Predicting the Contamination between
Sites of Sediment Core Measurement in Lake Ontario

by

Daniel J. Jakubek

A Research Paper

Presented to Ryerson University

in partial fulfillment of the requirements for the degree of

Master of Spatial Analysis

A joint program with the University of Toronto

Toronto, Ontario, Canada

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Author’s Declaration

I hereby declare that I am the sole author of this Research Paper.

I authorize Ryerson University to lend this Research Paper to other institutions or individuals for the purposes of scholarly research.

_________________________
Daniel J. Jakubek
Abstract

Research activities involving the collection of sediment core samples are extremely time consuming and expensive to fund. This research utilized data from the Environment Canada Great Lakes Sediment Assessment Program. A total of 32 contaminants were measured at 70 sediment core-sampling locations in Lake Ontario. Various methods for the estimation of sediment contamination levels were investigated. The Sediment Quality Index (SQI) was calculated and assessed as being a satisfactory measure for areas where sediment quality is frequently threatened or impaired.

Ordinary kriging was identified as the optimal spatial interpolation model. Individual prediction maps were successfully produced for 20 of the contaminants and cross-validation was used to further assess 'ordinary kriging' as an appropriate method for predicting the spatial distribution of sediment contamination. The results rendered from cross-validation provided an assessment of the relative success of each of the interpolation procedures. Limitations including the limited number of sampling sites, and minimal data at Lake Ontario Areas of Concern, were the main factors for inaccurate prediction surfaces.
Acknowledgements

I would like to express appreciation to my graduate faculty advisor Dr. Wayne Forsythe for his ongoing support and guidance while completing this research project, and throughout the Master of Spatial Analysis program. As a result of his assistance, personal goals set throughout the program were attainable. I would also like to thank Dr. Marie Truelove, Director of the M.S.A. program, whose organization enabled me to acquire professional experience at the Canadian Ministry of Environment through the practicum component of the program.

While completing the required practicum with The Ecosystem Health Division, I was supplied with data that was utilized to perform the analysis in this research project. Sediment contamination levels were initially measured for use in the Environment Canada Great Lakes Sediment Assessment Program. Specifically, I would like to thank the following individuals for their support:

Scott Painter, Watershed Scientist, Ecosystem Health Division, Canada Centre for Inland Waters, Canadian Ministry of the Environment

Alice Dove, Watershed Scientist, Ecosystem Health Division, Canada Centre for Inland Waters, Canadian Ministry of the Environment
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<th>Acronym</th>
<th>Description</th>
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</thead>
<tbody>
<tr>
<td>AOC</td>
<td>Area of Concern</td>
</tr>
<tr>
<td>ARCS</td>
<td>Assessment and Remediation of Contaminated Sediment</td>
</tr>
<tr>
<td>ASE</td>
<td>Average Standard Error</td>
</tr>
<tr>
<td>CCME</td>
<td>Canadian Council of Ministers of the Environment</td>
</tr>
<tr>
<td>CWQI</td>
<td>Canadian Water Quality Index</td>
</tr>
<tr>
<td>DDT</td>
<td>Dichloro-Diphenyl-Trichloroethane</td>
</tr>
<tr>
<td>EC</td>
<td>Environment Canada</td>
</tr>
<tr>
<td>EPA</td>
<td>Environmental Protection Agency</td>
</tr>
<tr>
<td>ERL</td>
<td>Effects Range Low</td>
</tr>
<tr>
<td>ERM</td>
<td>Effect Range Median</td>
</tr>
<tr>
<td>ESRI</td>
<td>Environmental Systems Research Institute</td>
</tr>
<tr>
<td>GIS</td>
<td>Geographic Information Systems</td>
</tr>
<tr>
<td>GLA</td>
<td>Great Lakes Atlas</td>
</tr>
<tr>
<td>GLFS</td>
<td>The Great Lakes Forecasting System</td>
</tr>
<tr>
<td>GLIN</td>
<td>The Great Lakes Information Network</td>
</tr>
<tr>
<td>GLWQA</td>
<td>Great Lakes Water Quality Agreement</td>
</tr>
<tr>
<td>GPI</td>
<td>Global Polynomial Interpolation</td>
</tr>
<tr>
<td>HCB</td>
<td>Hexachlorobenzene</td>
</tr>
<tr>
<td>IDW</td>
<td>Inverse Distance Weighting</td>
</tr>
<tr>
<td>IJC</td>
<td>International Joint Commission</td>
</tr>
<tr>
<td>LEL</td>
<td>Lowest Effect Level</td>
</tr>
<tr>
<td>LEWQ</td>
<td>Lake Erie Water Quality</td>
</tr>
<tr>
<td>LOLMP</td>
<td>Lake Ontario Lakewide Management Plan</td>
</tr>
<tr>
<td>LOTMP</td>
<td>Lake Ontario Toxics Management Plan</td>
</tr>
<tr>
<td>LPI</td>
<td>Local Polynomial Interpolation</td>
</tr>
<tr>
<td>LWD</td>
<td>Low Water Datum</td>
</tr>
<tr>
<td>MEL</td>
<td>Minimum Effect Levels</td>
</tr>
<tr>
<td>MOE</td>
<td>Ministry of the Environment</td>
</tr>
<tr>
<td>MPE</td>
<td>Mean Prediction Error</td>
</tr>
<tr>
<td>NOAA</td>
<td>National Oceanic and Atmospheric Administration</td>
</tr>
<tr>
<td>nse</td>
<td>Normalized Sum of Excursions</td>
</tr>
<tr>
<td>NYSDEC</td>
<td>New York State Department of Environmental Conservation</td>
</tr>
<tr>
<td>OC</td>
<td>Organochlorine</td>
</tr>
<tr>
<td>PAH</td>
<td>Polycyclic Aromatic Hydrocarbon</td>
</tr>
<tr>
<td>PCB</td>
<td>Polychlorinated Biphenyls</td>
</tr>
<tr>
<td>PCDD</td>
<td>Polychlorinated Dibenzo-p-Dioxins and dibenzofurans</td>
</tr>
<tr>
<td>PCDF</td>
<td>Polychlorinated Dibenzofurans</td>
</tr>
<tr>
<td>PEL</td>
<td>Probable Effect Level</td>
</tr>
<tr>
<td>POP</td>
<td>Persistent Organic Pollutants</td>
</tr>
<tr>
<td>RBF</td>
<td>Radial Basis Functions</td>
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<tr>
<td>RMSPE</td>
<td>Root-Mean-Square-Prediction Error</td>
</tr>
<tr>
<td>RVT</td>
<td>Regionalized Variable Theory</td>
</tr>
<tr>
<td>SMPE</td>
<td>Standardized Mean Prediction Error</td>
</tr>
<tr>
<td>Acronym</td>
<td>Description</td>
</tr>
<tr>
<td>---------</td>
<td>--------------------------------------------------</td>
</tr>
<tr>
<td>SQI</td>
<td>Sediment Quality Index</td>
</tr>
<tr>
<td>SQT</td>
<td>Sediment Quality Triad</td>
</tr>
<tr>
<td>SRMSPE</td>
<td>Standardized Root Mean Square Prediction Error</td>
</tr>
<tr>
<td>TEL</td>
<td>Threshold Effect Level</td>
</tr>
<tr>
<td>USEPA</td>
<td>United States Environmental Protection Agency</td>
</tr>
</tbody>
</table>
Chapter 1: Introduction

1.1 Introduction

Over the past century, Lake Ontario has experienced contamination of sediment, water, and biota as a result of anthropogenic activities, including mass development along the Canadian portion of its western shoreline known as the Golden Horseshoe (Figure 1.1).

The Golden Horseshoe includes the major cities of Toronto, Mississauga, Oakville, Hamilton, St. Catharines, Niagara Falls, Burlington, and Oshawa. Half the population of Ontario lives in or around these cities (ASG, 2002). With a general objective to “restore the overall health of the Great Lakes ecosystem” (LOLMP, 1998), Canadian and American government institutions combined resources in the development of a Lakewide Management Plan. As a result of these actions, toxic contamination in the Lake Ontario basin has decreased, however, contaminants remain in the ecosystem with the capacity to
bioaccumulate (accumulate in aquatic organisms to levels that are harmful to human health) (LOLMP, 1998). It is due to the persistence of these toxic contaminants that research regarding the sediment and water quality in the Great Lakes continues.

1.2 Pollution in the Great Lakes

The Great Lakes Basin (Figure 1.2) consists of approximately 23,000 km$^3$ of fresh water, representing 18 % of the world’s supply (GLA, 1995).

![Figure 1.2 – The Great Lakes Basin](image)
(Source: U.S. Army Corps of Engineers, Detroit District in GLIN, 2002)

Toxic pollutants can be found in the aquatic system due to the re-suspension of sediment, cycling through biological food chains, and continuing pollution-causing processes. These pollutants are human-made organic chemicals and heavy metals that can be toxic
in small amounts and cause negative effects in minute concentrations over a long period of time. The first Europeans to settle in this region in the 1600’s limited their exploitation of the natural resources to wildlife in the system. However, the arrival of new immigrant groups caused ecological changes through processes including logging, farming, and commercial fishing.

After the turn of the 20th century, growing urbanization and industrial development in the Great Lakes Basin caused widespread bacterial contamination and added to the floating debris produced by activities such as logging and agriculture. Continued industrialization and intensified agricultural practices were the causes for the development of new chemical substances. Polychlorinated Biphenyls (PCBs) and Dichloro-Diphenyl-Trichloroethane (DDT) were developed for use as pesticides in agricultural activity in the 1920s and 1940s respectively (Hodgson and Levi, 1997). Toxic runoff produced by these pesticides, the use of synthetic fertilizers developed to further enhance crop yield, existing sources of nutrient rich pollutants (untreated human waste from urban areas), and phosphate detergents accelerated the rate of biological production in the system (GLA, 1995). Eutrophic-imbalance was first documented in Lake Erie in the 1950’s. The imbalance was characterized by a depletion of dissolved oxygen and the formation of massive algal blooms in this body of water (GLA, 1995). By 1980, the International Joint Commission (IJC) estimated that approximately 2500 chemicals were in common use in the Great Lakes Basin.
The major industries located in the Great Lakes region include steel production, pulp and paper, chemicals, automobiles, and manufactured goods. The most significant urban areas were developed at the mouths of Great Lakes tributaries due to transportation needs and freshwater resources for domestic and industrial use (GLA, 1995). Lake Erie features the smallest water volume of the Great Lakes and it is significantly affected by urbanization and agricultural practices in its surrounding area. Agricultural lands in the Lake Erie basin are extremely fertile. As a result, contaminants enter the lake as runoff from intensive farming practices in southwestern Ontario, Ohio, Indiana, and Michigan. Furthermore, 17 metropolitan areas featuring populations greater than 50,000 surround Lake Erie and act as sources of contamination to the system (LEWQ, 1989). Lake Ontario is similarly affected by urbanization and agricultural practices, although, industrialization is also a main pollution factor. As early as 1870, water could not be drawn from Hamilton Harbour or local wells due to high contamination levels in this area (GLA, 1995).

Lake Michigan is the second largest of the Great Lakes and is located entirely within the United States. The population is sparse in its northern basin, however this area receives wastes from the world’s largest concentration of pulp and paper mills. Furthermore, the southern basin consists of U.S. metropolitan areas including Milwaukee and Chicago. Combined, these cities account for 20% of the human population in the Great Lakes Basin (GLA, 1995). Lakes Huron and Superior are the least contaminated entities in the Great Lakes system. The Saginaw River basin is the largest contributor of contamination in Lake Huron resulting from agricultural practices. The Lake Superior
basin features a cool climate with poor soils. These factors coupled with its northern location make it the cleanest of the Great Lakes. The main source for contamination is atmospheric deposition.

1.3 Study Area

Lake Ontario is located in the southeastern part of the Great Lakes Basin. It has an area of approximately of 19010 square kilometres, and is the smallest of the Great Lakes (GLFS, 2002). With a mean surface elevation of 75 meters above sea level, it has the lowest elevation of the Great Lakes (GLFS, 2002). It does however feature the highest ratio of watershed area to lake surface area among all of the Great Lake basins (LOLMP, 1998). A full account of Lake Ontario’s physical characteristics (including depth) can be found in Table 1.1 and Figure 1.3.

Table 1.1: Physical Characteristics of Lake Ontario

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Metric Units</th>
</tr>
</thead>
<tbody>
<tr>
<td>Low Water Datum (LWD)</td>
<td>74.00 m</td>
</tr>
<tr>
<td>Length</td>
<td>311.00 km</td>
</tr>
<tr>
<td>Width</td>
<td>85.3 m</td>
</tr>
<tr>
<td>Shoreline Length</td>
<td>1168.38 km</td>
</tr>
<tr>
<td>Total Surface Area</td>
<td>19010.42 km²</td>
</tr>
<tr>
<td>Surface Area in Canada</td>
<td>10049.11 km²</td>
</tr>
<tr>
<td>Surface Area in US</td>
<td>8961.31 km²</td>
</tr>
<tr>
<td>Water Volume at LWD</td>
<td>1638.08 km³</td>
</tr>
<tr>
<td>Average Depth Below LWD</td>
<td>86.26 m</td>
</tr>
<tr>
<td>Maximum Depth Below LWD</td>
<td>244.45 m</td>
</tr>
<tr>
<td>Average Surface Elevation</td>
<td>74.61 m</td>
</tr>
<tr>
<td>Maximum Surface Elevation</td>
<td>75.61 m</td>
</tr>
<tr>
<td>Minimum Surface Elevation</td>
<td>73.59 m</td>
</tr>
</tbody>
</table>

(Source: The Great Lakes Forecasting System: Lake Bathymetry Data, 2002)
Lake Ontario’s drainage basin covers portions of the Canadian Province of Ontario and New York State in the United States. It is fed primarily by the waters of Lake Erie through the Niagara River. The average inflow discharge is approximately 7000 m$^3$/s (Atkinson et al., 1994), and this flow accounts for nearly 80 percent of the total inflow into Lake Ontario (Blair et al., 1993). Additional inflow (14 percent) stems from other Lake Ontario basin tributaries including the Genesee, Oswego, and Black Rivers in New York, and the Trent River in Ontario (LOLMP, 1998). The remaining inflow enters as precipitation and represents approximately six percent of the water body’s total volume (LOLMP, 1998). Approximately 93 percent of the water in Lake Ontario is drained to the northeast by the St. Lawrence River, with the remaining seven percent lost through evaporation (LOLMP, 1998). The outflow discharge rate into the St. Lawrence River...
averages 7400 m$^3$/second (Rukavina et al., 1990). The Lake Ontario Drainage Basin is represented in Figure 1.4.

The entire Great Lakes Basin can be characterized as having a temperate and humid climate (USEPA et al., 1987 in LOLMP, 1998). Warm, humid air masses originating in the Gulf of Mexico influence the Lake Ontario Basin in the summer months, whereas Arctic and Pacific air masses influence the area in the winter. Due to heat transfer processes, near shore areas feature temperate climates uncommon to Lake Ontario’s northern latitude. When atmospheric temperatures are high, radiant energy is absorbed by the water and subsequently released when temperatures are lower (LOLMP, 1998).
1.3.1 Data Samples and Sampling Locations

The geology within the Lake Ontario Basin is classified as either non-depositional consisting of material such as bedrock common to inshore areas, or depositional materials such as glacial till and fine-grained particulates including silts and clays that accumulate in deeper offshore areas. In this analysis, field research conducted under the Environment Canada Great Lakes Sediment Assessment Program provided sediment contamination data for 32 variables measured at 70 specific sampled sites in 1998 (Figure 1.5). The sites were selected at intervals of approximately 30 km.

The headings chosen to create the grid of measured locations were east/west and north/south. Deviations from the grid formation were made in order to assess Lake Ontario Areas of Concern (AOCs) including Hamilton Harbour and the mouth of the Niagara River. The U.S-Canada Great Lakes Water Quality Agreement define AOCs as "geographic areas that fail to meet the general or specific objectives of the agreement where such failure has caused or is likely to cause impairment of beneficial use of the area's ability to support aquatic life." Due to time and funding restraints, the spatial distribution of sediment contamination throughout Lake Ontario could not be more thoroughly measured. However, deterministic and geostatistical interpolation techniques can be used to estimate the spatial distribution of sediment contamination within Lake Ontario.
Lake Ontario Sediment Sampling Locations

Legend
- Sediment Sampling Locations

Figure 5 - Lake Ontario Sediment Sampling Locations
Surficial sediment samples were collected using a mini-box core sampling procedure. The samples collected during the survey consisted of fine-grained sediments classified as clay, sand, silt, or mud. The initial 3 centimetres of the sediment was sub-sampled in order for analyses of persistent organic pollutants (POPs), metals, particle size characterization, and nutrients to be performed (Marvin et al., 2002). Table 1.2 documents the specific contaminants that were measured at each of the locations within Lake Ontario and their corresponding federal guideline levels.

