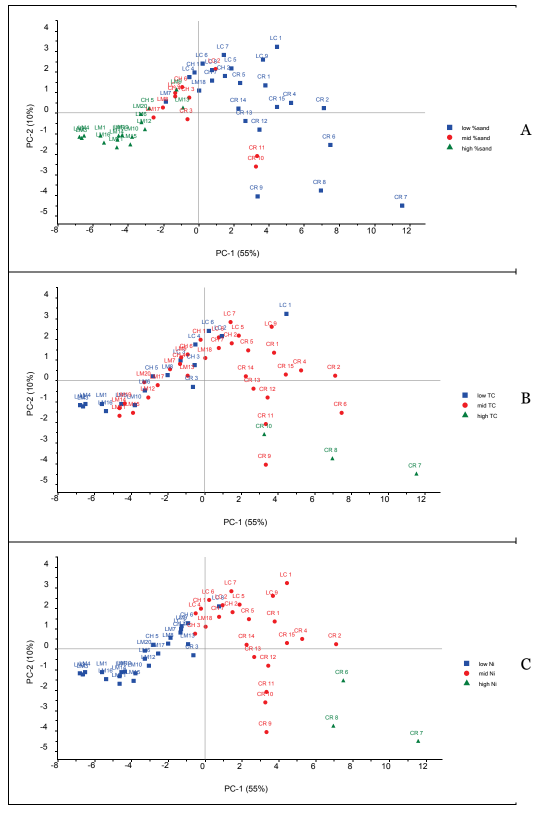


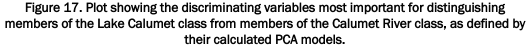
Figure 15. (A) Score and (B) loading plots from PLS-DA of the Calumet River samples. Here, important variables for Factors 1-2 are found within the outer ellipse of the loading plot.

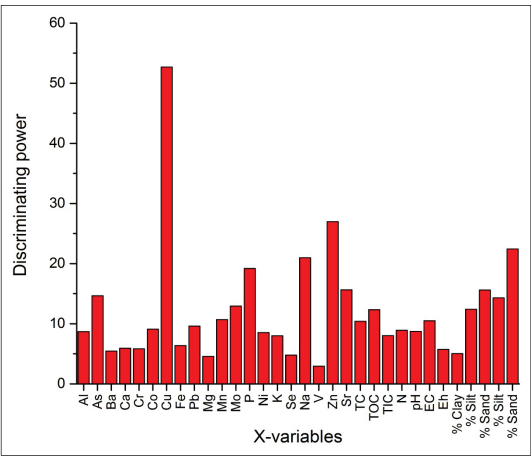




The results of the SIMCA calculations showed that samples from Lake Calumet, Calumet River, Calumet Harbor, and Lake Michigan each formed a unique class. A table displaying these classification results can be found in Appendix F. No classes were assigned to the PC and IC samples as they were previously determined in the overall PCA to exist in their own unique classes.

The SIMCA results also elucidated the most important variables for discriminating between each class. For example, Figure 17 shows that Cu was the most important variable distinguishing members in the Lake Calumet class from members in the Calumet River class. In general, heavy metal concentrations served as the discriminating variables between the Calumet River class and the Lake Calumet and Calumet Harbor classes. Concentration and texture (%clay, silt, sand) were the most important variables discriminating the Lake Calumet class from the Calumet Harbor class.





4 Summary

The multiple geochemical parameters that were measured on the sediment samples collected from the Calumet River system provided a robust dataset that was useful in quantifying and grouping sediments into classes. The multivariate PCA, PLS-DA, and SIMCA analyses were able to distinguish sediment groupings from each other based on these geochemical parameters and to fingerprint a sediment type that correlated with six geospatial locations (Lake Calumet, Calumet River, Calumet Harbor, Lake Michigan, Pullman Creek, and Indian Creek). These fingerprints were then used to evaluate if sediment from these locations was being deposited within the stretch of the Calumet River between Thomas J. O'Brien lock and Lake Michigan.

Within these groupings, sediments from Pullman Creek and Indian Creek were identified as outliers to the rest of the system. The previously conducted CSM showed no evidence of sediment input into the Calumet River coming from Indian Creek even during/after heavy rainfall events. The fact that the Calumet River sediments near the mouth of Indian Creek did not show similar geochemical signatures as Indian Creek samples further support that conclusion. While the CSM did indicate Pullman Creek as being a primary source of sediment into Lake Calumet following large precipitation events, it is important to point out that during those conditions the creek could be transporting a significant portion of fine sediment in suspension. At the time of the sampling, flow in Pullman Creek was low, and the bottom sediments that were collected may not be representative of what is typically transported into Lake Calumet during precipitation events.

