

Phase II GIS-Based Sediment Quality Database for the St. Louis River Area of Concern (AOC)

Help Section for Database Users

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Disclaimer

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List of Acronyms

| | |
|--------------------|--|
| AOC | Area of Concern |
| ARDC | Arrowhead Regional Development Commission |
| AVS | Acid Volatile Sulfide |
| CAS | Chemical Abstracts Service |
| CD | Compact Disk |
| CPRD | Coastal Protection and Restoration Division |
| DBF | Database File |
| DDT | Dichloro-diphenyl-trichloroethane |
| ESRI | Environmental Systems Research Institute |
| GIS | Geographic Information System |
| GLNPO | Great Lakes National Program Office |
| IDL | Instrument Detection Limit |
| IJC | International Joint Commission |
| MARPLOT | Mapping Application for Response, Planning and Local Operational Tasks |
| MDL | Method Detection Limit |
| MESL | MacDonald Environmental Sciences Ltd. |
| MN | Minnesota |
| MPCA | Minnesota Pollution Control Agency |
| MS | Microsoft |
| N | Nitrogen |
| NAD | North American Datum |
| NH ₃ -N | Ammonia Nitrogen |
| NOAA | National Oceanic and Atmospheric Administration |
| PAH | Polycyclic Aromatic Hydrocarbons |
| PCB | Polychlorinated Biphenyls |
| PEC | Probable Effect Concentration |
| PEC-Q | Probable Effect Concentration-Quotient |
| ppm | parts per million |
| PQL | Practical Quantitation Limit |
| QA/QC | Quality Assurance/Quality Control |
| RAP | Remedial Action Plan |
| SEM | Simultaneously Extracted Metals |
| SQL | Structured Query Language |
| SQT | Sediment Quality Target |
| USEPA | United States Environmental Protection Agency |
| UTM | Universal Transverse Mercator |
| WDNR | Wisconsin Department of Natural Resources |
| WI | Wisconsin |

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Chapter 1. Introduction

The St. Louis River constitutes the second largest tributary to Lake Superior. The headwaters begin in northeastern Minnesota (MN), and the lower estuary, which covers an area of approximately 12,000 acres, bisects the border between Duluth, MN and Superior, WI (MPCA and WDNR 1992). The lower estuary culminates in the Duluth-Superior Harbor, which is one of the largest inland seaports in the world and the most heavily used port in the Great Lakes basin.

The middle and lower portions of the estuary support a variety of industrial, commercial, residential, and recreational activities. In addition, these areas provide essential habitats for aquatic organisms (e.g., walleye) and aquatic-dependent wildlife species (e.g., bald eagle). However, aquatic habitats in some of these areas have been adversely affected by economic development of the St. Louis River over the past 130 years.

In 1987, concerns over environmental quality conditions prompted the International Joint Commission (IJC) to designate 43 Areas of Concern (AOCs) in the Laurentian Great Lakes region between the United States and Canada (IJC 1989). The lower 72 nautical kilometers of the St. Louis River from Cloquet, MN to the Duluth, MN and Superior, WI entries to Lake Superior were designated as one of these AOCs. Contaminated sediments contribute to several use impairments in the St. Louis River AOC, including the issuance of fish advisories, restrictions on dredging, and habitat impairments to bottom-feeding organisms. A number of ecosystem health indicators have been selected to support the assessment of sediment quality conditions within the St. Louis River AOC, including sediment chemistry, sediment toxicity, benthic macroinvertebrate community structure, tissue chemistry, the physical characteristics of sediments, and biomarkers in fish (Crane *et al.* 2000). Investigations conducted using data on multiple indicators provide a weight-of-evidence approach for assessing the effects of contaminated sediments on the beneficial uses of this aquatic ecosystem.

As part of the Remedial Action Plan (RAP) process for the St. Louis River AOC, stakeholders identified a need to compile the sediment quality data collected from the St. Louis River in a database format. As a first step, the Arrowhead Regional Development

Commission (ARDC) developed a sediment quality database in 1990 that included all available sediment quality data (31 studies) from the early 1970s to 1990 (ARDC 1990). The sources of these studies included the United States Army Corps of Engineers, United States Environmental Protection Agency (USEPA), Minnesota and Wisconsin (WI) state agencies, contractors, and university researchers. However, no attempt was made to evaluate quality assurance/quality control (QA/QC) procedures in these studies due to a lack of resources. In addition, accurate locational information was not available for most of these sampling stations so the data could not be plotted on maps based on geographic information system (GIS) software.