Ouyang et al. (2002) concluded that heavy metal concentrations in sediment including lead, copper, zinc, and cadmium that were located above a sediment depth of 1.5 metres posed a threat to the health of aquatic organisms. Furthermore, it is important to note that the influence of the particle size on contaminant concentrations in sediment usually shows an inverse correlation with grain size (Ouyang et al., 2002).

1.4 The Problem

The estimation of contaminant loading into Lake Ontario and identification of the sources for this loading are difficult tasks. Furthermore, research activities which exist in order to measure contaminant levels at specific locations throughout this body of water are both extremely time-consuming and expensive to fund. In order to identify the potential ‘hotspots’ (areas creating ecosystem risk) for sediment contamination in Lake Ontario, the Sediment Quality Index (SQI) was used. The SQI performs risk assessment
### Table 1.2: Table of Contaminants and Federal Guidelines

<table>
<thead>
<tr>
<th>Contaminant</th>
<th>Threshold Effect Level TEL</th>
<th>Probable Effect Level PEL</th>
</tr>
</thead>
<tbody>
<tr>
<td>Arsenic</td>
<td>5.9 ug/g</td>
<td>17 ug/g</td>
</tr>
<tr>
<td>Cadmium</td>
<td>0.6 ug/g</td>
<td>3.53 ug/g</td>
</tr>
<tr>
<td>Chromium</td>
<td>37.3 ug/g</td>
<td>90 ug/g</td>
</tr>
<tr>
<td>Copper</td>
<td>35.7 ug/g</td>
<td>196.6 ug/g</td>
</tr>
<tr>
<td>Lead</td>
<td>35 ug/g</td>
<td>91.3 ug/g</td>
</tr>
<tr>
<td>Nickel</td>
<td>16 ug/g</td>
<td>75 ug/g</td>
</tr>
<tr>
<td>Zinc</td>
<td>123 ug/g</td>
<td>314.8 ug/g</td>
</tr>
<tr>
<td>Mercury</td>
<td>0.17 ug/g</td>
<td>0.486 ug/g</td>
</tr>
<tr>
<td>Alpha-HCH or BHC</td>
<td>6 ng/g</td>
<td>200 ng/g</td>
</tr>
<tr>
<td>hexachlorobenzene (HCB)</td>
<td>20 ng/g</td>
<td>480 ng/g</td>
</tr>
<tr>
<td>Beta-HCH (Lindane)</td>
<td>0.94 ng/g</td>
<td>1.38 ng/g</td>
</tr>
<tr>
<td>Heptachlor Epoxide</td>
<td>0.6 ng/g</td>
<td>2.74 ng/g</td>
</tr>
<tr>
<td>Alpha-Chlordane</td>
<td>4.5 ng/g</td>
<td>8.87 ng/g</td>
</tr>
<tr>
<td>Dieldrin</td>
<td>2.85 ng/g</td>
<td>6.67 ng/g</td>
</tr>
<tr>
<td>pp’ – DDE</td>
<td>1.42 ng/g</td>
<td>6.75 ng/g</td>
</tr>
<tr>
<td>Endrin</td>
<td>2.67 ng/g</td>
<td>62.4 ng/g</td>
</tr>
<tr>
<td>pp’ – DDD</td>
<td>3.54 ng/g</td>
<td>8.51 ng/g</td>
</tr>
<tr>
<td>op’ - + pp’ DDT</td>
<td>1.19 ng/g</td>
<td>4.77 ng/g</td>
</tr>
<tr>
<td>Mirex</td>
<td>7 ng/g</td>
<td>2600 ng/g</td>
</tr>
<tr>
<td>polychlorinated biphenyls (PCBs)</td>
<td>34.1 ng/g</td>
<td>277 ng/g</td>
</tr>
<tr>
<td>Dioxins and Furans</td>
<td>0.85 ng/g</td>
<td>21.5 ng/g</td>
</tr>
<tr>
<td>phenanthrene</td>
<td>41.9 ng/g</td>
<td>515 ng/g</td>
</tr>
<tr>
<td>Anthracene</td>
<td>46.9 ng/g</td>
<td>245 ng/g</td>
</tr>
<tr>
<td>Fluoranthene</td>
<td>111 ng/g</td>
<td>2,355 ng/g</td>
</tr>
<tr>
<td>Pyrene</td>
<td>53 ng/g</td>
<td>875 ng/g</td>
</tr>
<tr>
<td>benzo[a]anthracene</td>
<td>31.7 ng/g</td>
<td>385 ng/g</td>
</tr>
<tr>
<td>Chrysene</td>
<td>57.1 ng/g</td>
<td>862 ng/g</td>
</tr>
<tr>
<td>benzo[b/k]fluoranthene</td>
<td>240 ng/g</td>
<td>26,800 ng/g</td>
</tr>
<tr>
<td>benzo[a]pyrene</td>
<td>31.9 ng/g</td>
<td>782 ng/g</td>
</tr>
<tr>
<td>benzo[1,2,3-cd]pyrene</td>
<td>200 ng/g</td>
<td>6400 ng/g</td>
</tr>
<tr>
<td>dibenzo[a,h]anthracene</td>
<td>6.22 ng/g</td>
<td>135 ng/g</td>
</tr>
<tr>
<td>benzo[g,h,i]perylene</td>
<td>170 ng/g</td>
<td>6400 ng/g</td>
</tr>
<tr>
<td>total polycyclic aromatic hydrocarbons (PAHs)</td>
<td>4,000 ng/g</td>
<td>200,000 ng/g</td>
</tr>
</tbody>
</table>

(Source: Canadian Council of Ministers of the Environment, 1999)

by providing a general description of sediment quality on the basis of whether existing federal contamination guidelines are exceeded.
1.4.1 Sediment Contamination Research

The United States Environmental Protection Agency’s (USEPA) Assessment and Remediation of Contaminated Sediment (ARCS) program exists in order to determine the nature and extent of sediment contamination in the Great Lakes (Burton et al., 1996). In ARCS analyses, benthic invertebrate communities are often used as bioindicators for sediment contamination. Since the habitat of benthic communities coincides with lake-bottom sediment, small crustaceans commonly referred to as Amphipod Diporeia have been used as potential bioindicators for contaminants including metals in sediment (Song and Breslin, 1998). Many studies have reported changes in benthic community composition resulting from sediment contamination (Hilsenhoff, 1987; Waterhouse and Farrell, 1985; Clements et al., 1992 in Canfield et al., 1996). Processes such as the Sediment Quality Triad (SQT) approach (Chapman, 1992 in Canfield et al., 1996) are also used in evaluations of how benthic community composition measures can aid in the assessment of contaminated and uncontaminated sediment (Canfield et al., 1996). The SQT is a weight of evidence approach that incorporates measures of sediment chemistry, sediment toxicity, and benthic community composition in evaluations of sediment quality (Canfield et al., 1996).

Guidelines, objectives, and criteria represent numerical sediment quality assessment values used in sediment toxicology. Examples of such assessment values include the Threshold Effect Level (TEL) and Probable Effect Level (PEL), which were used in this analysis. The TEL represents “the concentration below which adverse
biological effects are expected to occur rarely,” whereas, the PEL defines the level above which adverse effects are expected to occur frequently (CCME, 1999). These assessment values are developed using a weight of evidence approach in which biological and chemical data from modeling, laboratory, and field studies performed on fresh water sediments are analyzed (Smith et al., 1996). The calculation of two such assessment values defines three ranges of chemical concentration: those that are rarely, occasionally, and frequently associated with adverse biological effects.

In past research, numerical quality assessment values were created and used in a comparison to assess the TEL and PEL as good indicators of the severity of sediment contamination. In these analyses, the TEL was compared to the four following assessment values: (1) Ontario’s Provincial Sediment Quality Guidelines (Lowest effect Levels or LEL) (Persaud et al., 1992); (2) the Minimum Effect Levels (MEL) developed for the St. Lawrence River (MENVIQ/EC, 1992 in Smith et al., 1996); (3) the Effects Range Low (ERL), created by the National Oceanic and Atmospheric Administration (NOAA); and (4) the sediment effect concentrations for benthic communities (Ingersol et al., 1996 in Smith et al., 1996). The PEL was similarly compared to the four following values: (1) the Severe Effect Levels (SEL) (Persaud et al., 1992); (2) the Toxic Effect Levels (MENVIQ/EC, 1992 in Smith et al., 1996); (3) the Effect Range Median (ERM) values (Long, 1992 in Smith et al., 1996); and (4) the probable effect levels reported for benthic communities (Ingersol et al., 1996 in Smith et al., 1996).
Sediment quality can be assessed by utilizing the Sediment Quality Index (SQI). This measure of sediment quality is derived from the Canadian Water Quality Index (CWQI). The SQI formula, based on the CWQI, was developed by the British Columbia Ministry of Environment, Lands and Parks, and modified by the Ministry of Environment in Alberta (CCME, 1999). A sediment quality index is a means of summarizing complex sediment contamination data mathematically, by combining all existing measures of contamination to provide a general description of sediment quality within a body of water. The index is useful in assessing sediment quality relative to its desired state, defined by specific objectives. Additionally, this index addresses the degree to which water quality is affected by human activity.

Marvin et al. (2002) applied the SQI to an assessment of sediment quality in Lakes Erie and Ontario on the basis of the federal PEL guideline (contaminants such as Mirex that reside within sediment in minute concentrations utilized the provincial LEL guideline). They did not, however, use interpolation procedures to estimate sediment quality throughout the lakes; SQI scores were calculated on the basis of point data values in order to examine the spatial pattern of sediment quality. The spatial trends in sediment quality in the two lakes reflected the trends for individual contaminant classes such as mercury and PCBs. The Lake Ontario data is featured in this analysis.
1.4.2 Objectives

In this study, the characteristics and spatial distribution of 32 contaminants in Lake Ontario sediment are investigated using field measurements. The specific objectives in this analysis can be summarized as:

1) Assess the Sediment Quality Index as a satisfactory measure for the areas in Lake Ontario where sediment quality is frequently threatened or impaired;

2) Identify whether deterministic or geostatistical interpolation methods are more appropriate methods for predicting spatial distributions of contaminants in sediment using SQI scores;

3) Assess existing interpolation procedures and identify an optimal method for the 32 specific contaminants measured in the Lake Ontario sediment samples.
Chapter 2: Literature Review

2.1 Sedimentation Processes in Lake Ontario

Lake Ontario is composed of two main sedimentary basins: (1) the Kingston basin (a relatively shallow basin located at the northeastern end of Lake Ontario; and (2) a main basin including the Niagara, Mississauga, and Rochester Basins (relatively deep sub-basins bordered by shallow inshore zones extending along the entire perimeter of the main basin) (LOLMP, 1998). Figure 2.1 displays the Lake Ontario sedimentary basins.

Analyses performed on suspended sediment concentrations demonstrated that the Niagara River supplies approximately 1.8 million tons of sediment to Lake Ontario annually (Joshi et al., 1992). A physical feature located at the mouth of the Niagara River named the Niagara Bar is an example of a shallow inshore zone created by the inflow of sediment. Sediment is deposited at this junction because the velocity of the current in
Lake Ontario is lower than that of the Niagara River. As a result, shear stress on the river bottom is increased, and buoyant discharge conditions result in a defined surface plume. Due to inertia and buoyancy, and after a moderate distance, the Coriolis acceleration, the plume is turned clockwise in an easterly direction (Atkinson et al., 1994).

The majority of water circulation in Lake Ontario occurs within its main sub-basins and eastern shore. However, discharges by rivers and estuaries can modify the circulation of Lake Ontario waters, due to baroclinic pressure gradients caused by buoyancy input and through the initial momentum flux of the discharge (Atkinson et al., 1994). It is arguable that a lakewide circulation with an eastern heading may be initiated from the Niagara River inflow. Furthermore, the area is affected by prevailing winds from a west-northwest direction (LOLMP, 1998). The combination of these factors results in water circulation that moves in a counter-clockwise motion (Sly, 1990), and significantly less net flow along the north inshore zone of Lake Ontario (LOLMP, 1998). Figure 2.2 displays the annual current circulation in Lake Ontario featuring a general west-east heading.

2.2 Lake Ontario’s Lakewide Management Plan

Lake Ontario is vulnerable to human activities that have occurred throughout the Lake Superior, Michigan, Huron, and Erie basins, since it is located at the bottom end of the Great Lakes system. Over the past century, the Lake Ontario ecosystem has experienced negative changes as a result of toxic pollution originating from the excessive
development of the Great Lakes region. Major industrial centres including Hamilton, Toronto, Oshawa, and Kingston are situated on its Canadian shoreline to the north. The cities of Rochester and Oswego are located on its American shore in the state of New York to the south. These point sources of pollution, combined with dredging practices in the upstream Great Lakes tributaries such as the Niagara River, have been major contributors to poor sediment and water quality in Lake Ontario (GLA, 1995). In 1972, the Canadian and United States governments agreed that water quality was to be improved in the Great Lakes, and future pollution input levels were to be decreased (Zarull et al., 1999). The Great Lakes Water Quality Agreement was renewed in 1987 in order to ban and control the contaminants entering the Great Lakes and restore the health of the Great Lakes ecosystem (LOLMP, 1998). In addition, a Lakewide Management Plan (LMP) was developed for each of the Great Lakes and signed by the four parties involved in its implementation: Region II of the USEPA, Environment Canada (EC), the New York State Department of Environmental Conservation (NYSDEC), and the Ontario Ministry of the Environment (MOE). Given the abundance of toxins identified in the
Niagara River and Lake Ontario, it was necessary for the four involved parties to develop a Lake Ontario Toxics Management Plan (LOTMP). The main purpose of LOTMP was to “define the toxics problem in Lake Ontario and to develop and implement a plan to eliminate the problem through both individual and joint agency actions” (LOLMP, 1998).

To initiate implementation of LOTMP, it was necessary to identify the priority toxic chemicals residing in Lake Ontario. This was accomplished on the basis of ‘impairment of beneficial use(s),’ which “is a change in the chemical, physical, or biological integrity of the Great Lakes System sufficient to cause any of the following (Hartig et al., 1990):

(1) Restrictions on fish and wildlife consumption;
(2) Tainting of fish and wildlife flavour;
(3) Degradation of fish and wildlife populations;
(4) Fish tumours or other deformities;
(5) Bird or animal deformities or reproductive problems;
(6) Degradation of benthos;
(7) Restrictions on dredging activities;
(8) Eutrophication or undesirable algae;
(9) Restrictions on drinking water consumption, or taste and odor problems;
(10) Closing of beaches;
(11) Degradation of aesthetics;
(12) Added costs to agriculture or industry;
(13) Degradation of phytoplankton and zooplankton populations;
(14) Loss of fish and wildlife habitat.”

In a study conducted by Zarull et al. (1999), all use impairments (excluding tainting of fish and wildlife flavour, restrictions on drinking water consumption, and the closing of
beaches) are potentially associated with contaminated sediment. It was also necessary to identify the critical pollutants in Lake Ontario “that persist at levels that, singly or in synergistic or additive combination, are causing, or are likely to cause, impairment of beneficial uses past application of regulatory controls due to their (GLA, 1995):

(1) Presence in open lake waters;
(2) Ability to cause or contribute to a failure to meet Agreement objectives through their recognized threat to human health and aquatic life;
(3) Ability to bioaccumulate.”

In the Niagara River and Lake Ontario, the Lakewide Critical Pollutants identified as the focus of reduction activities are as follows (LOLMP, 1998):

(1) polychlorinated biphenyls (PCBs);
(2) 1,1,1-Trichloro-2,2-bis(P-chlorophenyl)ethane (DDT) and its metabolites;
(3) Mirex;
(4) Dioxins/Furans;
(5) Mercury; and
(6) Dieldrin.

These toxins were identified as the critical pollutants in the Lake Ontario basin because they bioaccumulate and remain in water, sediment, and biota for long periods of time.