While the remaining sediment groupings (Lake Michigan, Calumet Harbor, Lake Calumet, Calumet River) all displayed unique fingerprints, SIMCA results showed that samples from Lake Calumet, Calumet River, and Calumet Harbor shared some similarity that was not seen in the Lake Michigan samples. This result also aligns with the CSM observation of sediments being suspended and transported from both Lake Calumet and Calumet Harbor into the Calumet River during periods of high wind/wave events. Radioisotope activities also indicated limited transport of sediments from Lake Calumet and Calumet Harbor into their respective adjoining areas of the Calumet River. It is suspected that wave events also suspend fine sediments in the shallow, nearshore waters of Lake Michigan. However, previous high wind and wave events that are common in late winter and early spring could have already suspended and transported this material prior to the sampling in May 2014. The predominately sandy texture and the inconsistency in measurable ²¹⁰Pb activities of the LM samples support this conclusion.

Though similarities between Calumet Harbor, Lake Calumet, and Calumet River samples were observed, the PCA and PLS-DA score data clearly showed the Calumet River group to plot primarily in a different region from the other groupings. Heavy metal concentrations were found to be one of the primary causes for this difference. The fact that similar levels of metals were not found in the harbor or Lake Calumet suggests that the source of these metals is within the reach of the Calumet River between Lake Calumet and Calumet Harbor. This suggests that the residence time of bottom sediments in Calumet River is sufficient enough to pick up this additional heavy metal signature. Analysis showed that Cu and Zn are the primary loadings responsible for the geochemical difference between CR sediments and those from LC and CH. Additionally, mineralogy results showed certain areas of the river to contain chromite that was not found elsewhere in the system. While identifying the exact source of this contamination was beyond the scope of this project, the common industrial usage of Cr, Cu, and Zn suggest that anthropogenic activities along this stretch of the river are likely responsible for this signature.

In addition to providing information about likely sources of sediment and contamination into the Calumet River, the geochemical fingerprinting also produced a tool that allows for the monitoring of sediment geochemical characteristics over time. Previous, historical data obtained from dredge material sampling was utilized to showcase how the geochemical fingerprint of the dredged material from the Calumet River changed with time. The PCA, PLS-DA, and SIMCA analyses performed on the samples collected in 2014 produced six classifications of sediment types. Future sediment samples processed with the same methods and models used in this study would allow for the spatial and temporal monitoring of any change in the chemical nature of these sediments.

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Appendix A: Grain Size Data

Table A1. Weight percentage sand, silt, and clay are presented for the bottom samples collected in May 2014. PC=Pullman Creek, IC=Indian Creek, LC=Lake Calumet, CR=Calumet River, CH=Calumet Harbor, LM=Lake Michigan.

Somelo	0/Sand				-	0/0:4	%Clay		
Sample	%Sand	%Silt	%Clay	Sample	%Sand	%Silt	%Clay		
PC1	86	12	2	CH1	21	63	16		
PC2	70	26	4	CH2	12	76	12		
PC Average	78	19	3	CH3	65	28	7		
IC1	57	41	2	CH4	39	53	7		
IC2	56	41	2	CH5	82	15	3		
IC Average	57	41	2	CH6	48	45	7		
LC1	26	58	16	CH7	16	73	11		
LC2	42	48	10	CH Average	41	50	9		
LC3	39	48	13	LM1	93	6	1		
LC4	36	52	13	LM2	100	0	0		
LC5	8	73	18	LM3	98	2	0		
LC6	32	52	16	LM4	99	1	0		
LC7	10	72	18	LM5*	99	-	-		
LC8	17	67	16	LM6*	97	-	-		
LC9	5	82	14	LM7	36	57	7		
LC Average	24	61	15	LM8	50	45	5		
CR1	9	76	15	LM9	73	24	3		
CR2	13	69	19	LM10*	99	-	-		
CR3	62	31	7	LM11	92	8	0		
CR4	16	66	18	LM12	92	7	0		
CR5	29	55	16	LM13	70	26	4		
CR6	19	67	14	LM14	92	8	0		
CR7	19	64	18	LM15	87	13	0		
CR8	30	60	10	LM16	98	2	0		
CR9	35	57	9	LM17	47	49	4		
CR10	48	43	9	LM18	24	61	15		
CR11	38	54	8	LM19	87	13	0		
CR12	29	57	14	LM21*	100	0	0		
CR13	36	51	13	LM Average	79	19	2		
CR14	24	59	17	*Grain size analysis performed by dry sieve method. No clay and silt analysis performed.					
CR15	16	67	17						
CR Average	28	58	14						

Appendix B: Light Element Data

Table B1. Weight percent TOC, TC, TN along with atomic OC/N ratios for the bottom samples collected in May 2014 are presented. PC=Pullman Creek, IC=Indian Creek, LC=Lake Calumet, CR=Calumet River, CH=Calumet Harbor, LM=Lake Michigan, BD=Below Detection, N/A=Not Applicable.

	Harbor, LM=Lake Michigan, BD=Below Detection, N/A=Not Applicable.												
Sample	%TOC	%TC	%TN	OC/N(a)	Sample	%TOC	%TC	%TN	OC/N(a)				
PC1	2.31	5.13	BD	N/A	CH1	2.01	4.87	BD	N/A				
PC2	1.43	4.39	BD	N/A	CH2	1.48	6.04	0.14	12.33				
PC Average	1.87	4.76	BD	N/A	CH3	0.8	3.46	BD	N/A				
IC1	2.1	5.87	0.21	11.67	CH4	1.47	4.96	0.08	21.44				
IC2	1.7	7.54	0.15	13.22	CH5	1.02	3.99	BD	N/A				
IC Average	1.9	6.71	0.18	12.31	CH6	1.17	5.41	0.1	13.65				
LC1	1.46	4.31	0.15	11.36	CH7	1.32	6.11	0.15	10.27				
LC2	1.27	4.15	0.12	12.35	CH Average	1.32	4.97	0.12	12.83				
LC3	1.2	3.77	BD	N/A	LM1	0.9	2.49	BD	N/A				
LC4	1.05	3.6	0.06	20.42	LM2	BD	1.05	BD	N/A				
LC5	1.57	4.89	0.12	15.26	LM3	BD	1.05	BD	N/A				
LC6	1.09	3.72	BD	N/A	LM4	BD	1.34	BD	N/A				
LC7	1.46	4.81	0.11	15.48	LM5	1.05	2.3	BD	N/A				
LC8	1.58	4.86	0.11	16.76	LM6	1	3.6	BD	N/A				
LC9	2.64	5.63	0.18	17.11	LM7	2.09	5.48	BD	N/A				
LC Average	1.48	4.42	0.12	14.39	LM8	2.7	4.54	BD	N/A				
CR1	2.25	5.35	0.19	13.82	LM9	1.25	4.97	0.11	13.26				
CR2	2.76	5.43	0.26	12.38	LM10	2.11	4.27	BD	N/A				
CR3	2.76	3.77	0.11	29.27	LM11	2.9	5.02	BD	N/A				
CR4	2.64	5.34	0.28	11.00	LM12	1.46	5.43	BD	N/A				
CR5	1.55	4.7	0.15	12.06	LM13	3.18	5.87	0.15	24.73				
CR6	3.27	6.53	0.23	16.59	LM14	2.11	4.85	BD	N/A				
CR7	4.57	10.47	0.29	18.39	LM15	3.38	5.65	BD	N/A				
CR8	6.82	11.59	0.23	34.59	LM16	1.49	2.88	BD	N/A				
CR9	7.96	7.98	0.17	54.63	LM17	3.6	5.47	BD	N/A				
CR10	3.11	8.42	0.17	21.34	LM18	2.89	6.08	BD	N/A				
CR11	4.12	6.21	0.14	34.33	LM19	2.28	5.12	BD	N/A				
CR12	2.55	5.78	0.14	21.25	LM21	2.57	5.29	BD	N/A				
CR13	0.89	6.13	0.24	4.33	LM Average	2.17	4.14	0.13	19.51				
CR14	3.04	5.97	0.2	17.73	BD = Below Detection limit, 0.1%								
CR15	2.51	7.68	0.26	11.26	N/A= Not Applicable								
CR Average	3.39	6.76	0.2	19.78									

Appendix C: Trace Element Data

Table C1. Total metal concentrations (mg/kg) of aluminum, arsenic, barium, cadmium,
calcium, chromium, cobalt, and copper for the bottom sediment samples collected in May
2014. PC=Pullman Creek, IC=Indian Creek, LC=Lake Calumet, CR=Calumet River.