2.3 Critical Pollutants and Sources of Loading in Lake Ontario

Estimating the critical pollutant loading that enters Lake Ontario via sources including rivers/tributaries, precipitation, sewage treatment facilities, waste sites, agricultural runoff, and other sources is a difficult task. The sources for environmental toxicants can generally be categorized as point sources or non-point sources. A point
source of loading are discharges of chemicals that can be identified or measured, including industrial or municipal effluent outfalls, chemical or petroleum spills and dumps, smokestacks, and other stationary atmospheric charges (Hodgson et al., 1997). Non-point sources include more diffuse inputs over large areas, that do not have a identifiable point of entry, such as pesticide and fertilizer runoff, mobile source emissions, atmospheric deposition, and contaminated sediment or mine tailings (Hodgson et al., 1997). The Great Lakes Water Quality Agreement (GLWQA) categorizes critical pollutant loadings into four general groups:

1. Loadings from sources outside the Lake Ontario Basin
2. Loadings from sources inside the Lake Ontario basin
3. Atmospheric loadings and
4. Releases from Lake Ontario to the St. Lawrence River and volatilization to the atmosphere (LOLMP, 1998).

Loading estimates are often inaccurate due to the accumulation of data from various sources and variations in the data collection methodologies employed by different monitoring programs. With inaccurate data, contaminants measured as outflow from the lake, including PCBs and DDT, may be measured at higher levels than the inflow of these contaminants. An explanation for such an event is that contaminants are being released into Lake Ontario from sediments located at the bed of the water body (LOLMP, 1998). The water retention time within Lake Ontario is estimated at approximately seven years (Sly, 1991). However, contaminants have a tendency to bind to sediments on lake bottoms. If this process occurs, these contaminants can be covered over by additional sediment, and remain in the system for an indefinite period of time. Thus, toxic
substances have the ability to remain in a lake ecosystem for periods of time extending past a seven-year retention span (Sly, 1991).

While toxic substances have the capacity to remain in lake-bottom sediment for extended periods of time, they also may be re-suspended into the water column by the processes of bioturbation and re-suspension due to storm events and dredging activities. If this process occurs, these contaminants may be transferred to higher trophic levels in the food chain. This is possible due to the presence of ‘benthos’, organisms whose habitat coincides with lake-bottom sediments (Song and Breslin, 1998). Benthic organisms can be a significant source of food for aquatic organisms.

The Lake Ontario AOCs located in New York State include Eighteen Mile Creek, the Niagara River, Oswego River/Harbour, the Rochester Embayment, and the St. Lawrence River at Massena. In the Province of Ontario, AOCs include the Bay of Quinte, Port Hope, the Toronto Waterfront, and Hamilton Harbour. Sediment contamination levels have been reported at higher levels in these regions than in open-water areas (EPA, 2000). In these AOCs, benthic communities have been degraded.

Lake Ontario’s benthic communities are dominated by small crustaceans (Diporeia spp.) and worms (Stylodrillus heringianus) (Song and Breslin, 1998). The health of benthic organisms in Lake Ontario is a good indicator of the lake’s environmental quality. This is because such organisms require habitats of cold water that
are well-oxygenated and free from toxic pollutants. Thus, poor sediment and water quality are precursors for an unhealthy benthic organism.

The presence of contaminated bottom sediment is also a concern in nearshore areas that practice dredging activities. In such areas, dredging is necessary in order to maintain channels for freighters used in shipping goods and for the operation of personal watercraft. The most significant issue in the dredging process is the disposal of toxic contaminants that may exist within the sediment. Disposal areas include offshore, upland, and confined regions of the basin. There is no assurance that contaminants will be disposed of properly, and so toxins may re-enter the aquatic system.

2.3.1 Polychlorinated Biphenyls (PCBs)

The manufacturing of PCBs occurred between the years 1929 and 1977. PCBs were utilized as an industrial safety product in processes that required high heat inputs, and/or were fire hazards. After 1977, the production of PCBs no longer continued after the discovery that “PCBs released into the environment were bioaccumulating to levels of concern in a wide range of organisms” (GLA, 1995). Following the banning of PCB production, this contaminant is still considered a critical pollutant. As seen in Table 2.1, its levels continue to exceed human health standards and its levels may pose health and reproduction problems in aquatic and terrestrial wildlife (LOLMP, 1998).
Table 2.1: Origin of PCB Loadings to Lake Ontario

<table>
<thead>
<tr>
<th>Origin of Loading</th>
<th>Loading (kg/yr)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Upstream Great Lakes Basins</td>
<td>302 kg/yr</td>
</tr>
<tr>
<td>Niagara River Basin</td>
<td>138 kg/yr</td>
</tr>
<tr>
<td>Lake Ontario Basin (Point and Non-Point Source)</td>
<td>100 kg/yr</td>
</tr>
<tr>
<td>Atmospheric Loading</td>
<td>64 kg/yr</td>
</tr>
</tbody>
</table>

(Source: LOLMP, 1998)

2.3.2 DDT and Its Metabolites

DDT is primarily a pesticide that was developed in the 1940’s. Between the years 1946 and 1972, it was the most commonly used pesticide in North America (GLA, 1995). After the discovery that “DDT and its breakdown products were causing widespread reproductive failures in various wildlife species,” its agricultural use in North America was banned (LOLMP, 1998).

The levels of DDT in the Great Lakes have decreased significantly since the banning of the pesticide. It is hypothesized that a significant amount of its remaining tributary loadings consist of atmospheric deposition (LOLMP, 1998). Thus, it is difficult to decipher the amount of DDT each source contributes to its total loading within the Lake Ontario Basin. This ambiguity explains why atmospheric and point/non-point sources of contamination exist as one measurement value in Table 2.2.
Table 2.2: Origin of DDT Loadings to Lake Ontario

<table>
<thead>
<tr>
<th>Origin of Loading</th>
<th>Loading (kg/yr)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Upstream Great Lakes Basins</td>
<td>96 kg/yr</td>
</tr>
<tr>
<td>Atmospheric Deposition and Sources within the Lake Ontario Basin</td>
<td>33.5 kg/yr</td>
</tr>
</tbody>
</table>

(Source: LOLMP, 1998)

2.3.3 Mirex

Mirex is a chlorinated organic compound that was used throughout the United States as an insecticide to control the imported fire ant. Mirex is most widely produced for its use as a flame retardant in industrial, manufacturing, and military applications (Sergeant et al., 1993). It is also widely known for its use as a pesticide. Its use and production is now banned in North America. Mirex is identified as a critical pollutant, as seen in Table 2.3, because its “levels in some Lake Ontario fish continue to exceed human health standards” (LOLMP, 1998) and for decades it has been identified as a contaminant that accumulates in aquatic ecosystems and affects sediments, fish, and birds (Scrudato and DelPrete, 1982).

Table 2.3: Origin of Mirex Loadings to Lake Ontario

<table>
<thead>
<tr>
<th>Origin of Loading</th>
<th>Loading (kg/yr)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Niagara River Basin</td>
<td>1.8 kg/yr</td>
</tr>
<tr>
<td>Oswego River</td>
<td>0.9 kg/yr</td>
</tr>
</tbody>
</table>

(Source: LOLMP, 1998)

2.3.4 Dioxins and Furans

Dioxins and furans are waste by-products from industrial processes such as paper production (Pearson et al., 1998). Processes that typically produce dioxins and furans
include the operation of incinerators and internal combustion engines. Low levels of these contaminants are also produced in forest fire and wood burning ovens (Pearson et al., 1998). Dioxins are identified as critical pollutants because “levels of these contaminants exceed human health standards in some Lake Ontario fish and because these chemicals may limit the full recovery of the Lake Ontario bald eagle, mink, and otter populations by reducing the overall fitness and reproductive health of these species” (LOLMP, 1998).

The largest known source of dioxins and furans is atmospheric deposition that accounts for approximately 5 grams per year (LOLMP, 1998). Potential non-atmospheric sources exist because of impurities in industrial chemicals (Pearson et al., 1998). Due to low levels of these contaminants in the Great Lakes Basins, quantifying their presence is a difficult task. The Niagara River, through data from sediment cores, mussels, and fish (Spottail Shiners), is identified as a source of dioxins and furans, and thus, it acts as a source of the contaminants into Lake Ontario.

2.3.5 Mercury

Mercury is a naturally occurring element that can be found within most rocks and soils. Unlike heavy metals such as copper and zinc, which are essential biological micronutrients required for the growth of organisms, mercury is considered to be extremely toxic with respect to human health and aquatic life (Ouyang et al., 2002). Mercury was initially used as an additive to paints in order to control the creation of
mildew. Presently, its most common uses include medical and dental products and thermometers, and it can be found within batteries (LOLMP, 1998).

A case study regarding Minamata Bay in Japan can be used as an example of the effects of mercury on aquatic ecosystems. During the 1950’s and 1960’s, wastes (including mercury) were created at a chemical and plastics plant and drained into Minamata Bay. This mercury was converted into absorbed methyl mercury by bacteria found residing within aquatic sediments. By 1970, 107 human deaths were attributed to mercury poisoning due to the consumption of fish and shellfish by the local population (Hodgson et al., 1997).

2.3.6 Dieldrin

Dieldrin is a former pesticide that is presently banned from use throughout North America. Through a natural breakdown process, another pesticide named Aldrin transforms into Dieldrin. Aldrin is used to control pests in soil such as termites on corn and potato crops (GPA, 2002). This is considered a critical pollutant because, as seen in Table 2.4, “Dieldrin concentrations in water and fish tissue exceed the Great Lakes Water Quality Initiative criteria throughout the lake” (LOLMP, 1998).

<table>
<thead>
<tr>
<th>Origin of Loading</th>
<th>Loading (kg/yr)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Upstream Great Lakes Basins</td>
<td>43 kg/yr</td>
</tr>
<tr>
<td>Atmospheric Deposition</td>
<td>13 kg/yr</td>
</tr>
<tr>
<td>Point and Non-Point Sources within the Lake Ontario Basin</td>
<td>9 kg/yr</td>
</tr>
</tbody>
</table>

(Source: LOLMP, 1998)
2.4 Application of the Sediment Quality Index in Lake Ontario

Contamination associated with sediments in bodies of water such as Lake Ontario impedes attempts to conserve natural ecosystems found in the Great Lakes Basin. To uncover the role that humans actively play in discharging harmful contaminants into these ecosystems, Environment Canada conducts surveys in the Great Lakes in order to identify the specific occurrence and spatial distribution of toxic contaminants in these environments. The surveys provide the ability to observe sediment concentration data within the context of sediment quality guidelines, and act as indicators of areas identified as potential risks to surrounding environments. The Canadian Council of Ministers of the Environment (CCME) has adopted Canadian Sediment Quality Guidelines in order to protect aquatic ecosystems from contaminants including PAHs, organochlorine pesticides (OCs), PCBs, and polychlorinated dibenzo-p-dioxins and dibenzofurans (PCDDs/PCDFs) (Marvin et al., 2002).

The availability of data depicting sediment contamination in Lake Ontario presents the opportunity for the evaluation of sediment quality by using the Canadian Sediment Quality Guidelines (CCME, 1999). The specific guidelines were designed as aids in the identification of potential ecosystem risk, and in order to assist in the prioritization of sediment quality concerns (Marvin et al., 2002). Using sediment contamination levels and spatial interpolation among the 70 stations in Lake Ontario, provides a means for assessing the relative risk for contamination between sites of sediment quality measurement. The basis for risk assessment is the individual site’s
departure from the ‘Threshold Effect Levels’ or ‘Probable Effect Levels’ (whichever are being used in the specific analysis). Fundamentally, the SQI is an index of sediment quality over space (Marvin et al., 2002). The SQI is effective because it is sensitive to the degree of contamination above or below a set guideline. Given this sensitivity, poorer scores should result at sites with sediment contamination exceeding the TELs and PELs. For Mirex, a contaminant for which Canadian Federal guidelines have not been published, the Ontario Provincial lowest effect level (LEL) was used (Marvin et al., 2002). The ‘LEL’ for the province of Ontario can be defined as “a level of contamination that has no effect on the majority of sediment-dwelling organisms and where the sediment is clean-to-marginally polluted” (Persaud et al., 1993 in Marvin et al., 2002).

2.5 Spatial Interpolation – Kriging

Kriging techniques were initially developed by a South African mining geologist named D.G. Krige (Bailey et al., 1995). Kriging methods utilize statistical models that incorporate autocorrelation among a group of measured points to create prediction surfaces (Johnston et al., 2001). Specifically, weights are assigned to measurement points on the basis of distance in which spatial autocorrelation is quantified in order to weight the spatial arrangement of measured sampling locations (Johnston et al., 2001). By accounting for statistical distance with a variogram model, as opposed to Euclidean distance utilized in deterministic interpolation, customization of the estimation method to a specific analysis is possible. Isaaks and Srivastava (1989) state that if the pattern of spatial continuity of the data can be described visually using a variogram model, it is
difficult to improve on the estimates that can be derived in the kriging process. Furthermore, kriging accounts for both the clustering of nearby samples and for their distance to the point to be estimated (Isaaks and Srivastava, 1989). Given the statistical properties of this method, measures of certainty or accuracy of the predictions can be produced in the cross-validation process that will be documented in a later section. It is arguable that kriging is the optimal interpolation method on the basis of its functionality and its ability to assess error statistically, when forming predicted surfaces. In the following sub-sections, kriging models including ordinary, simple, universal, indicator, co-kriging, and probability kriging are outlined.

2.5.1 Ordinary Kriging

Ordinary kriging is the most flexible kriging model because it functions under the assumption that the mean $u$ is an unknown constant, and thus, the random errors at the data locations are unknown (Johnston et al., 2001). Ordinary kriging is most appropriate for data that have a spatial trend and, furthermore, this system can easily be applied to block (average) estimation from point estimation. Thus, the average of a specific number of point estimates can be represented as a direct block estimate if one wishes to group the data values (Isaaks and Srivastava, 1989).
2.5.2 Simple Kriging

The assumption of the model when using simple kriging is similar to ordinary kriging. However, the mean $u$ in the equation refers to a known constant, rather than an unknown constant. Furthermore, it is based on the assumption of second order stationarity (Olea, 1991 in Burrough and McDonnell, 1998). Second order stationarity refers to the assumptions that the data come from a random process with a constant mean, and that spatial covariance only depends on the distance and direction separating any two locations (Johnston et al., 2001). If it is assumed that the mean $u$ is a known constant, then the random errors at the data locations are also known. If the random error at each data location is known, the estimation of autocorrelation among the data locations is optimal (Johnston et al., 2001). However, the assumption of knowing the exact mean is often unrealistic, (Johnston et al., 2001) and data from the physical environment is often too restrictive to assume second order stationarity. In such datasets, ordinary kriging is usually applied (Burrough and McDonnell, 1998).

2.5.3 Universal Kriging

Universal kriging follows a similar model to ordinary and simple kriging. However, the mean $u$ is replaced by empirical regression transfer models or deterministic functions such as second-order polynomials to form the trend (Burrough and McDonnell, 1998). The random errors produced using this method are obtained by subtracting the
deterministic function from the data representing the original locations (Johnston et al., 2001). Using this model, autocorrelation is modeled from the random errors.

2.5.4 Indicator Kriging

Indicator kriging is based on the following model:

\[ I(s) = u + e(s) \]  

where \( u \) is assumed to be an unknown constant, \( e(s) \) represents the random error at the location \( s \), and \( I(s) \) is a binary variable (Johnston et al., 2001). This model works similarly to ordinary kriging, however, binary data can be created through thresholds such as the PEL in this analysis. Due to the indicator variables being 0 or 1, the interpolation maps display the probabilities of a specific variable (e.g., Mercury) being in a class that is indicated by the binary number 1. An example of its use is an analysis used to produce maps of the probability that the lead concentration at particular sites in the city of Sydney, Australia were greater than local regulatory guidelines (Markus and McBratney, 2001).

2.5.5 Cokriging

In this analysis, contamination levels for 32 variables were measured at each of the 70 locations throughout Lake Ontario. However, in alternative analyses, one variable may not be measured as frequently due to reasons such as a lack of funding. If these variables are spatially correlated with one another, co-kriging can be used to apply the
spatial variation of one variable to aid in the mapping of a second (Burrough and McDonnell, 1998). However, Cokriging requires a larger amount of estimation, including the estimate of autocorrelation for each variable and its cross-correlations. Each estimation adds variability to the prediction (Johnston et al., 2001).