Sample	AI	As	Ba	Cd	Ca	Cr	Со	Cu
PC1	3580	9.4	63.9	1.12	40300	19.7	3.77	28.7
PC2	5210	18.3	102	2	65100	42.4	3.93	36.3
PC Average	4395	13.85	82.95	1.56	52700	31.05	3.85	32.5
IC1	3860	8.71	91.7	1.08	69200	172	3.64	25.5
IC2	1910	33.2	145	1.81	160000	23.3	3.19	50
IC Average	2885	20.96	118.35	1.45	114600	97.65	3.42	37.75
LC1	7250	16.3	53.5	0.42	96900	36.4	8.13	18.9
LC2	5210	35.7	32.9	0.45	68100	13	7.75	18.8
LC3	3580	20.6	16.8	0.64	41100	8.21	7.19	15
LC4	6130	20.1	30.1	0.34	57800	12	8.55	14.3
LC5	6640	26.3	31.4	0.69	56700	15.7	9	23.9
LC6	7510	14	34.6	0.46	48900	13.2	8.42	16.9
LC7	6950	20.9	33.3	0.7	53700	17.1	9.13	23.2
LC8	5470	22.8	30.9	0.9	47900	16.6	7.46	22.6
LC9	8500	25.9	53.8	1.41	52900	24.5	10.4	34.4
LC Average	6360	22.51	35.26	0.67	58222	17.41	8.45	20.89
CR1	9900	17.2	54.2	1.08	57400	27.4	9.99	31.5
CR2	11900	19.3	81.7	1.48	51200	52.1	11.7	42.2
CR3	5010	7.82	34.3	0.57	53000	18.9	6.81	21.6
CR4	11000	12.7	74.7	1.24	51000	35.6	9.98	38.8
CR5	7840	9.39	50.9	0.69	61000	22.1	8.25	328
CR6	10300	19.1	88.7	2.26	56700	56.4	12.3	60.1
CR7	12300	19.2	88.5	2.41	63100	79.6	12.8	69.2
CR8	8450	15	66.1	2.01	40100	54.2	9.67	64.7
CR9	5330	13.1	44.2	1.49	36500	40.7	7.93	52.8
CR10	7930	10.1	51.7	1.03	49500	43.6	8.89	36.3
CR11	7560	13.6	61.8	1.06	54800	32.8	9.23	37.6
CR12	8000	12.6	76.1	0.91	54100	26.2	8.81	31.4
CR13	7470	12.2	59.2	0.74	50100	22.2	9.28	36.7
CR14	6730	13.3	48.8	0.52	55600	18.8	8.69	28.8
CR15	10200	13.1	64.9	0.71	59500	28.3	9.6	29.6
CR Average	8661	13.85	63.05	1.21	52907	37.26	9.6	60.62

Sample	AI	As	Ba	Cd	Ca	Cr	Co	Cu
•			-					
CH1	5250	12.7	23.2	0.18	109000	11.7	6.57	19
CH2	8270	9.01	40.3	0.52	83100	19.1	7.93	30.5
CH3	6150	14	28.7	0.24	44100	12.1	9.78	21
CH4	4200	7.65	21.1	0.38	74000	10.8	5.57	12.8
CH5	3120	9.86	16.4	0.19	52600	8.18	6.14	12.1
CH6	4940	7.77	27.9	0.38	79500	13.8	5.66	13.9
CH7	7450	8.39	36.6	0.42	84700	17.3	7.41	19.6
CH Average	5626	9.91	27.74	0.33	75286	13.28	7.01	18.41
LM1	1290	3.81	6.67	0.13	59000	4.25	2.78	2.97
LM2	901	4.52	5.27	BD	19400	2.63	2.5	2.12
LM3	920	3.57	5.12	BD	25100	2.97	2.26	2.02
LM4	761	2.75	5.31	BD	23300	2.42	1.81	2.02
LM5	2460	12.6	14.3	0.19	30800	5.6	5.13	10
LM6	2400	16.6	17.1	1.07	77600	6.21	5.53	11.6
LM7	3700	5.38	20.1	0.31	72300	11.2	4.9	12.6
LM8	3150	4.65	20.2	0.42	99800	13.9	4.64	9.8
LM9	3500	6.14	27.8	0.32	97300	11.7	5.75	9.15
LM10	2400	16.1	14.8	0.11	59400	7.35	5.43	6.09
LM11	1210	3.18	6.8	0.23	75600	4.65	2.8	2.84
LM12	3060	5.06	20.1	0.3	76300	10.9	3.83	8.4
LM13	4290	7.51	30.3	0.3	90500	13.3	5.51	11.6
LM14	1420	3.69	8.54	0.13	79100	5.41	3.07	3.83
LM15	1520	3.37	8.3	0.15	94300	6.15	3.11	4.03
LM16	944	3.15	5.35	BD	50000	4.54	2.52	2.89
LM17	2910	5.27	16.5	0.26	102000	9.11	3.98	10.7
LM18	5690	9.55	24.1	0.24	62400	10.8	9.52	23.5
LM19	1440	3.62	8.14	0.14	77200	5.46	3.32	5.26
LM21	2170	20.1	23.4	0.15	78900	5.9	6.79	9.93
LM Average	2307	7.03	14.41	0.28	67515	7.22	4.26	7.57

Table C2. Total metal concentrations (mg/kg) of aluminum, arsenic, barium, cadmium, calcium, chromium, cobalt, and copper for the bottom sediment samples collected in May 2014. CH=Calumet Harbor, LM=Lake Michigan, BD=Below Detection.

Sample	Fe	Pb	Mg	Mn	Мо	Р	Ni	К
PC1	9250	118	16700	348	1.95	408	11.2	562
PC2	19600	155	28200	743	3.58	545	15.5	777
PC Average	14425	136.5	22450	545.5	2.77	476.5	13.35	669.5
IC1	27600	95.8	12700	3410	15.7	644	19.9	450
IC2	10600	441	10600	500	30	251	15.8	379
IC Average	19100	268.4	11650	1955	22.85	447.5	17.85	414.5
LC1	39800	24.9	20300	1230	39.9	421	25.7	1970
LC2	25400	33.3	23600	569	7.57	420	22.5	1680
LC3	15600	17.8	21600	453	3.85	322	15.4	993
LC4	17600	31.6	30000	462	3.67	294	20.5	1700
LC5	23600	52.7	28700	592	5.32	344	23.5	1870
LC6	20100	30.2	24900	514	3.53	264	21.9	2020
LC7	21900	51.5	28600	586	4.56	310	24	1960
LC8	18500	56.4	23800	502	3.02	330	19.2	1460
LC9	24900	123	27200	654	4.69	337	27.6	2200
LC Average	23044	46.82	25411	618	8.46	338	22.26	1761
CR1	26200	82.9	26900	922	3.04	631	27.2	2550
CR2	37100	174	22400	1620	3.23	869	32.9	2690
CR3	17700	57.7	24600	609	1.95	509	18.5	1170
CR4	30500	122	23900	1040	2.55	1020	29.5	2610
CR5	23600	132	25200	634	3.31	413	25.6	2010
CR6	42200	263	23900	2030	4.68	1030	35.3	1990
CR7	62300	308	22600	2780	9.95	1400	49.8	2140
CR8	50900	239	18400	1470	7.77	696	37.6	1700
CR9	46300	168	15800	1120	6.28	447	29.9	1010
CR10	33400	120	21600	1340	5.32	415	33.9	1280
CR11	39800	149	24200	1430	6.28	440	33.3	1380
CR12	34300	148	22300	1110	6.38	539	27.3	1750
CR13	32300	131	24800	1040	5.32	375	25.9	1830
CR14	38800	69.2	27200	876	7.04	392	29.8	1680
CR15	52900	86.9	27100	1410	3.43	518	24.4	2310
CR Average	37887	150.05	23393	1295.4	5.1	646.27	30.73	1873

Table C3. Total metal concentrations (mg/kg) of iron, lead, magnesium, manganese, molybdenum, phosphorous, nickel, and potassium for the bottom sediment samples collected in May 2014. PC=Pullman Creek, IC=Indian Creek, LC=Lake Calumet, CR=Calumet River.

Sample	Fe	Pb	Mg	Mn	Мо	Р	Ni	K
CH1	25700	83.6	49700	655	5.57	242	21.3	1380
CH2	22300	60	39800	639	2.26	413	21.4	2100
CH3	25600	26.8	22900	384	9.88	259	24.5	1890
CH4	14100	23.4	36000	446	2.54	298	15.4	1150
CH5	18700	16.5	27400	324	5.53	226	15	951
CH6	16800	30.1	36200	459	1.54	319	15.4	1270
CH7	18800	31.9	41300	570	1.99	396	20.2	1960
CH Average	20286	38.9	36186	496.71	4.19	307.57	19.03	1529
LM1	6220	7.36	32700	261	0.59	239	6.73	340
LM2	5050	3.56	11100	122	0.42	101	4.19	214
LM3	4480	6.95	14100	141	0.35	184	4.25	226
LM4	4040	3.67	12000	120	0.43	126	3.97	187
LM5	12400	28.5	15200	257	2.63	143	11.6	765
LM6	14700	13.2	37000	436	3.1	226	14.1	699
LM7	12800	37.3	36000	403	1.35	298	13.6	983
LM8	14800	25.1	48300	476	1.07	359	14.5	771
LM9	14400	20.7	31100	397	1.42	393	14.6	914
LM10	15500	10.4	31100	378	1.39	228	11.5	467
LM11	8110	6.66	42800	275	0.33	421	7.89	269
LM12	10700	20.7	37200	378	0.78	312	11.2	824
LM13	13900	23.6	36100	427	1.08	417	15.4	1100
LM14	7880	9.5	44600	314	0.36	302	8.6	310
LM15	8630	8.61	53000	336	0.44	442	10.3	346
LM16	7010	7.11	28600	236	0.33	294	6.43	198
LM17	9760	17.7	53600	460	1.12	306	13.2	777
LM18	18700	55.5	33800	408	7.12	240	26	1840
LM19	7120	32.7	43000	308	0.42	331	8.79	333
LM21	17700	12.5	40300	449	3.8	236	15.6	697
LM Average	10695	17.57	34080	329.1	1.43	279.9	11.12	613