2.5.6 Probability Kriging

Probability kriging is a compilation of both indicator kriging and co-kriging. If this method is used in analysis, a prediction map is produced displaying the probability that a specific attribute exceeds a set threshold (Burrough and McDonnell, 1998). Probability kriging strives to perform the same predictions as indicator kriging, however, co-kriging is used in order to improve the accuracy of the predictions (Johnston et al., 2001).
Chapter 3: Methodology

In order to predict the sediment contamination levels at unknown locations throughout Lake Ontario, it was necessary to assess deterministic and geostatistical methods of spatial interpolation. In completing this task, deterministic interpolation methods including Inverse Distance Weighting (IDW), Radial Basis Functions (RBF), Global Polynomial Interpolation (GPI), and Local Polynomial Interpolation (LPI) were investigated relative to geostatistical (kriging) methods. In order to compare these methods, SQI scores were utilized for interpolation between the 70 sampling sites throughout Lake Ontario.

Identification of regions with sediment that is frequently threatened or impaired was performed using the Sediment Quality Index. The SQI scores were then used as general indicators of sediment contamination. Existing kriging methods were examined and an assessment of the optimal kriging method was performed using the Environmental Systems Research Institute’s (ESRI) ArcGIS 8.1 software (ESRI, 2001). The analysis in this research paper was limited to this software due to its efficiency and the large quantity of spatial interpolations necessary to accomplish the desired objectives. Furthermore, using a single software package for all interpolations reduces the possibility of discrepancies between results produced using alternative methods. Prediction maps featured as ArcGIS contour surfaces were created for each individual contaminant. It was on the basis of the resulting contour prediction maps and cross-validation measures that
the optimal interpolation method for indicating the sediment contamination in Lake Ontario could be determined.

3.1 Calculation of the Sediment Quality Index (SQI)

In order to calculate the SQI, the specific body of water for which the index applies and the specific variables and objectives (contaminant concentrations exceeding their PELs) applying to the study must be defined. The Sediment Quality Index, which computes an index score on a per site basis, with no grouping of sites (Marvin et al., 2002), is based on a two-component equation including ‘scope’ and ‘amplitude’ only. A third ‘frequency’ component, used to calculate the Canadian Water Quality Index (CWQI), is excluded in the calculation of the SQI because frequency (the amount of tests exceeded within a group of sites) is equivalent to scope when applied to a single site with no temporal data (Marvin et al., 2002). Therefore, ‘frequency’ would be redundant if applied in the SQI calculation, in which the components are described as:

\[ \text{F}_1 = \frac{\text{Number of failed variables}}{\text{Total number of variables}} \times 100 \]  

\[ \text{F}_1 = (\text{Number of failed variables} / \text{Total number of variables}) \times 100 \quad (2) \]
F$_2$ – The ‘amplitude’ is a representation of the amount by which failed test values do not meet their objectives (exceed the PEL for contaminant concentration in sediment) (Marvin et al., 2002). Three steps are necessary to calculate F$_2$:

Step 1: First, excursions are calculated. The term ‘excursion’ refers to the number of times an individual concentration is greater than the objective. For the case in which the test value must not exceed the objective, the calculation is as follows:

$$\text{Excursion}_i = (\frac{\text{Failed Test Value}_i}{\text{Objective}_j}) - 1$$  \hspace{1cm} (3)

For the case in which the test value must not fall below the objective, the calculation is:

$$\text{Excursion}_i = (\frac{\text{Objective}_j}{\text{Failed Test Value}_i}) - 1$$  \hspace{1cm} (4)

Step 2: The collective amount by which individual tests are out of compliance is calculated by summing the excursions of individual tests from their objectives and dividing by the total number of tests (this includes both those meeting objectives and those not meeting objectives). This variable is referred to as the ‘normalized sum of excursions, or nse and can be calculated with the following equation:

$$nse = \frac{\text{the summation of excursion}_i}{\# \text{ of tests}}$$  \hspace{1cm} (5)
Step 3: Using excursions, the amplitude, $F_2$ is calculated. $F_2$ can be calculated utilizing an asymptotic function, which scales the normalized sum of the excursions from objectives to produce a range between $0 – 100$. The calculation is as follows:

$$F_2 = \left(\frac{nse}{0.01} * nse + 0.01\right)$$

(6)

Once these two factors have been obtained, **The Sediment Quality Index (SQI)** can be calculated as:

$$SQI = 100 - \sqrt{(F_1)^2 + (F_2)^2 / 1.414}$$

(7)

The value 1.414 is used to normalize the resulting values to a range between 0 and 100. This value is generated because $[100^2 + 100^2]^{0.5} = 141.4$ (Marvin et al., 2002). According to this scale, a water quality of 100 is the ‘best’ and a water quality of 0 is the ‘worst’ (CCME, 2001).

$F_1$ and $F_2$ are combined to produce a single value (between 0 and 100) that describes sediment quality in the following categories:

**Excellent**: (SQI Value of 95 – 100) – sediment is devoid of any contaminant-related impairment and is indicative of ambient environmental background conditions. Index values within this range are achieved when practically all measurements fall within the guideline values.
**Good**: (SQI Value 80 – 94) – only a minor degree of sediment impairment is indicated. Most measurements fall within guideline values and rarely deviate from ambient environmental background levels.

**Fair**: (SQI Value 60 – 79) - sediment quality is usually protected but occasionally threatened or impaired. Some measurements deviate from ambient environmental background levels.

**Marginal**: (SQI Value 45 – 59) - sediment quality is frequently threatened or impaired. Some measurements deviate from ambient environmental background levels.

**Poor**: (SQI Value 0 – 44) - sediment quality is almost always threatened or impaired. Most measurements deviate substantially from ambient environmental background levels (Marvin et al., 2002).

‘Categorization’ is the term for the assignment of the SQI values to the four categories for any particular contaminant and is based on three factors including the most reliable information available for each specific application, leading expert opinions, and the expectation of sediment quality by the general public (CCME, 2001).

The basic SQI formula encompasses all key components of sediment quality, can be easily calculated, and is flexible in various applications. These applications include direct comparisons among sites that employ identical variables and objectives. Comparison among sites can be problematic when variables and objectives vary at each site. If this issue arises, it is most beneficial to compare sites on the basis of their ability to meet relevant objectives.
3.2 Spatial Interpolation: Deterministic Methods

When utilizing deterministic methods for spatial interpolation, contour surfaces are created from measured points (i.e. stations for the measurement of sediment contamination) on the basis of their extent of similarity (Inverse Distance Weighting) or the extent to which the surface is smoothed (Radial Basis Functions) (Johnston et al., 2001). Furthermore, deterministic methods can be separated into global and local groups. Global interpolation techniques utilize an entire dataset in order to calculate predictions, whereas local techniques use small regions (known as neighbourhoods) that are designated within the larger spatial areas in order to calculate predictions (Johnston et al., 2001). Another distinction for deterministic interpolation methods is whether they are exact or inexact interpolators. Exact interpolators are used to predict values that are identical to the measured values at sampling locations, whereas inexact interpolators predict values that differ from values measured at sampling locations (Johnston et al., 2001). Table 3.1 outlines the deterministic methods for spatial interpolation that are offered within the Geostatistical Analyst extension of ArcGIS.

Table 3.1: Deterministic Methods for Spatial Interpolation

<table>
<thead>
<tr>
<th></th>
<th>Global Interpolator</th>
<th>Local Interpolator</th>
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</thead>
<tbody>
<tr>
<td><strong>Exact Interpolator</strong></td>
<td></td>
<td>Inverse Distance Weighting, Radial Basis Functions</td>
</tr>
<tr>
<td><strong>Inexact Interpolator</strong></td>
<td>Global Polynomial</td>
<td>Local Polynomial</td>
</tr>
</tbody>
</table>

In order to predict the degree of sediment contamination in areas other than the 70 measurement sites for sediment core sampling in Lake Ontario, deterministic methods
featured in Table 3.1 were each assessed independently utilizing the Sediment Quality Index values for interpolation.

3.2.1 Inverse Distance Weighted Interpolation (IDW)

IDW is an interpolation method that assumes that entities located in close proximity to one another are more similar than those located farther apart. Measurements of contaminant concentrations closer to the prediction locations are weighted accordingly and have a higher degree of influence on the final predictions than sites located at larger distances. Thus, the local influence of each measured point decreases with distance (Isaaks and Srivastava, 1989). IDW interpolation is a common method used in Geographic Information Systems (GIS) in order to create raster overlays from point data (Burrough and McDonnell, 1998). Since it is identified as an exact interpolator, lines can be woven across interpolated values creating either vector contour maps or raster shaded maps. The term vector refers to the representation of spatial data by points, lines, and polygons. The term raster refers to a regular grid of cells covering an area (Burrough and McDonnell, 1998).

Given the appropriate dataset, IDW interpolation can prove to be a powerful prediction tool, however, shortcomings do exist with this method. First, IDW must be an exact interpolator due to the circumstance that if the value for an interpolated point coincides with that of a data point, the unsmoothed value must be replaced (Burrough and McDonnell, 1998). Furthermore, the form of the contour map is dependent on how the
sampling sites are clustered and the presence of outliers. IDW interpolations often feature bulls-eye patterns occurring around sampling locations with values differing significantly from their surrounding points (Burrough and McDonnell, 1998). Finally, IDW interpolation features no methods for testing the quality of predictions. Therefore, the only method for assessment of the accuracy of predictions is by performing additional observations (Burrough and McDonnell, 1998).

The formula necessary to perform IDW interpolation in this analysis is as follows:

\[ Z(s_o) = \sum_{i=1}^{N} \lambda_i Z(s_i) \]  

(8)

In this formula, \( Z(s_o) \) represents the unknown value at a location \( (s_o) \); \( N \) represents the number of measured sample points that surround a prediction location utilized in the prediction; \( \lambda_i \) represents the assigned weights for each measured point used in the interpolation. Given the nature of IDW, these weights will decrease with increased distance; \( Z(s_i) \) represents the observed value at a location \( (s_i) \) (Johnston et al., 2001).

In order to perform any interpolation, two critical decisions must be made. First, the weighting must be assigned. In the Geostatistical Wizard, within the Geostatistical Analyst extension of ArcGIS 8.1, this is called the ‘power function.’ Second, the neighbourhood must be defined by choosing an appropriate shape and number of neighbours.

To attain the optimal power parameter, a method named ‘cross-validation’ was used in order to determine the root-mean-square-prediction error (RMSPE) for the
prediction surface. The RMSPE is a statistic that quantifies the error of the prediction surface. Given IDW interpolation, as the distance between points increases, the weight assigned to the neighbours is reduced by a factor of ‘p’ (Isaaks and Srivastava, 1989). Cross-validation is a method in which measured points “are removed and compared to the predicted value for the specific location” (Johnston et al., 2001). The RMSPE is a summary statistic produced by cross-validation in order to quantify error of the prediction surface. The optimal power parameter is one that produces the minimum RMSPE statistic.

In determining the optimal search neighbourhood for the analysis, it was necessary to consider the possible directional influences on how the data were weighted. As discussed in Chapter 2, the majority of sedimentation in Lake Ontario stems from the Niagara River. Due to factors including shear stress, buoyant discharge conditions, inertia, and Coriolis acceleration, a surface plume is created in an easterly direction. Thus, the search neighbourhood was adjusted to an elliptical shape, orienting the major axis parallel to the eastern direction of the sedimentation. This is an important process in IDW interpolation because locations east of a specific measured point in Lake Ontario will be more similar at further distances than points that are perpendicular to the specific sedimentation pattern (Johnston et al., 2001). The second step in determining the neighbourhood involves defining the number of data points within it that will be used for creation of the prediction surface. In this analysis, a maximum of five neighbours, and a minimum of one neighbour, was used in each prediction. This particular number of neighbours was rendered appropriate given the total number of sampled sites and their
dispersion throughout Lake Ontario. Experimental processes testing various
neighbourhood search areas by means of identifying the most optimal cross-validation
results, supported the decision for five data locations within each neighbourhood. Five
data locations within each neighbourhood resulted in the lowest RMSPE results after an
appropriate model was fit to the semivariogram. In order to include a balanced
representation of points in all directions, the ellipse was divided into four sectors. Each of
these sectors included the selection of an equal number of points. Additional reasoning
for partitioning the elliptical neighbourhood is because the sediment contamination levels
for Lake Ontario have been collected in a grid pattern. Division of the neighbourhood
into four sectors reduces the possibility that the five nearest neighbours included in the
interpolation are located along one transect (Johnston et al., 2001).

3.2.2 Radial Basis Functions (RBF)

Radial Basis Functions are exact interpolation methods in which a given surface is
forced through each sample value that is measured in the specific analysis (Johnston et
al., 2001). The goal of RBF is to fit a line through a group of sampled values, while
minimizing the overall curvature of the surface (Sutton et al., 1998). The equations for
this curve fitting have been termed ‘kernel functions’. The Geostatistical Analyst offers
five kernel functions: thin-plate spline, spline with tension, completely regularized spline,
multiquadric function, and inverse multiquadric spline. The formula necessary to perform
RBF interpolation in this analysis is as follows:
\[ Z(s_0) = \sum \omega_i \phi(\|s_i - s_0\|) + \omega_{n+1} \]  

(9)

In this formula, \( Z(s_0) \) represents the unknown value at a location \( s_0 \), \( \phi \) is a radial basis function, \( \|s_i - s_0\| \) is Euclidean distance between the prediction location \( s_0 \) and each data location \( s_i \) and \( \omega_i : i = 1,2,\ldots,n+1 \) are weights to be estimated (Johnston et al., 2001).

Given that RBF and IDW are exact interpolators (used to predict values that are identical to the measured values at sampling locations), they differ from one another in that RBF will predict values above the maximum measured value or below the minimum measured value in an analysis, whereas IDW will not (Johnston et al., 2001). However, similar to IDW interpolation, RBF features no methods for testing the quality of predictions.

An RBF interpolation is typically employed if one is predicting smooth surfaces from an abundance of data points (Sutton et al., 1998). RBF interpolation is not an appropriate method when “large changes in the surface values occur within a short horizontal distance” (Johnston et al., 2001) or when the sample data are prone to error or uncertainty. Typically, surfaces such as elevation that vary mildly render good interpolation results. This analysis incorporates sediment contamination data that features significant changes within short horizontal distances. Thus, RBF may produce inaccurate results using this dataset.
In order to create the optimal predicted surface using RBF interpolation, the five kernel functions were tested. The optimal kernel function was identified as the ‘Multiquadric,’ on the basis of the lowest RMSPE value. The identical search neighbourhood was utilized for the RBF predictions as was used in IDW interpolation.

3.2.3 Global Polynomial Interpolation (GPI)

Global Polynomial Interpolation involves fitting a polynomial or mathematical function of spatial coordinates of the sample sites, to the observed data at these sites, using least square regression (Bailey et al., 1995). Using this technique, the surface being fit to the sample sites rarely passes through them, defining GPI as an inexact interpolator (Johnston et al., 2001). However, the number of sample sites above the surface medium is usually equivalent to the number below it. GPI attempts to create smooth surfaces, which mathematically depict trends that vary gradually over the study area. GPI is also used to “examine or remove the effects of long-range or global trends,” (Johnston et al., 2001) a process called trend surface analysis (Bailey et al., 1995). The data samples in this analysis feature a global trend do to the east-west circulation patterns of water in Lake Ontario and the eastern sedimentation pattern stemming from the Niagara River. The major flaw with GPI is that more complex polynomials (i.e. fourth order or quartic) are prone to outliers. The term outliers refers to excessively high and/or low values predicted over the study area. These outliers are especially a problem near the edges of a study area. In order to predict sediment contamination levels throughout Lake Ontario using GPI, it was necessary to define the polynomial function that would produce the lowest
RMSPE. The formula necessary to perform GPI interpolation in this analysis is as
follows:

\[ Z(x_i, y_i) = \beta_0 + \beta_1 x_i + \beta_2 y_i + \varepsilon(x_i, y_i) \]  

(10)

In this formula, \( Z(x_i, y_i) \) represents the datum at location \( (x_i, y_i) \), \( \beta \) are parameters, and \( \varepsilon(x_i, y_i) \) is a random error (Johnston et al., 2001).

3.2.4 Local Polynomial Interpolation (LPI)

Local Polynomial Interpolation involves fitting a succession of polynomial
functions to specified neighbourhoods over a surface area, rather than fitting a single
polynomial to an entire surface (Johnston et al., 2001). The Geostatistical Analyst allows
the user to take advantage of the *Searching Neighbourhood* dialog box in order to define
the shape, maximum and minimum number of points to use, along with specifying a
sector configuration for the given analysis. The reason that these options exist is in order
to exercise an interpolation method that considers local variation in the sample sites.

Local Polynomial Interpolation is often applied to case studies in the domain of
geological sciences because the variables of interest often have a short-range variation
(Johnston et al., 2001). The most accurate LPI surface is that producing the lowest
RMSPE. The formula necessary to perform LPI interpolation in this analysis is identical
to that used in GPI. The difference is that data are used within localized windows, rather
than the entire data set (Johnston et al., 2001).
3.3 Spatial Interpolation: Geostatistical Methods

In preceding sections, deterministic interpolation methods were detailed by which the smoothness of predicted surfaces were created on the basis of mathematical formulas. Geostatistical interpolation methods utilize statistical models incorporating autocorrelation or statistical relationships among a group of measured points to create prediction surfaces (Johnston et al., 2001). To achieve these prediction maps, kriging is used.

3.3.1 Regionalized Variable Theory (RVT) and Kriging

In order to describe kriging as an interpolation technique, it is necessary to outline the Regionalized Variable Theory (RVT). Kriging is based on the RVT, which assumes that the spatial variation of a variable represented at specific measurement locations is statistically homogeneous throughout the defined surface (Buttner et al., 1998). RVT functions under the assumption that the spatial variation of a particular variable is expressed as the sum of three specific components forming the following equation:

\[ Z(x) = m(x) + e'(x) + e'' \]  

(11)

In this equation, the value for the variable Z at the spatial coordinate x is given by the sum of: \( m(x) \) representing a deterministic function that describes the structural component of Z at x, \( e'(x) \) representing the regionalized variable described as random, but spatially correlated, and \( e'' \), a residual term or spatially correlated random noise (Isaaks and Srivastava, 1989).
3.3.2 Ordinary Kriging

The equation used to perform ordinary kriging is:

\[ Z(s) = u + e(s) \] (12)

where \( Z(s) \) represents the value for the unknown variable at a spatial location \( s \), \( u \) represents an unknown constant mean for the data (thus no trend), and \( e(s) \) represents the random errors (Johnston et al., 2001). When considering the random errors, it is important to note that the random process is intrinsically stationary. Intrinsic stationarity is defined as “an assumption that the data come from a random process with a constant mean, and a semivariogram that only depends on the distance and direction separating any two locations” (Johnston et al., 2001). A discussion regarding the semivariogram will follow in the next two sections.

The next step is calculating the predictor, which is the weighted sum of the data, featured in the following equation:

\[ Z(s_0) = \sum_{i=1}^{N} \lambda_i Z(s_i) \] (13)

Where \( Z(s_i) \) is the measured value at the \( i \)th location, \( \lambda_i \) is an unknown weight for the measured value at the \( i \)th location, \( (s_0) \) are the coordinates of the prediction location, and \( N \) represents the number of measured points used in the prediction of a value for an unknown location (Johnston et al., 2001). This predictor is similar to IDW interpolation, although weighting is dependant on distances from the prediction location, along with the semivariogram, and the spatial relationship of measured values surrounding the prediction location.
It is necessary to ensure that the prediction is unbiased. When predictions are made at numerous locations, some predictions will be greater than the actual values, and some will be below the values. On average, the difference between the actual and predicted values should be zero.

3.3.3 The Empirical Semivariogram

It is necessary to create an empirical semivariogram in order to examine the structure of the data. A semivariogram is a graph which plots half the difference squared between pairs of locations (the averaged semivariogram values) on the y-axis, relative to the distance that separates them on the x-axis (Johnston et al., 2001). Averaged values can be used due to the assumption of intrinsic stationarity. The ability to plot all pairs in a manageable time frame is a difficult task. ArcGIS utilizes a technique in which pairs of locations are grouped based on specified ranges of distances and directions. This process is referred to as ‘binning,’ by which the average empirical semivariance for all pairs of points is recorded. Another binning method (not used by ArcGIS 8.1) is based on grouping pairs of locations into radial sectors. The word “empirical” means that a certain quantity is dependant on data, observations, or experiment and is not a model. In order to calculate the distance between two locations, the formula for Euclidean distance is used:

$$d_{ij} = \sqrt{(x_i - x_j)^2 + (y_i - y_j)^2} \quad (14)$$

In order to calculate the empirical semivariance (y-axis), the following formula is utilized:
Semivariance = 0.5 * average[(value at location I – value at location j)²]

(15)

(Johnston et al., 2001).

3.3.4 Fitting A Model to the Empirical Semivariogram

Once the average semivariance values are plotted against the average distances of the ‘bins,’ it is necessary to fit a model to the empirical semivariogram. Figure 3.1 is a semivariogram fit with an exponential model representing SQI scores.

**Semivariogram: Exponential Model Fitted to SQI Measurements**

Figure 3.1 – Exponential Model Fitted to SQI Measurements

Before a model could be fit to the empirical semivariogram, it was necessary to determine whether the Lake Ontario sediment contaminants were normally distributed. If the distribution was skewed, it would have been necessary to perform the appropriate
transformations. A normal distribution is not necessary to obtain prediction maps in the ordinary kriging process. However, kriging is the best predictor among unbiased predictors when data are normally distributed (Johnston et al., 2001).

The semivariogram accounts for spatial autocorrelation in the data; however, it lacks an explanation for autocorrelation in various directions and does not ensure that positive kriging variances will result from the kriging predictions. Thus, it is necessary to fit a model in order to determine semivariogram values at different distances. Within the Geostatistical Analyst, the following functions exist in order to model the empirical semivariogram: circular, spherical, tetraspherical, pentaspherical, exponential, Gaussian, rational quadratic, Hole Effect, K-Bessel, J-Bessel, and stable. In this analysis, the spherical, exponential, and Gaussian models are used for reasons explained in section 4.3.

The Spherical model generally depicts decreasing spatial autocorrelation or an increase of semivariance. At a specific point in the empirical variogram, the model plateaus, no longer displaying autocorrelation. The distance at which the model initially levels out is known as the ‘range’. The semivariogram value that the model displays at the range is known as the ‘sill’ (Bailey et al., 1995). Finally, ‘the nugget effect’ occurs when the chosen model does not cross through the origin, but intercepts the y-axis at some value greater than zero. It can be attributed to either measurement error or variation when distances are smaller than assigned sampling intervals (Johnston et al., 2001).
The Gaussian model has a parabolic shape near the origin and is used if the semivariogram depicts autocorrelation at short distances between data points. This model incorporates this shape in order to make use of the closest samples in the prediction (Isaaks and Srivastava, 1989).

The exponential model displays a decrease in spatial autocorrelation with distance. An autocorrelation value of zero only occurs at an infinite distance (Johnston et al., 2001). In order to calculate the semivariance at a specific distance, the following formula is used:

\[ \text{Semivariance} = \text{Slope} \times \text{Distance} \]  
\[ (16) \]

in which the slope refers to the specific slope of the model used and distance is equal to the distance between pairs of locations (Johnston et al., 2001). At this point, it is necessary to solve for \( \lambda \) (the weights to assign to the measured values surrounding the prediction location) in which the matrix formula for ordinary kriging must be used:

\[ g = \Gamma \ast \lambda \]  
\[ (17) \]

in which \( \Gamma \) represents the ‘gamma matrix’ and \( g \) represents the g-vector which is applied to the unmeasured location that is being predicted (Johnston et al., 2001). The g-vector is calculated using the aforementioned ‘semivariance’ equation. The distances to be used in this calculation are dependant on the distance from the unmeasured location to the measured points defined by the specific search neighbourhood for the interpolation.

In order to calculate the value at an unknown location, the final step is the following equation:
Kriging Predictor = \Sigma ( \lambda_i \ast i) \quad (18)

where \( \lambda \) represents the weight for each measured value and \( i \) represents the measured value (Johnston et al., 2001).

3.3.5 Anistrophy: Directional Influence of Autocorrelation

Anistrophy can be defined as “a property of a spatial process or data where spatial dependence or autocorrelation changes with both the distance and the direction between two locations” (Johnston et al., 2001). The cause for such directional influences are often unidentifiable, therefore it is modeled as random error. Anistropic influences can be quantified and accounted for using ArcGIS 8.1 software.

3.3.6 Determining the Search Neighbourhood

As measured data points become located at greater distances from the prediction locations, they become less spatially autocorrelated with one another. The ability to set the specific size of the search neighbourhood, and assign the specific number of measured locations to be used in making a prediction, allows for the elimination of locations that have minimal influence on the overall prediction. Furthermore, a search neighbourhood customized to fit the spatial arrangement of a specific dataset increases the speed at which predictions can be made because sampling locations that are not spatially autocorrelated are excluded from the prediction process.
In this kriging analysis, the same search neighbourhood was used as in the IDW interpolation. An elliptical search neighbourhood was chosen to account for the eastern sedimentation patterns stemming from the Niagara River. The major and minor axes were assigned values of 1 and 0.5 respectively, and the ellipse was divided into four sectors in which the maximum number of neighbours was limited to 5 and the minimum was limited to 1. This is similar to Buttner et al., (1998) who used ordinary kriging to predict the spatial distributions of 12 elements in an acidic mining lake using an isotropic search neighbourhood including the ten nearest neighbours from each prediction location. A ten neighbour search neighbourhood was appropriate because a total of 47 sampling sites were measured within three basins approximately 120 to 140 metres in width and 250 to 400 metres in length (Buttner et al., 1998). The ratio of sampled sites with respect to the surface area of the lake was the reason for a search neighbourhood featuring 10 neighbours. The shorter distances between sampling locations increased the probability of autocorrelation between their contamination concentrations. The ratio between the surface area of Lake Ontario and the grid of 70 sampling locations separated by intervals of 30 km is the reason the maximum number of neighbours was limited to 5 in this analysis. Additional neighbours would surpass the range of the semivariogram, and thus, they would lack spatial autocorrelation.

3.3.7 Cross-Validation: Identification of the Best Model

In order to identify the degree of accuracy that the semivariogram parameters and the search neighbourhood possess in predicting the unknown locations, the Geostatistical
Analyst can be used to perform cross-validation. Cross-validation is a method that removes each measured location one at a time in order to predict their values on the basis of the measured values in the entire dataset. On the basis of cross-validation results, accuracy can be determined regarding the chosen model and search neighbourhood for each prediction. This method was used to quantify the uncertainty of 2,3,7,8-tetrachlorodibenzo-p-dioxin concentrations in the Passaic River in New Jersey, after an indicator kriging interpolation was performed (Barabas et al., 2001). This study is significant because the cross-validation process provided sufficient indication regarding the accuracy of the prediction surface produced.

In order to visualize these results, the Geostatistical Analyst offers a scatter plot of predicted values against measurement values, an error plot in which true values are subtracted from the predicted values, and a standard error plot in which the measurement values are subtracted from the predicted values and divided by the estimated kriging standard errors. These three plots, which depict how well the kriging process is predicting at unknown locations, are featured as Figures 3.2 through 3.4. For example, Figure 3.2 representing predicted vs. measured values, shows a blue line (best fit) of the predicted values superimposed over the measured values (red dots). The black dashed line represents the 1:1 line. Measured values are not scattered around this line because kriging tends to under-predict large values and over-predict small values (Johnston et al., 2001).
Predicted Vs. Measured Values

Figure 3.2 – Predicted (blue line) Vs. Measured (red dots) Values

Figure 3.3 is similar to Figure 3.2, however true values are subtracted from predicted values in order to show the error generated from the interpolation.

Error Plot

Figure 3.3 – Error Plot

In Figure 3.4, the predictions are assessed relative to whether they are centred on the measurement locations. It is apparent that this prediction pertaining to SQI scores throughout Lake Ontario is relatively unbiased.
A fourth plot identified as Figure 3.5 below, illustrates the difference between the
predicted and measured values divided by the estimated kriging standard errors and the
corresponding quantiles from a standard normal distribution (Johnston et al., 2001). This
plot indicates a relatively normal distribution because the measured values stray
minimally from the 1:1 line.

Cross-validation also provides values including Mean Prediction Error (MPE),
Standardized Mean Prediction Error (SMPE), Root-Mean Squared Prediction Error
(RMSPE), Average Standard Error (ASE) and Standardized Root-Mean-Squared
Prediction Errors (SRMSPE), which assess the accuracy of the chosen model. A good
model will calculate MPE and SMPE values near a value of zero to show that predictions are unbiased or are centred on the measured locations. Additionally, low RMSPE values identify that predictions are close to their true values. The ASE values are used to assess the variability in the predictions from the measurement values. Therefore, the average standard error must be similar to the root-mean square prediction error in order to correctly assess the variability in the prediction. For instance, if the ASE value is greater than the RMSPE value, the variability of the prediction is being overestimated. The SRMSPE value also provides another method to assess variability. If the prediction standard errors are valid, the SRMSPE values should be close to 1 (Johnston et al., 2001). However, standardized RMSPE values greater than 1 translate to an underestimation in the variability of the predictions.
Chapter 4: Results and Discussion

This chapter first examines whether the Sediment Quality Index is a satisfactory measure for areas within Lake Ontario where sediment quality is frequently threatened or impaired. This assessment is performed in relation to the critical pollutants exceeding federal guidelines at the sampled sites. Second, deterministic interpolation methods including IDW, RBF, GPI, and LPI are assessed relative to the ordinary kriging model. Third, an assessment of existing kriging models including ordinary, simple, universal, indicator, co-kriging, and probability kriging is conducted. Finally, the best interpolation model is used for predicting the spatial distribution of 32 individual contaminants within Lake Ontario.

4.1 Sediment Quality Index as a General Indicator of Sediment Contamination

In order to determine whether the SQI is a satisfactory measure for determining the areas in aquatic ecosystems where sediment quality is frequently threatened or impaired, the critical pollutants including PCBs, DDT and its metabolites, Mirex, dioxins and furans, mercury, and Dieldrin were assessed relative to the SQI scores. Figure 4.1 displays four measured locations throughout Lake Ontario that are classed are classified as marginal with SQI values ranging from 45 - 59. Since critical pollutants are of highest concern because of their negative impacts on Lake Ontario as an aquatic ecosystem, the locations labelled as frequently threatened or impaired based on low SQI values should indicate the presence of the critical pollutants at high concentrations (near or above the
Figure 4.1 - Lake Ontario Contaminated Sediment Hotspot Locations
Probable Effect Level). Table 4.1 indicates the relationship between low SQI scores and high concentrations of critical pollutants. The reason Mirex was not included in this table is because values of ‘0’ were measured at all sites for this contaminant. Measurement error must account for these 0 ng/g values for two reasons: (1) Mirex is identified as a Critical Pollutant in the Lake Ontario Basin and (2) Concentrations of Mirex in sediment are typically minute, therefore, traces of the pollutant may not have been identified.

**Table 4.1: SQI and Critical Pollutants**

<table>
<thead>
<tr>
<th>Critical Pollutant</th>
<th>Threshold Effect Level</th>
<th>Probable Effect Level</th>
<th>Station 22</th>
<th>Station 33</th>
<th>Station 40</th>
<th>Station 69</th>
</tr>
</thead>
<tbody>
<tr>
<td>PCBs</td>
<td>34.1 ng/g</td>
<td>277 ng/g</td>
<td>177 ng/g</td>
<td>208 ng/g</td>
<td>232 ng/g</td>
<td>167 ng/g</td>
</tr>
<tr>
<td>DDT</td>
<td>1.19 ng/g</td>
<td>4.77 ng/g</td>
<td><strong>19.58 ng/g</strong></td>
<td><strong>10.28 ng/g</strong></td>
<td><strong>15.08 ng/g</strong></td>
<td>1.88 ng/g</td>
</tr>
<tr>
<td>Dioxins/Furans</td>
<td>0.85 ng/g</td>
<td>21.5 ng/g</td>
<td>0 ng/g</td>
<td><strong>244.5 ng/g</strong></td>
<td><strong>243.6 ng/g</strong></td>
<td>0 ng/g</td>
</tr>
<tr>
<td>Mercury</td>
<td>0.17 ug/g</td>
<td>0.486 ug/g</td>
<td><strong>1.38 ug/g</strong></td>
<td><strong>1.00 ug/g</strong></td>
<td><strong>0.78 ug/g</strong></td>
<td><strong>0.51 ug/g</strong></td>
</tr>
<tr>
<td>Dieldrin</td>
<td>2.85 ng/g</td>
<td>6.67 ng/g</td>
<td>2.27 ng/g</td>
<td>2.35 ng/g</td>
<td>2.93 ng/g</td>
<td>1.17 ng/g</td>
</tr>
<tr>
<td>Total Pollutants Exceeding PEL</td>
<td></td>
<td></td>
<td>13</td>
<td>15</td>
<td>14</td>
<td>17</td>
</tr>
</tbody>
</table>

Given this analysis, it is evident that the SQI considers two factors in the categorization of sediment quality: (1) the total number of contaminants exceeding the TEL and the PEL; and (2) the amount by which each threshold is exceeded. For example, the results for Station 33 display three critical pollutants greatly exceeding the PEL (highlighted in bold font), and one greatly exceeding the TEL. Additionally, a total 15 of the 32 contaminants were measured as exceeding the PEL, thus creating a low SQI score. In comparison, Station 69 features one critical pollutant exceeding the PEL, however a
total of 17 contaminants were found to exceed this measure at the location. On the basis of these results, the SQI can be considered a satisfactory measure of sediment contamination in Lake Ontario.