Table C4. Total metal concentrations (mg/kg) of iron, lead, magnesium, manganese,
molybdenum, phosphorous, nickel, and potassium for the bottom sediment samples collected
in May 2014. CH=Calumet Harbor. LM=Lake Michigan

Sample Se Na Ag TI Sn V Zn Sr									
Sample	Se	Na	Ag	TI	Sn		Zn	Sr	
PC1	0.53	536	BD	0.29	7.05	10.5	158	35.4	
PC2	0.67	702	0.11	0.32	7.7	15	250	53.5	
PC Average	0.6	619	0.11	0.31	7.38	12.75	204	44.45	
IC1	0.74	1190	BD	BD	3.63	38.7	157	94.6	
IC2	1.14	519	0.21	0.2	10.8	12.6	417	225	
IC Average	0.94	854.5	0.21	0.2	7.22	25.65	287	159.8	
LC1	0.79	639	BD	0.47	0.14	35	72.6	84.6	
LC2	0.71	352	BD	0.53	0.14	14.9	100	42.5	
LC3	0.72	182	BD	0.55	0.72	9.87	113	18.8	
LC4	0.33	250	BD	0.54	0.19	13.7	92.3	26.7	
LC5	0.97	279	BD	0.71	0.26	14.9	160	26.8	
LC6	0.8	258	BD	0.59	0.35	14	89.1	24.2	
LC7	0.8	265	BD	0.75	3.14	14	161	25.5	
LC8	0.88	350	BD	0.65	0.87	12.6	171	24.7	
LC9	1.03	472	BD	1.01	1.25	16.3	242	29.2	
LC Average	0.78	338.56	BD	0.64	0.78	16.14	133.44	33.67	
CR1	1	295	0.11	0.59	0.45	21	276	31	
CR2	1.67	371	0.28	0.49	0.92	27.8	543	31.9	
CR3	0.63	232	0.12	0.31	0.5	12.7	197	23.5	
CR4	1.26	357	0.3	0.47	0.78	21.6	379	29.6	
CR5	0.74	295	0.16	0.55	4.08	15	281	35.3	
CR6	1.23	374	0.59	0.51	2.13	25.1	866	37.8	
CR7	1.39	375	0.78	0.57	3.05	48.5	1020	41.3	
CR8	1.12	274	0.83	0.59	1.65	28.9	1040	27	
CR9	0.91	165	0.4	0.54	2.07	22.1	692	23.9	
CR10	0.63	240	0.27	0.58	3.7	30.5	364	28.3	
CR11	0.82	262	0.23	0.74	1.7	31	348	30.1	
CR12	1.11	279	0.11	0.75	0.51	22.1	248	35.6	
CR13	0.91	228	BD	0.63	0.4	19.2	229	27.3	
CR14	0.82	202	0.12	0.71	0.33	16.4	155	26.4	
CR15	0.94	240	0.12	0.49	0.53	23.8	230	30.9	
CR Average	1.01	279.27	0.32	0.57	1.52	24.38	457.87	30.66	

Table C5. Total metal concentrations (mg/kg) of selenium, sodium, silver, thallium, tin, Vanadium, zinc, and strontium for the bottom sediment samples collected in May 2014. PC=Pullman Creek, IC=Indian Creek, LC=Lake Calumet, CR=Calumet River, BD=Below Detection.