With the exception of Hamilton Harbour, the sampling locations where sediment quality is frequently threatened or impaired are located within the deep lake basins. The explanation for the location of high sediment contamination levels in these areas is described in the following process. First, the major factor contributing to the loading of pollutants into Lake Ontario is surface runoff. The clearing of original forested areas for agricultural purposes and logging, results in less soil stability and erosion/runoff into Lake Ontario and its tributaries (GLA, 1995). These processes increase the transport of soil particles and pollutants as suspended soil particulates in water, and the sediment is deposited to near-shore areas or near the mouths of tributaries (GLA, 1995). Once deposited, contaminated particles may become buried in deep sediment below a 10 cm depth. Particles buried at this depth are often considered to be lost to the aquatic system (Zarull et al., 1999).

Two processes exist that have the capacity to reintroduce contaminated sediment back into the water column. Bioturbation is the first process that results from the activity of benthic invertebrates in which sediment can be recycled from as deep as 40 cm from the active surface layer (Zarull et al., 1999). The second process is the resuspension of sediment as a result of major storm events, internal waves, currents, and vessel traffic (Zarull et al., 1999). Such external forces are causes for high flow events, which have
been observed to cause significant mass loadings of contaminants from a river into a lake (Cardenas and Lick, 1996 in Zarull et al., 1999). As expressed in section 4.2.3, the sediment originating in near shore areas can eventually accumulate in the deep water basins. Furthermore, the sediment plume developed at the mouth of the Niagara River leads directly to areas with high estimated sediment contamination levels.

4.2 Deterministic Methods

Having determined the SQI to be a satisfactory measure for sediment contamination, deterministic interpolation models including IDW, RBF, GPI, and LPI were assessed using the data from the sampling locations. The RMSPE values documented in Table 4.2 assess optimal functionality for each of the four methods. The optimal prediction surface created using each method is identified by the lowest RMSPE value generated using each specific function.

### Table 4.2: Results for Deterministic Methods of Spatial Interpolation

<table>
<thead>
<tr>
<th>Deterministic Method</th>
<th>Power</th>
<th>Kernel Function</th>
<th>Mean</th>
<th>Root-Mean-Squared-Error</th>
</tr>
</thead>
<tbody>
<tr>
<td>Inverse Distance Weighting</td>
<td>2nd</td>
<td>N/A</td>
<td>-1.63</td>
<td>13.22</td>
</tr>
<tr>
<td>Radial Basis Function</td>
<td>N/A</td>
<td>Multiquadric</td>
<td>-0.5637</td>
<td>12.22</td>
</tr>
<tr>
<td>Global Polynomial Interpolation</td>
<td>1st</td>
<td>N/A</td>
<td>0.02772</td>
<td>13.81</td>
</tr>
<tr>
<td>Local Polynomial Interpolation</td>
<td>2nd</td>
<td>N/A</td>
<td>-0.8688</td>
<td>12.99</td>
</tr>
</tbody>
</table>
Figures 4.2 to 4.5 represent the four deterministic interpolation maps including IDW, RBF, GPI, and LPI.

4.2.1 Inverse Distance Weighting

Utilizing IDW interpolation, cross-validation was used in order to attain the optimal power parameter and thus, to determine the lowest RMSPE for the prediction surface. As a result, the lowest RMSPE value was recorded applying the second power function and rendering a RMSPE of 13.22.

Given that IDW interpolation contours depend on the clustering of the sampling sites, bulls-eye patterns were created around locations featuring values differing from their surrounding points (Figure 4.2). These patterns prevent the possibility of making accurate predictions in between measured sites of sediment contamination because IDW does not account for the spatial autocorrelation of the surrounding points.

4.2.2 Radial Basis Functions

Radial Basis Functions use cross-validation for identification of the optimal kernel function in order to create a contour map. Given the SQI scores, the most optimal kernel function was identified as the ‘Multiquadric,’ on the basis of an RMSPE value of 12.22.
Figure 4.2 - Inverse Distance Weighting, SQI Scores

Legend

<table>
<thead>
<tr>
<th>Sediment Quality Index Categories</th>
<th>Prediction Intervals</th>
</tr>
</thead>
<tbody>
<tr>
<td>Marginal: SQI Value of 45 - 59</td>
<td>47 - &lt;55</td>
</tr>
<tr>
<td>Fair: SQI Value of 60 - 79</td>
<td>55 - &lt;60</td>
</tr>
<tr>
<td>Good: SQI Value of 80 - 94</td>
<td>60 - &lt;65</td>
</tr>
<tr>
<td>Excellent: SQI Value of 95 - 100</td>
<td>65 - &lt;70</td>
</tr>
<tr>
<td></td>
<td>70 - &lt;75</td>
</tr>
<tr>
<td></td>
<td>75 - &lt;80</td>
</tr>
<tr>
<td></td>
<td>80 - &lt;85</td>
</tr>
<tr>
<td></td>
<td>85 - &lt;90</td>
</tr>
<tr>
<td></td>
<td>90 - &lt;95</td>
</tr>
<tr>
<td></td>
<td>95 - 100</td>
</tr>
<tr>
<td></td>
<td>&lt;95 - &lt;100</td>
</tr>
<tr>
<td></td>
<td>No Data</td>
</tr>
</tbody>
</table>
Figure 4.3 - Radial Basis Functions, SQI Scores
Figure 4.4 - Global Polynomial Interpolation, SQI Scores
Figure 4.5 - Local Polynomial Interpolation, SQI Scores
The RBF prediction surface is shown in Figure 4.3. It is obvious that RBF are exact interpolation methods in which the SQI prediction surface was created by forcing contour lines through many sample values that were measured throughout Lake Ontario. Being an exact interpolator, the goal of RBF is to fit a line through a group of sampled values, while minimizing the overall curvature of the surface. Given the smooth nature of this interpolation model and the spatial differentiation of contamination concentrations over short horizontal distances, RBF was not identified as the most optimal prediction method.

4.2.3 Global Polynomial Interpolation (GPI)

Global Polynomial Interpolation is an inexact interpolation method that uses mathematical functions of spatial coordinates of sample sites in order to create prediction surfaces. In this analysis, a linear surface or first-order polynomial produced a predicted surface with the lowest RMSPE value calculated as 13.81. Since GPI attempts to account for global trends, the interpolated surface provides minimal variation in the concentration of sediment contamination.

Although the first-order polynomial produced the surface featuring the lowest RMSPE, performing GPI using the SQI dataset is not optimal for prediction of sediment contamination in Lake Ontario. This statement is supported by the lack of predicted SQI score variation featured in Figure 4.4. Minimal variation in the prediction surface is a result of GPIs inability to account for significant differences in contaminant concentrations over short horizontal distances. Specifically, this lack of variation exists
within the main sedimentary basin including the Niagara, Mississauga, and Rochester sub-basins. Within these sub-basins the range of SQI scores is predicted between 75 – 90, however, a large majority of measured values in these areas feature SQI scores below 75. Thus, GPI is not the optimal model to predict sediment contamination throughout Lake Ontario because it creates slowly varying surfaces in order to account for some global trend (eg. sedimentation with an eastern heading).

4.2.4 Local Polynomial Interpolation (LPI)

Given the domain of this analysis, short-range variations in SQI scores exist, making LPI an appropriate method for generating prediction surfaces. Rather than fitting a single polynomial to an entire area, LPI fits various polynomial functions to defined neighbourhoods over a surface area. In this analysis, a quadratic surface or second-order polynomial produced a predicted surface (Figure 4.5) with the lowest RMSPE value calculated as 12.99.

The resultant surface generated by the LPI model is more reasonable than that produced using GPI. The main reason is because it displays appropriate variation in SQI scores in the main sedimentary basin of Lake Ontario. Given this interpolation surface, SQI values ranging from 60 - 85 were featured in Lake Ontario’s main sedimentary basin, in which the variations account for significant differences in contaminant concentrations over short horizontal distances given the existing measured values that the predictions were based on. However, the northern extent of Lake Ontario is featured with
SQI values ranging from 95 - 100. Given that the Toronto Harbourfront is a Lake Ontario AOC, high SQI values along the northern shoreline appear to be too high. This is probably a result of two factors; the first due to the limitations of LPI as a prediction model; and the second related to the low number of measured sampling locations in the northern shoreline areas. For example, a sediment contamination measurement location at the Toronto Harbourfront AOC would have influenced the prediction along the northern shoreline of Lake Ontario. AOCs generally feature lower SQI scores, and thus, lower scores would be predicted if the measurements were taken at such locations. Although LPI produced a reasonable prediction surface for Lake Ontario, it was not utilized due to the limitations in determining the accuracy of the predictions through cross validation results measuring the bias and variability in its production.

4.3 Geostatistical Methods

Geostatistical interpolation models including ordinary, simple, universal, indicator, co-kriging and probability kriging were outlined and assessed for predicting sediment contamination levels in Lake Ontario. Ordinary kriging was chosen as the best model for prediction. Table 4.3 represents the semivariogram models and search neighbourhoods applied for the prediction of each contaminants distribution.

The process of semivariogram modelling is necessary because the semivariogram only provides information describing the spatial autocorrelation of datasets. It is
Table 4.3: Lake Ontario Sediment Contaminants: Semivariogram Models and Neighbourhood Search Size

<table>
<thead>
<tr>
<th>Contaminant</th>
<th>Semivariogram Model</th>
<th>Neighbourhood Search Size (# of neighbours)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sediment Quality Index (SQI)</td>
<td>Exponential</td>
<td>5</td>
</tr>
<tr>
<td>Arsenic</td>
<td>Spherical</td>
<td>5</td>
</tr>
<tr>
<td>Cadmium</td>
<td>Exponential</td>
<td>5</td>
</tr>
<tr>
<td>Chromium</td>
<td>Spherical</td>
<td>5</td>
</tr>
<tr>
<td>Copper</td>
<td>Exponential</td>
<td>5</td>
</tr>
<tr>
<td>Lead</td>
<td>Gaussian</td>
<td>5</td>
</tr>
<tr>
<td>Nickel</td>
<td>Gaussian</td>
<td>5</td>
</tr>
<tr>
<td>Zinc</td>
<td>Exponential</td>
<td>5</td>
</tr>
<tr>
<td>Mercury</td>
<td>Gaussian</td>
<td>5</td>
</tr>
<tr>
<td>Alpha-HCH or BHC</td>
<td>Gaussian</td>
<td>5</td>
</tr>
<tr>
<td>Hexachlorobenzene (HCB)</td>
<td>Exponential</td>
<td>5</td>
</tr>
<tr>
<td>Beta-HCH (Lindane)</td>
<td>Exponential</td>
<td>5</td>
</tr>
<tr>
<td>Heptachlor Epoxide</td>
<td>Exponential</td>
<td>5</td>
</tr>
<tr>
<td>Alpha-Chlordane</td>
<td>Exponential</td>
<td>5</td>
</tr>
<tr>
<td>Dieldrin</td>
<td>Exponential</td>
<td>5</td>
</tr>
<tr>
<td>pp’– DDE</td>
<td>Gaussian</td>
<td>5</td>
</tr>
<tr>
<td>Endrin</td>
<td>Exponential</td>
<td>5</td>
</tr>
<tr>
<td>pp’– DDD</td>
<td>Exponential</td>
<td>5</td>
</tr>
<tr>
<td>op’- + pp’ DDT</td>
<td>Exponential</td>
<td>5</td>
</tr>
<tr>
<td>Mirex</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Polychlorinated biphenyls (PCBs)</td>
<td>Exponential</td>
<td>5</td>
</tr>
<tr>
<td>Dioxins and Furans</td>
<td>Spherical</td>
<td>5</td>
</tr>
<tr>
<td>Phenanthrene</td>
<td>Spherical</td>
<td>5</td>
</tr>
<tr>
<td>Anthracene</td>
<td>Spherical</td>
<td>5</td>
</tr>
<tr>
<td>Fluoranthene</td>
<td>Exponential</td>
<td>5</td>
</tr>
<tr>
<td>Pyrene</td>
<td>Spherical</td>
<td>5</td>
</tr>
<tr>
<td>Benzo[a]anthracene</td>
<td>Spherical</td>
<td>5</td>
</tr>
<tr>
<td>Chrysene</td>
<td>Spherical</td>
<td>5</td>
</tr>
<tr>
<td>Benzo[b/k]fluoranthene</td>
<td>Spherical</td>
<td>5</td>
</tr>
<tr>
<td>Benzo[a]pyrene</td>
<td>Spherical</td>
<td>5</td>
</tr>
<tr>
<td>Indeno[1,2,3.cd]pyrene</td>
<td>Exponential</td>
<td>5</td>
</tr>
<tr>
<td>Dibenzo[a,h]anthracene</td>
<td>Exponential</td>
<td>5</td>
</tr>
<tr>
<td>Benzo[g,h I]perylene</td>
<td>Exponential</td>
<td>5</td>
</tr>
<tr>
<td>Total PAHs</td>
<td>Exponential</td>
<td>5</td>
</tr>
</tbody>
</table>
necessary to fit a semivariogram model in order to provide information for all possible directions and distances, and to ensure that kriging predictions have positive kriging variances (Johnston et al., 2001).

The three models utilized in this analysis are identified as the spherical, exponential, and gaussian models. The prediction of unknown values is most significantly influenced when the shape of the curve near the origin differs. For instance, the Gaussian model is used if the semivariogram depicts autocorrelation at short distances between data points, given its parabolic shape near the origin. The Gaussian model incorporates this shape in order to make use of the closest samples in the prediction (Isaaks and Srivastava, 1989). However, the parabolic shape often causes a screening effect in which one data sample falls in between another sample and the point being estimated. In such an instance, negative weights can be produced (Isaaks and Srivastava, 1989). If a particularly high sample value is associated with a negative weight, there is a high possibility of a negative estimate. This explains why the Spherical and Exponential models were applied more readily in this analysis than the Gaussian model. These models are more linear in nature and as a result, they produce fewer negative weights. Earth science analyses such as pollutant concentrations should never produce negative estimation values because the lowest amount measurable is a value of 0 (Isaaks and Srivastava, 1989).
4.3.1 Cross-Validation Results for Ordinary Kriging

As outlined in section 3.3.7, the cross-validation procedure provides measures of accuracy for the predictions made using the ordinary kriging method. The measures produced include MPE, SMPE, RMSPE, ASE and SRMSPE. Values calculated for these measures are documented in Table 4.4, where ordinary kriging is assessed as a predictor of sediment contamination for 32 different contaminants.

Given each specific contaminant, results are most variable when considering RMSPE values and when comparing RMSPE values with ASEs. Before the interpolated surfaces can be assessed, it is necessary to document possible explanations for inaccuracies that occurred in the kriging process.

4.3.2 Limitations of the data

The main reason kriging is a valuable asset in the domain of environmental science is because the process of collecting and analyzing samples of many types is both a tedious and an expensive task. Due to budget and time restraints in research programs, data collected for entities such as Lake Ontario are often limited. In this analysis, kriging offers the possibility for predicting sediment contamination throughout Lake Ontario on the basis of 70 sampling sites. A large percentage of the results (rendered by kriging the 32 contaminants in this analysis) were positive. Negative results featuring high RMSPE scores may have been calculated due to an inappropriate number of measured sampling
locations. In the previously mentioned kriging analysis in Germany, it was concluded that 47 sampling sites were not sufficient to reduce the spatial variance distributions to acceptable values in order to make accurate predictions (Buttner et al., 1998). As stated in section 3.3.6, 47 sampling sites were measured over a relatively small area, compared to 70 sites throughout Lake Ontario, which has a large surface area. However, knowledge of the sedimentation and current circulation patterns in Lake Ontario was useful in estimating the sediment dispersal within the system.

As displayed in Figure 1.3, the sampling locations are not evenly dispersed throughout the lake. The region for highest concern of inaccurate predictions is obviously the entire northern shoreline of Lake Ontario. Unlike the southern shoreline, no data were generated from sampling locations in close proximity to the northern shoreline.

With the exception of the Niagara River and Hamilton Harbour, sediment samples were not analyzed at other Areas of Concern (AOC) such as the Toronto Harbourfront or major contaminating rivers flowing into Lake Ontario along the northern shoreline. Instead, high SQI scores were predicted in these regions based on similar scores from measured sites located within their neighbourhood search areas. These results are inaccurate because AOCs possess high sediment contamination levels, and thus, low SQI values. Sediment quality predictions made in the south-western extent of the lake are generally more accurate because two samples were taken in Hamilton Harbour and one at the mouth of the Niagara River. Due to the abundance of inflow from the Niagara River,
a sampling location at its mouth represents sediment contamination in the region more thoroughly.

Finally, using ordinary kriging produces portions of Lake Ontario’s area as featuring no data. Initially, a rectangular prediction surface approximately covered the entire area of Lake Ontario. Areas located along the south-west and north-east edges of Lake Ontario feature no data due to limitations in the extent of sampling site locations.

4.3.3 Sediment Quality Index (SQI)

In Figure 4.6, SQI categories are overlaid on top of the SQI contamination score prediction surface. SQI categories range from points that display Marginal (red) to Excellent (green) sediment quality. Furthermore, brown to yellow colour intervals represent predictions of low to high SQI scores respectively. When analyzing cross-validation results, it is important to concentrate on the negative aspects including the worst errors, areas consistently displaying bias, or areas that have been misclassified (Isaaks and Srivastava, 1989). It is also important to note that successful cross-validation results do not necessarily guarantee an accurate prediction surface. However, poor results are good indicators of inaccuracies in predicted surfaces (Isaaks and Srivastava, 1989). Figure 4.6 displays the lowest SQI scores in the region of Hamilton Harbour and central regions of the Niagara, Mississauga, and Rochester basins. Highest SQI values are estimated along the northern shoreline of Lake Ontario. However, they may be inaccurate
given the lack of measured sampling locations in this area. Furthermore, in Lake Ontario, sediment from near shore areas eventually accumulates in the deep lake basins or moves through the St. Lawrence River to the Atlantic Ocean (Allan, 1984; Sorokin, 1966; Karickhoff and Morris, 1995 in Zarull et al., 1999). Therefore, it was necessary to sample the sediment in the deep basins where this contaminated material is known to migrate.

The cross-validation results derived from the SQI score interpolation results document the RMSPE, ASE, SMPE and SRMSPE as 13.14, 13.68, -0.03784, and 0.9635 respectively. Thus, the predictions are unbiased and reasonably close to the measured locations. The variability assessed by the deviations of the ASE from the RMSPE and the SRMSPE from the value 1, is slightly overestimated. Through examination of the cross-validation results and the predicted surface, ordinary kriging proved to be an appropriate method for predicting sediment contamination utilizing the SQI scores.

On the basis of the RMSPE produced for deterministic methods and the ordinary kriging model, results were similar ranging from 12.22 to 13.81. Kriging was determined the most appropriate interpolation method tested because unlike deterministic models, it quantifies spatial autocorrelation in order to weight the spatial arrangement of measured points (Johnston et al., 2001). Additional reasons include the following:

(1) kriging accounts for both the clustering of nearby samples and their distance from the estimation points;
(2) values and plots assessing the accuracy of the predictions are calculated in the cross-validation process;
(3) multiple kriging models exist, and can be used depending on the desired output surfaces and the nature of the data set.
In Figure 4.6, ordinary kriging results (using SQI scores) show higher contamination levels in the deep lake basins. Kriging accounted for this sedimentation process where low SQI scores are situated within the deepest extent of each basin, and in close proximity to Lake Ontario AOCs. Interpretation of the five contour surfaces (Figure 4.7) relative to the distribution of depth and inflow throughout Lake Ontario supports the kriging model as an improvement over the alternative deterministic methods. Given the general water circulation patterns and inflows from rivers and tributaries, sediment has the capacity to be re-suspended, and deposited in deeper extents of the Lake Ontario basin. Deterministic methods did not account for the depositional processes in this aquatic system as accurately.

Following the assessment of existing kriging models in ArcGIS 8.1, ‘ordinary kriging’ was selected as the most appropriate kriging method in order to predict sediment contamination levels at unknown locations throughout Lake Ontario. The reasons for this decision include:

1. There is no reason to reject the assumption of an ‘unknown’ constant mean;
2. The data plotted in the semivariogram appeared to have a spatial trend. Thus, ordinary kriging could be applied to this analysis because prediction weights could be derived from a fitted variogram model;
3. Simple kriging was not reasonable because the assumption of a ‘known’ constant mean \( u \), was not possible given the dataset. Furthermore, the assumption of second-order stationarity was not possible given the physical nature of the data;
4. Universal Kriging is similar to ordinary kriging, however, a deterministic trend replaces the unknown constant mean. This model could not be applied because the assumption of an unknown mean \( u \) rendered more optimal results in the cross-validation process than assigning a deterministic trend;
Deterministic Interpolation Vs. Kriging

Image(s)/Figure not available.
(5) Indicator kriging produces interpolation maps displaying the probability that a specific variable is in a defined class indicated by a binary number. The interest of this analysis is predicting contaminant levels throughout Lake Ontario. Therefore, indicator kriging would not render the desired results;

(6) Cokriging could not produce improved results, because its purpose is to aid spatial interpolation when utilizing incomplete datasets. This analysis was performed using a complete dataset for 32 contaminants residing in Lake Ontario sediment;

(7) Probability kriging is a compilation of both indicator and co-kriging. Rejection of these methods is grounds for utilizing ordinary kriging in this analysis.

Prediction maps were created using the ordinary kriging method for the 32 contaminants that were analyzed in the core samples. The intervals defined for the point data measurements represent the federal concentration guidelines specific to each contaminant. These TELs and PELs for 32 variables are featured in Table 1.2. Prediction surfaces are represented in following sections.

Measured contaminant values are overlaid onto these prediction maps in order to assess the estimations surrounding the contamination values. Each contaminant was classed based on its contamination level relative to the federal TEL and PEL guidelines. Statistics were generated in the cross-validation process including MPE, SMPE, RMSPE, ASE, and SRMSPE that validated ordinary kriging as an appropriate interpolation method for sediment contamination in Lake Ontario.
Table 4.4: Cross-Validation Results for Ordinary Kriging

<table>
<thead>
<tr>
<th>Contaminant</th>
<th>Mean</th>
<th>Root-Mean-Square Error</th>
<th>Average Standard Error</th>
<th>Mean Standardized</th>
<th>RMS Standardized</th>
</tr>
</thead>
<tbody>
<tr>
<td>Arsenic</td>
<td>-0.03715</td>
<td>12.31</td>
<td>11.25</td>
<td>-0.002141</td>
<td>1.09</td>
</tr>
<tr>
<td>Cadmium</td>
<td>0.151</td>
<td>1.543</td>
<td>1.514</td>
<td>0.09463</td>
<td>1.023</td>
</tr>
<tr>
<td>Chromium</td>
<td>0.8721</td>
<td>20.07</td>
<td>19.95</td>
<td>0.04434</td>
<td>1.0</td>
</tr>
<tr>
<td>Copper</td>
<td>1.938</td>
<td>27.09</td>
<td>28.38</td>
<td>0.05492</td>
<td>0.9448</td>
</tr>
<tr>
<td>Lead</td>
<td>3.091</td>
<td>42.72</td>
<td>42.26</td>
<td>0.07361</td>
<td>1.011</td>
</tr>
<tr>
<td>Nickel</td>
<td>1.098</td>
<td>24.42</td>
<td>22.65</td>
<td>0.03799</td>
<td>1.077</td>
</tr>
<tr>
<td>Zinc</td>
<td>3.998</td>
<td>172.5</td>
<td>148</td>
<td>0.01515</td>
<td>1.159</td>
</tr>
<tr>
<td>Mercury</td>
<td>0.01983</td>
<td>0.3452</td>
<td>0.3584</td>
<td>0.05032</td>
<td>0.9693</td>
</tr>
<tr>
<td>Alpha-HCH or BHC</td>
<td>0.002521</td>
<td>0.1612</td>
<td>0.1607</td>
<td>0.01426</td>
<td>1.009</td>
</tr>
<tr>
<td>Hexachlorobenzene (HCB)</td>
<td>0.4908</td>
<td>14.46</td>
<td>14.65</td>
<td>0.02495</td>
<td>0.9689</td>
</tr>
<tr>
<td>Beta-HCH (Lindane)</td>
<td>0.008849</td>
<td>0.2457</td>
<td>0.2264</td>
<td>0.02652</td>
<td>1.076</td>
</tr>
<tr>
<td>Heptachlor Epoxide</td>
<td>0.00339</td>
<td>0.1296</td>
<td>0.1219</td>
<td>0.02388</td>
<td>1.048</td>
</tr>
<tr>
<td>Alpha-Chlordane</td>
<td>0.03205</td>
<td>0.5431</td>
<td>0.5349</td>
<td>0.04861</td>
<td>1.038</td>
</tr>
<tr>
<td>Dieldrin</td>
<td>0.03634</td>
<td>0.7788</td>
<td>0.776</td>
<td>0.04062</td>
<td>0.9901</td>
</tr>
<tr>
<td>pp’ – DDE</td>
<td>1.379</td>
<td>14.94</td>
<td>14.19</td>
<td>0.09391</td>
<td>1.058</td>
</tr>
<tr>
<td>Endrin</td>
<td>0.01257</td>
<td>0.1746</td>
<td>0.1742</td>
<td>0.06992</td>
<td>1.007</td>
</tr>
<tr>
<td>pp’ – DDD</td>
<td>1.085</td>
<td>12.66</td>
<td>12.97</td>
<td>0.08294</td>
<td>0.9771</td>
</tr>
<tr>
<td>op’ - + pp’ DDT</td>
<td>0.3099</td>
<td>4.488</td>
<td>4.314</td>
<td>0.0706</td>
<td>1.046</td>
</tr>
<tr>
<td>Mirex</td>
<td>N/A</td>
<td>N/A</td>
<td>N/A</td>
<td>N/A</td>
<td>N/A</td>
</tr>
<tr>
<td>Polychlorinated biphenyls (PCBs)</td>
<td>3.813</td>
<td>65.52</td>
<td>68.1</td>
<td>0.04724</td>
<td>0.9617</td>
</tr>
<tr>
<td>Dioxins and Furans</td>
<td>0.2669</td>
<td>76.71</td>
<td>83.79</td>
<td>0.008826</td>
<td>0.9465</td>
</tr>
<tr>
<td>Phenanthrene</td>
<td>0.965</td>
<td>143.3</td>
<td>108.6</td>
<td>0.004604</td>
<td>1.308</td>
</tr>
<tr>
<td>Anthracene</td>
<td>0.3568</td>
<td>31.26</td>
<td>21.57</td>
<td>0.00676</td>
<td>1.473</td>
</tr>
<tr>
<td>Fluoranthene</td>
<td>11.95</td>
<td>276.8</td>
<td>168.6</td>
<td>0.03379</td>
<td>1.963</td>
</tr>
<tr>
<td>Pyrene</td>
<td>2.052</td>
<td>233.2</td>
<td>160.6</td>
<td>0.003045</td>
<td>1.462</td>
</tr>
<tr>
<td>Benzo[a]anthracene</td>
<td>1.828</td>
<td>149.7</td>
<td>100.7</td>
<td>0.003765</td>
<td>1.519</td>
</tr>
<tr>
<td>Chrysene</td>
<td>1.189</td>
<td>193.5</td>
<td>146.5</td>
<td>0.0002555</td>
<td>1.32</td>
</tr>
<tr>
<td>Benzo[b/k]fluoranthene</td>
<td>4.631</td>
<td>365.9</td>
<td>278.3</td>
<td>-0.0008384</td>
<td>1.34</td>
</tr>
<tr>
<td>Benzo[a]pyrene</td>
<td>1.267</td>
<td>146.9</td>
<td>119.7</td>
<td>0.002388</td>
<td>1.226</td>
</tr>
<tr>
<td>Indeno[1,2,3.cd]pyrene</td>
<td>9.004</td>
<td>172.5</td>
<td>181.8</td>
<td>0.04856</td>
<td>0.9544</td>
</tr>
<tr>
<td>Dibeno[a,h]anthracene</td>
<td>1.397</td>
<td>31.24</td>
<td>32.34</td>
<td>0.04203</td>
<td>0.9722</td>
</tr>
<tr>
<td>Benzo[g,h I]pyrene</td>
<td>7.947</td>
<td>133.9</td>
<td>136.4</td>
<td>0.05662</td>
<td>0.9874</td>
</tr>
<tr>
<td>Total PAHs</td>
<td>54.41</td>
<td>1865</td>
<td>1354</td>
<td>0.01011</td>
<td>1.49</td>
</tr>
</tbody>
</table>
4.3.4 Metals

Displayed as Figures 4.8 through 4.15 respectively, ordinary kriging was used to predict metal concentrations for Mercury, Cadmium, Arsenic, Chromium, Lead, Nickel, Copper, and Zinc in Lake Ontario. Due to a high RMSPE value, a large deviation between the RMSPE and ASE, and a standardized RMSPE that deviated significantly from the value 1, an accurate prediction surface could not be produced for Zinc concentrations. A biased prediction for Zinc concentrations is displayed with a MPE value of 3.998.

Figure 4.8 estimates the locations for the highest Mercury concentrations in the deep central regions of both the Mississauga and Rochester sub-basins. Unlike the prediction maps produced for the SQI values, federal guideline intervals representing measured concentrations either below or exceeding the TEL or PEL are superimposed over the contour surfaces in Figures 4.8 through 4.15. Furthermore, the specific choropleth schemes ranging from five to nine intervals are applied to all metals. The yellow pigment represents the lowest metal concentrations in the sediment and the brown represents the highest. The predicted surface for Mercury produced the most reliable cross-validation results, followed closely by Cadmium (Figure 4.9). Both results were relatively unbiased and rendered low RMSPE values. However, variability in the predictions was slightly overestimated for Mercury and underestimated for Cadmium. Measured concentrations for the presence of Cadmium exceeding the PEL were fewer
Figure 4.8 - Ordinary Kriging, Mercury Concentrations
Ordinary Kriging - Implementing Cadmium Concentrations

Legend

Cadmium Federal Concentration Guidelines | Prediction Intervals (µg/g)
--- | ---
Below TEL (0.0 - 0.5 µg/g) | 0.0 - 0.5
Exceeds TEL (0.6 - 3.5 µg/g) | 0.5 - 1.0
Exceeds PEL (3.5 - 5.8 µg/g) | 1.0 - 1.5

Figure 4.9 - Ordinary Kriging, Implementing Cadmium Concentrations
Ordinary Kriging - Implementing Chromium Concentrations

Legend

Chromium Federal Contamination Guidelines
- Below TEL (0.0 - <37.3 mg/kg)
- Exceeds TEL (37.3 - <60.0 mg/kg)
- Exceeds PEL (90.0 - 110.0 mg/kg)

Prediction Intervals (mg/kg)
- 0.0 - <10.0
- 10.0 - <20.0
- 20.0 - <30.0
- 30.0 - <40.0
- 40.0 - <50.0

Figure 411- Ordinary Kriging, Implementing Chromium Concentrations
Figure 4.12 - Ordinary Kriging, Lead Concentrations
Ordinary Kriging - Implementing Copper Concentrations

Legend

Copper Federal Contamination Guidelines
- Below TEL (0.0 - <35.7 ug/g)
- Exceeds TEL (35.7 - 119.0 ug/g)

Prediction Intervals (ug/g)
- 0.0 - <20.0
- 20.0 - <40.0
- 40.0 - <60.0
- 60.0 - <80.0
- 80.0 - <100.0
- 100.0 - 109.0
- No Data

Figure 4.14 - Ordinary Kriging, Implementing Copper Concentrations
Image/Figure not available.
than the case for Mercury, however, high contamination concentrations for both elements were in similar general locations.

A general trend for the spatial distribution of metals throughout the interpolated surfaces in this section is that highest distributions exist in the deep central lake basins including the Rochester, Mississauga, and in most instances, the Niagara basin. Variations in the distribution of these particles exist due to their physical properties such as particle size and atomic weights. The distribution of these elements can be explained due to the specific sedimentation and circulation patterns in Lake Ontario.