CH=Calumet Harbor, LM=Lake Michigan, BD=Below Detection.											
Sample	Se	Na	Ag	TI	Sn	V	Zn	Sr			
CH1	0.42	196	BD	0.61	2.75	12.3	48.4	37.1			
CH2	1.08	204	BD	0.4	0.98	17.6	101	32.4			
CH3	0.94	155	BD	0.93	0.17	16.3	91.1	21.4			
CH4	0.65	169	BD	0.31	0.19	11.7	67.2	27			
CH5	0.57	139	BD	0.35	0.12	11.6	69.4	19.1			
CH6	0.96	221	BD	0.23	0.52	14.4	92.1	32.7			
CH7	0.88	198	BD	0.33	0.24	16.6	81.7	31.6			
CH Average	0.79	183.14	BD	0.45	0.71	14.36	78.7	28.76			
LM1	0.18	123	BD	BD	0.22	8.49	32.6	17			
LM2	0.21	79.8	BD	BD	BD	5.01	13.7	8.68			
LM3	0.22	76.9	BD	BD	0.41	4.62	22.6	10.8			
LM4	0.19	75.3	BD	BD	BD	4.4	10.7	11.3			
LM5	0.31	95.5	1.19	0.25	0.92	8.72	50.6	13			
LM6	0.47	194	BD	0.23	0.46	11.4	592	30.7			
LM7	0.65	161	BD	0.21	1.13	12	66.5	26.3			
LM8	0.66	244	BD	0.14	1.73	16	112	34.2			
LM9	0.5	350	BD	0.16	0.6	12.2	111	51.4			
LM10	0.55	164	BD	0.1	0.12	10.9	43.8	17.7			
LM11	0.38	145	BD	BD	0.2	18.4	51.6	19.4			
LM12	0.62	161	BD	0.11	1.44	11.4	73.3	27.7			
LM13	0.69	311	BD	0.16	0.66	13.4	93.1	42.4			
LM14	0.38	162	0.91	BD	13.2	14.6	43.3	20.6			
LM15	0.74	189	BD	BD	0.39	21.5	46.7	23.3			
LM16	0.33	100	BD	BD	0.51	11.8	29.6	14.5			
LM17	0.57	204	BD	0.11	0.69	12.6	63.1	27.6			
LM18	0.66	166	BD	0.79	2.24	14.5	57.5	21.6			
LM19	0.39	155	BD	BD	9.97	14.3	45	20.3			
LM21	0.54	179	BD	0.31	0.22	9.67	50	31.3			
LM Average	0.46	166.78	1.05	0.23	1.95	11.8	80.44	23.49			

Table C6. Total metal concentrations (mg/kg) of selenium, sodium, silver, thallium, tin, Vanadium, zinc, and strontium for the bottom sediment samples collected in May 2014. CH=Calumet Harbor, LM=Lake Michigan, BD=Below Detection.

Appendix D: Electrical Conductivity (EC) and pH Data

Sample	рН	EC	Sample	рН	EC
PC1	7.59	2072.00	CH1	7.96	604.00
PC2	7.70	1300.00	CH2	7.75	419.70
PC Average	7.65	1686.00	СНЗ	7.86	454.10
IC1	7.53	4198.00	CH4	6.82	472.80
IC2	7.45	1370.00	CH5	7.58	444.50
IC Average	7.49	2784.00	CH6	7.81	401.80
LC1	7.89	575.70	CH7	7.64	349.10
LC2	7.82	662.60	CH Average	7.63	449.43
LC3	7.84	582.90	LM1	7.92	187.70
LC4	7.78	619.10	LM2	8.56	141.60
LC5	7.63	781.70	LM3	8.58	138.90
LC6	7.93	670.00	LM4	7.39	112.90
LC7	7.90	608.30	LM5	7.64	254.00
LC8	7.85	768.40	LM6	7.92	179.70
LC9	7.77	909.40	LM7	7.65	318.70
LC Average	7.82	686.46	LM8	7.75	322.90
CR1	7.43	812.50	LM9	7.65	815.60
CR2	7.55	855.20	LM10	7.68	130.80
CR3	7.47	803.10	LM11	7.94	118.20
CR4	7.42	740.30	LM12	7.54	316.20
CR5	7.57	401.70	LM13	7.55	645.10
CR6	7.26	572.50	LM14	7.83	126.10
CR7	7.50	621.90	LM15	8.30	168.30
CR8	7.57	611.80	LM16	7.23	156.40
CR9	7.12	542.00	LM17	7.76	309.00
CR10	7.63	421.80	LM18	7.86	452.40
CR11	7.65	525.10	LM19	7.64	167.20
CR12	7.31	478.30	LM21	8.84	115.50
CR13	7.64	409.80	LM Average	7.86	258.86
CR14	7.47	524.70			
CR15	7.36	366.00			
CR Average	7.46	579.11			

Table D1. pH and EC (μ S/cm) data for the bottom sediments collected in May 2014. PC=Pullman Creek, IC=Indian Creek, LC=Lake Calumet, CR=Calumet River, CH=Calumet Harbor, LM=Lake Michigan.