Figure 4.10 representing Arsenic, produced a similar RMSPE to the prediction surface created implementing SQI scores, and its deviation from the ASE was larger, resulting in an underestimation of variability in the predictions. The highest concentrations of Arsenic were predicted in the south-eastern extent of Lake Ontario (the deepest region in the Rochester Basin). High concentrations in this area are also predicted in the shallow inshore region of the basin. This suggests that Arsenic loadings stem from rivers in this area including the Genesee and the Oswego. Figures 4.11 through 4.14 represent Chromium, Lead, Nickel, and Copper respectively. Prediction surfaces representing Chromium and Lead near perfectly estimated the variability and featured standardized RMSPE values of 1.0 and 1.011, however, Lead features an RMSPE value of 42.72, more than doubling that of Chromium. With RMSPE values of greater than 20, predictions are straying quite far from the measured locations. Therefore, when analyzing these prediction surfaces, these errors must be taken into account. On the basis of these
predictions, the highest concentrations of Chromium are predicted in the general region of Hamilton Harbour. It seems that Chromium loadings stem from industrial processes performed in the Hamilton Harbour region and the Niagara River. Additionally, the highest concentrations remain in the shallow inshore regions surrounding the Niagara basin. As a result, it is possible to hypothesize that the physical properties of Chromium inhibit significant lateral movement throughout Lake Ontario. The prediction map representing Lead is obviously inaccurate given that the highest predicted concentration interval indicates Lead values ranging from 80 to 100 ug/g, while measured concentrations of Lead were documented at amounts well over 100 ug/g in the central regions of Lake Ontario, and approximately 200 ug/g near Hamilton Harbour. The highest metal concentrations including Lead stem largely from steel mills and associated practices along the shoreline of Hamilton Harbour (Ecowise, 2002). Hamilton Harbour is a reservoir for industrial and municipal wastes, including lead laden effluents, and acts as a port that receives 400 to 1000 vessels per year (Ecowise, 2002). As a result, these contaminants infiltrate the sediment and are deposited in the deep central basin over time. However, it is vital to reiterate that prediction results may not be reliable due to inaccurate cross-validation values and simple comparison of measured and predicted values.

The prediction maps created for Nickel and Copper display high concentrations of each element in the central regions of the Niagara and Mississauga sub-basins. Similar to the cross-validation results produced for Lead, these prediction surfaces are subject to inaccuracies due to high RMSPE values and standardized RMSPE values deviating from
1. The interpolation surfaces suggest that major rivers in the Lake Ontario ecosystem including the Humber, Don, and Trent on the northern shoreline, and the Niagara, Genesee, and Oswego on the southern shoreline are the main sources for these contaminants. High contamination levels within the deep regions of the Niagara, Mississauga, and Rochester basins support this hypothesis.

4.3.5 Critical Pollutants

Figures 4.16 to 4.21 represent interpolated surfaces for PCBs, Dioxins and Furans, Dieldrin, DDT, DDD, and DDE respectively. Figure 4.16 represents the prediction map for Polychlorinated biphenyls. PCBs have been banned from production for the past 25 years, but their concentration levels still continue to exceed human health standards. This contaminant was measured at high concentrations in the deep regions of the Mississauga and Rochester basins. Comparing the measured values to the filled contours throughout the basin, the predictions seem reasonable, but, high RMSPE results and large deviations from the ASE are the reason to suspect inaccuracies in the kriging analysis. A possible explanation for these cross-validation results is a biased prediction supported by a MPE value of 3.813. It is due to the severe impacts that PCBs continue to make on aquatic ecosystems that the prediction map was included in this analysis despite its errors.
Figure 4.16 - Ordinary Kriging, PCB Concentrations
Ordinary Kriging - Dioxins and Furans Concentrations

Legend

Dioxins and Furans
Federal Concentration Guidelines
- Below TEL (0.0 - <0.83 ng/g)
- Exceeds TEL (0.83 - <21.5 ng/g)
- Exceeds PEL (21.5 - 271.2 ng/g)

Prediction Intervals:
- 3.0 - <30.0
- 30.0 - <60.0
- 60.0 - <120.0
- 120.0 - <150.0
- 150.0 - <180.0
- 180.0 - <210.0
- 210.0 - <240.0
- 240.0 - <271.2
- No Data

Figure 4.17 - Ordinary Kriging, Dioxins and Furans Concentrations
Figure 4.18 - Ordinary Kriging, Dieldrin Concentrations
Figure 4.19 - Ordinary Kriging, DDT Concentrations
Figure 4.21 - Ordinary Kriging, DDE Concentrations
The majority of PCB loadings into Lake Ontario originate from the upstream Great Lakes basins and the Niagara River basin totalling 440 kg/yr (LOLMP, 1998). The Niagara River is the main source for this inflow. The highest concentrations of PCBs in sediment are located in the deep central basins of Lake Ontario originating from the mouth of the Niagara River. Point and non-point sources, combined with atmospheric deposition, contribute approximately 165 kg/yr of PCBs to the lake (LOLMP, 1998). The majority of this contaminant enters the lake through tributaries and rivers.

Figure 4.17 and 4.18 represent prediction maps for Dioxins and Furans and Dieldrin. Cross-validation results reported for Dioxins and Furans depict slightly biased predictions in which the variability is overestimated. The RMSPE is extremely high indicating predictions that are not close to the measured values. Dieldrin was predicted successfully, featuring a low RMSPE value, and unbiased predictions in which the variability was very slightly overestimated.

The measurements for the highest concentrations of Dioxins and Furans were sampled in the deep region of the Niagara, Mississauga, and Rochester basins. These contaminants tend to exist at low levels in the environment. The largest source of these contaminants is atmospheric deposition and it accounts for approximately 5 grams per year (LOLMP, 1998). The Niagara River is an additional source, along with Oswego River on the south-east shore and the Genesee River near Rochester. Similar to Dioxins and Furans, the highest Dieldrin concentrations are predicted in the central regions of the main sub-basins. The majority of Dieldrin loadings stem from the Niagara River (and
other upstream sources) representing approximately 43 kg/yr that is deposited in the deep lake basins over time (LOLMP, 1998). Atmospheric deposition accounts for approximately 13 kg/yr.

The final critical pollutant predicted in this analysis is DDT (Figure 4.19). The cross-validation results for DDT showed a relatively unbiased prediction, in which predictions were close to the measured locations and variability was assessed reasonably well. Highest concentrations were predicted near the urban centre of Oshawa. This portion of the Lake Ontario shoreline does however feature many farming communities that used DDT as a pesticide in past decades. It is apparent that DDT in sediment has reached the deep lake basin due to re-suspension of sediment over time. DDT loadings originate from both the upstream Great Lakes (Niagara River) and from sources including the Humber and Don Rivers on the northern shoreline, and Eighteen Mile Creek, the Genesee, Oswego, and Black Rivers on the southern shoreline (LOLMP, 1998). Atmospheric loadings stemming from arable farmland regions near the Trent River also account for a small percentage of DDT loading. Figures 4.20 and 4.21 include DDD and DDE respectively (which combined form DDT). Cross-validation results for these contaminants do not suggest predictions as close to the measured location when compared to DDT, however, high concentration results reside in similar zones for each contaminant.
4.3.6 Polycyclic Aromatic Hydrocarbons (PAHs)

In total, the sediment contamination levels for eleven different PAHs were predicted throughout Lake Ontario. Prediction attempts for ten of the eleven contaminants were unsuccessful due to mean prediction errors deviating significantly from 0, high RMSPE values, unacceptable deviations of these values from the ASEs, and standardized RMSPE values deviating significantly from 1. Figure 4.22 represents Anthracene, the sole PAH in which the results rendered were not completely unacceptable. The results are comparable to those calculated for Copper, however a RMSPE of greater than 30 and a large overestimation in the variability of the prediction, makes for an inaccurate prediction surface. The highest concentrations of Anthracene were predicted and measured in the region of Hamilton Harbour. It is obvious that loadings of this PAH originate from industrial processes occurring along the shores of the harbour. Figures 4.23 to 4.32 represent the statistically insignificant prediction surfaces created for the remaining PAHs including Phenanthrene, Fluoranthene, Pyrene, Benzo[a]anthracene, Chrysene, Benzo[b/k]fluoranthene, Benzo[a]pyrene, Indeno[1,2,3-cd]pyrene, Dibenzo[a,h]anthracene, and Benzo[g,h,I]perylene respectively. Negative prediction results were rendered for the following reasons:

1. the MPEs were significantly biased (deviated from the value 0), thus, the predictions were not centred on the measurement values;
2. high RMSPE values were a result of the cross-validation process showing a large deviation of the predictions from their measured values;
Ordinary Kriging - Implementing Anthracene Concentrations

Legend

Anthracene Federal Concentration Guidelines  Prediction Intervals (ng/g)
- Below TEL (0.0 - <44.9 ng/g)
- Exceeds TEL (46.9 - <245.0 ng/g)
- Exceeds PEL (245.0 - 399.0 ng/g)

Figure 4.22 - Ordinary Kriging, Implementing Anthracene Concentrations
Image/Figure not available.
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Image/Figure not available.
Image/Figure not available.
Image/Figure not available.
Image/Figure not available.
Image/Figure not available.
Image/Figure not available.
Image/Figure not available.
the variability in the predictions was not assessed correctly
the distances between sampling locations were inappropriate given the range
depicted by the semivariogram models chosen in each interpolation. A higher
sampling density would have led to more accurate predictions because measured
points would remain within the range of the semivariogram, and thus, show
spatial autocorrelation.

4.3.7 The Remaining Contaminants

Figure 4.33 and 4.34 represent Alpha-HCH or BHC and Endrin. Both cross-
validation results reported low RMSPE scores and almost perfect variability. Similarly,
Figures 4.35 through 4.37 representing Lindane, Heptachlor Epoxide and Alpha-
Chlordane respectively, featured predictions close to measured locations, however, the
variability of prediction was underestimated slightly in all three cases. Figures 4.33
through 4.37 are fine examples of statistically accurate results using ordinary kriging.
Furthermore, with the exception of Figure 4.35 (Lindane), all contamination
measurements were recorded and predicted below the TEL. Figure 4.38, representing
Hexachlorobenzene produced similar results to the implementation of the SQI with
geostatistical interpolation.

Similar to the metals in section 4.3.4, reports documenting the origin of pollutant
loadings for the contaminants in this section are minimal. Due to low concentration levels
of these variables in Lake Ontario sediment, it is difficult to hypothesize their origins.
Figure 4.33 - Ordinary Kriging, Alpha-HCH/BHC Concentrations
Figure 4.36 - Ordinary Kriging, Heptachlor Epoxide Concentrations
Figure 4.37 - Ordinary Kriging, Alpha-Chlordane Concentrations
4.4 Sampling Strategies

As displayed in Figure 3.1, an exponential model is fit to SQI measurement values in a semivariogram. In this example, the range is a distance of approximately 3.2, beyond which autocorrelation does not exist. In order to produce the best results given the limitations of the dataset, data sampling locations over the surface of Lake Ontario must be within this range. A distance conversion could not be made from the semivariogram value, into its physical distance on the surface of the earth due to limitations of the ArcGIS software.

In order to optimize predictions, it will be necessary to take sediment contamination samples at all Lake Ontario AOCs and major tributaries providing inflow into the system. These entities may include the Don and Humber Rivers near the urban centre of Toronto, and Eighteen Mile Creek located near the Niagara River. Finally, it is important that sampling locations are situated within the range of the semivariogram, to ensure autocorrelation between sites of sediment contamination. However, the grid pattern featured in this analysis did not produce the optimal prediction results. Improved results would be produced if the sampling locations were more spatially stratified. Spatial stratification is the opposite of a random pattern, in that there is some bias imposed on how a particular sampling scheme is chosen. Improved results would be produced in this analysis if the sampling scheme was stratified in intervals no larger than the range produced in the semivariogram for the particular contaminant of interest.
Chapter 5: Summary and Conclusions

The measurement of sediment contamination throughout Lake Ontario is an immense task if one is to make thorough observations about the aquatic ecosystem as a whole. In performing the assessment to determine the SQI’s legitimacy as a satisfactory measure for identifying areas in Lake Ontario where sediment quality is frequently threatened or impaired, the spatial distribution of the critical pollutants was documented. In addition, the total number of contaminants at each measurement location exceeding the TEL and the PEL was determined together with the amount by which each threshold was exceeded. The SQI was found to be a good general measure for frequently threatened or impaired sediment after performing the calculation of the SQI at each of the 70 measurement locations and reviewing the results. Supporting this conclusion were sampling locations featuring high numbers of contaminants exceeding the PEL and TEL’s. In other frequently threatened or impaired sediment samples, the amount by which each threshold was exceeded was extremely high.

Once conclusions were made regarding the SQI as a satisfactory measure for frequently threatened or impaired sediment in Lake Ontario, the spatial distribution of SQI values was analyzed. Limitations in the spatial distribution of the sediment contaminant data prevented the prediction of an accurate SQI score distribution without implementation of deterministic and/or geostatistical interpolation methods. Estimations of this distribution were performed using deterministic methods including Inverse Distance Weighting Interpolation, Radial Basis Functions, and Global Polynomial
Interpolation, and Local Polynomial Interpolation. A geostatistical method referred to as ordinary kriging was also assessed relative to these deterministic methods, in order to identify the optimal interpolation technique for this analysis. Cross-validation was utilized to perform this assessment by producing a root-mean-square prediction error for each of the five prediction methods, in which the lowest values represented predicted locations situated reasonably close to the measured location. Although the RMSPE values were similar for the methods, kriging was identified as the best prediction model because it uses statistical models that incorporate autocorrelation among the group of measured points to create a prediction surface. Also, cross-validation results including MPE, ASE, SMPE, and SRMSPE are produced exclusively in the kriging process. These results include measures relating to the level of bias in a prediction and estimations of variability in the production of interpolated surfaces. While deterministic interpolation models created reasonable prediction surfaces (with the exception of GPI), the models relied on linear functions rather than semivariogram modeling that offers the ability to customize the analysis to the nature of the data.

An assessment of existing kriging methods within the ArcGIS software and identification of the optimal method to be used given the nature of this specific case study was the next step. Ordinary kriging was chosen as the optimal method for reasons regarding its functionality for the specific analysis. The ordinary kriging process produced 20 acceptable prediction surfaces for a possible 32 contaminants.
After assessing the cross-validation results for the spatial distribution of metals in this analysis, the most statistically optimal prediction surfaces were created for mercury and cadmium, and adequate results were rendered for the remaining metals. Zinc was excluded from the results for the reason of poor cross-validation results documented previously. The results calculated for the critical pollutants were variable, with the most accurate values representing Dieldrin. The most inaccurate results were calculated for PCBs.

The group of eleven PAHs produced the most statistically inaccurate results, featuring biased results in the form of mean prediction errors deviating from the value zero, high RMSPEs, large deviations of these values from their corresponding ASEs, and deviations of standardized RMSPEs from the value 1. Anthracene is one PAH for which an acceptable prediction surface was produced. In comparison, the group of variables referred to as the ‘remaining contaminants’ produced statistically significant cross-validation results. Alpha-HCH or BHC and Endrin produced the most accurate prediction surfaces, whereas, Hexachlorobenzene produced similar results to the implementation of the SQI scores utilizing the ordinary kriging method.

Before the kriging estimates could be discussed, it was necessary to identify the limitations of the dataset that created inaccurate results in the resulting surfaces. The main limitation of the data was the insufficient spatial distribution of measured sampling locations along the northern shoreline of Lake Ontario. A more sufficient spatial distribution of sampling locations along the shoreline would have created more accurate
prediction values in the northern region of Lake Ontario. For example, samples measured at Lake Ontario Areas of Concern such as the Toronto Harbourfront would have added to the accuracy of the predictions, and displayed higher sediment contamination levels in the northern shoreline areas. However, sediment dispersal in the Toronto region is dependant on factors such as the W/SW direction of the offshore current in this area.

In the future, more resources should be expended to develop a sampling scheme that will account for the proper range in which autocorrelation exists between sampling sites. As described in section 4.4, the development of a specific stratification scheme could be based on the calculated range in the semivariogram process. If sampling locations follow this defined stratification process, a more statistically accurate prediction surface can be created. It is vital to note that the semivariogram distribution will change for each contaminant. Thus, the ranges produced for each contaminant will not be identical. Before creating a sampling strategy, it is necessary to decide which contaminant distribution is most important to the organization in order to produce the best possible interpolation results. In this analysis, it is equally important to take samples at Lake Ontario AOCs and other rivers and tributaries that are sources of contamination loading into Lake Ontario.
References
(all urls current as of September 2002)


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