Appendix E: Radioisotope Data

							Gamma	²¹⁰ Pb	Alpha ²	¹⁰ Pb -
	⁷ Be - dp	om/g	¹³⁷ Cs - 0	dpm/g	²³⁴ Th -	dpm/g	- dpm/g		dpm/g	
ID	⁷ Be	(+/-)	¹³⁷ Cs	(+/-)	²³⁴ Th	(+/-)	²¹⁰ Pb	(+/-)	²¹⁰ Pb	(+/-)
LM1	0.53	0.69	-0.01	0.06	-1.07	3.17	10.50	3.17	12.99	4.93
LM7	0.61	0.81	0.26	0.17	2.51	2.71	4.77	5.04	5.82	4.40
LM8	1.23	1.53	0.14	0.10	6.33	7.22	0.00	41.0	4.35	2.51
LM18	2.18	1.23	0.01	0.10	2.86	6.04	3.73	14.3	7.35	3.29
CH2	-0.23	1.49	0.00	0.23	2.26	2.09	6.26	2.11	9.72	3.15
CH3	0.08	1.19	0.07	0.11	3.09	5.19	-5.55	6.97	11.28	3.66
CH4	0.27	0.97	0.16	0.16	2.60	6.37	12.28	24.6	5.71	3.55
CH6	1.51	1.43	0.24	0.18	2.55	3.62	11.77	7.30	5.11	2.78
CH7	-0.59	1.39	0.32	0.21	3.09	3.49	8.24	3.55	9.88	3.69
CR1	0.14	1.61	0.04	0.16	4.55	3.13	6.44	3.37	9.15	3.60
CR3	1.12	0.91	0.10	0.11	-1.03	3.00	4.71	10.3	3.42	3.62
CR5	1.00	1.10	0.17	0.16	6.22	8.55	-9.21	28.8	3.11	2.62
CR6	1.03	1.21	0.00	0.13	0.94	2.75	4.75	2.38	6.04	3.29
CR8	2.13	1.73	0.49	0.12	4.42	5.86	0.00	27.9	8.77	4.93
CR10	1.31	1.60	0.07	0.15	-2.49	4.35	-6.97	18.3	7.66	3.77
CR12	0.06	1.05	0.15	0.13	5.53	5.59	-16.81	30.4	2.16	2.55
CR14	-0.66	1.44	0.09	0.15	1.87	4.40	6.75	3.75	9.24	4.13
CR15	1.25	1.12	0.26	0.14	6.11	6.82	17.45	17.8	10.86	3.62
LC1	0.40	1.01	0.02	0.15	4.55	1.90	6.02	1.78	4.88	2.97
LC2	1.42	1.13	0.05	0.12	2.69	2.91	4.80	1.99	0.80	2.05
LC4	1.34	1.92	0.13	0.11	3.89	2.14	2.44	1.74	4.06	3.62
LC5	0.37	1.16	0.21	0.15	4.24	6.50	-1.16	1.74	4.51	3.44
LC9	0.21	1.29	0.20	0.16	3.00	4.24	13.54	9.81	6.26	3.42

Table E1. Radioisotope activity (dpm/g) and error (\pm) for all samples. Significant, measurable activities and associated error are indicated by black font. Red font indicates that no significant radioisotope activities were measured.

Appendix F: Multivariate Statistic Data SIMCA Classification Results

uley were previously identified as oddiels and removed from the model.						
Sample	Lake Calumet	Calumet River	Calumet Harbor	Lake Michigan		
PC 2						
PC 1						
LC 9	*					
LC 7	*					
LC 8	*					
LC 6	*	*				
LC 5	*					
LC 4	*					
LC 3	*					
LC 2	*					
LC 1	*					
CR 1		*				
CR 2		*				
CR 3		*				
CR 4		*				
CR 5		*				
CR 6		*				
CR 7		*				
CR 8		*				
CR 9		*				
CR 10		*				
CR 11		*				
CR 12		*				
CR 13		*				
CR 14		*				
CR 15		*				
CH 1			*			
CH 2			*			
CH 3			*			
CH 4			*	*		
CH 5			*	*		
CH 6			*	*		
CH 7			*			
	L	((ſ		

Table F1. SIMCA classification results. No grouping was assigned to PC samples because they were previously identified as outliers and removed from the model.

Sample	Lake Calumet	Calumet River	Calumet Harbor	Lake Michigan
LM 1				*
LM 2				*
LM 3				*
LM 4				*
LM 5				*
LM 6				*
LM 7			*	*
LM 8				*
LM 9				*
LM 10				*
LM 11				*
LM 12				*
LM 13				*
LM 14				*
LM 15				*
LM 16				*
LM 17				*
LM 18				*
LM 19				*
LM 20				*
IC 1				
IC 2				

Table F2. SIMCA classification results. No grouping was assigned IC samples because they were previously identified as outliers and removed from the model.

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within a region of concern. To he Thomas J. O'Brien lock and Lake sources. Additionally, geochemic geochemical fingerprint for sedin used to successfully distinguish b metals and a geochemical signatu occurring locally within the river, contaminants in the dredge record successfully used to track change	Ip identify possible sources of contaminated sedim Michigan, multiple bottom sediment samples wer al data from historic dredging records spanning a 4 hents within this reach of the river. Geochemical m etween all six potential sediment sources. Calumet	0-year period were examined to develop a historic easurements and advanced multivariate statistics were River samples showed elevated levels of heavy nents, suggesting that heavy metal contamination was ow historic reductions in heavy metal and organic Clean Water Act. Geochemical fingerprints were the project area. These fingerprints could be used
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