A matching sediment chemistry and toxicity database was completed in 2000 to support an evaluation of the predictive ability of numerical sediment quality targets (SQTs) in the St. Louis River AOC (Crane *et al.* 2000; 2002). The Minnesota Pollution Control Agency (MPCA) sought additional funding to expand this database with a wider range of sediment quality data (e.g., bioaccumulation, fish tissue, and additional sediment chemistry and toxicity data). In October 2000, the MPCA obtained a grant from the U.S. EPA's Great Lakes National Program Office (GLNPO) to develop the first phase of a GIS-based sediment quality database for the St. Louis River AOC and an associated GIS-mapping component [i.e., GIS data were compiled in Environmental Systems Research Institute (ESRI) ArcView 3.2 format]. MacDonald Environmental Sciences Ltd. (MESL) was retained in April 2001 to assist the MPCA with this effort; MESL had developed the previous matching sediment chemistry and toxicity database for the MPCA. A Quality Assurance Project Plan was completed and approved by GLNPO in July 2001 (Crane 2001) so that work could commence on the project. In October 2001, MESL and MPCA staff met with over 60 stakeholders in Duluth and St. Paul to obtain input on the development of this GIS-based database. Stakeholders were asked to identify priority sediment quality indicators, sources of candidate data sets, and key types of GIS data for the St. Louis River watershed (MacDonald *et al.* 2001). Their input was very useful in producing Phase I of the GIS-based sediment quality database (Smorong and Crane 2003; Smorong *et al.* 2003).

Due to the large amount of sediment quality data that have been collected from the St. Louis River AOC since 1990, funding is being obtained by the MPCA and its collaborators in a phased approach to further the development of the GIS-based sediment quality database. The Duluth office of the MPCA recently changed the scope of GLNPO grant number

GL97540401-2 in order to develop a comprehensive sediment management plan for the lower St. Louis River AOC. Continuation of the GIS-based database (i.e., Phase II) was identified as one task of this grant, for which Judy Crane is the MPCA project manager and Brian Fredrickson is the MPCA grant manager. MESL was retained in September 2003 to complete additional Phase I tasks and to conduct the Phase II updates of the GIS-based database and ArcView 3.2 projects. In addition, MESL was tasked with converting the ArcView 3.2 projects to a version compatible with a more recent version of ESRI's mapping software (i.e., ArcMap 8.3). A Quality Assurance Project Plan was completed and approved by GLNPO in May 2004 (Crane 2004) so that the Phase II work could begin.

Phase III of the GIS-based sediment quality database was initiated in September 2004 through a grant from Minnesota's Lake Superior Coastal Program to the MPCA (grant/project manager is Judy Crane); MESL has been retained to work on this project. Additional sediment quality data sets from the Minnesota portion of the AOC will be added to the Phase III database. The St. Louis River Citizens Action Committee (CAC) in collaboration with the MPCA, Wisconsin Department of Natural Resources (WDNR), and MESL will submit a grant application to the Wisconsin Coastal Management Program in November 2004 to propose conducting Phase IV of the GIS-based sediment quality database. If funded, this project would begin in July 2005 under the project management of Judy Crane (MPCA); additional sediment quality data sets from the Wisconsin portion of the AOC would be added to the database under this proposed phase. In addition, the ArcView 3.2 projects and ArcMap 8.3 map documents would be updated with additional GIS data under the proposed Phase IV project. The expanded database, and associated GIS-mapping components, will support the assessment, preservation, and restoration of the lower St. Louis River AOC and adjoining Lake Superior ecosystems.

The purpose of this Help Section for Database Users is to provide an overview of the MS Access 2000 sediment quality database, a summary of the updates that were completed as part of the Phase II effort, and general instructions for retrieving data from this database. The Help Section is organized into nine chapters and is indexed in such a way as to provide a quick reference guide for users. For more detailed information regarding the content and organization of the GIS-based sediment quality database, users should refer to the accompanying Technical Documentation (Smorong *et al.* 2004a). The Technical

Documentation is available upon request by contacting Judy Crane (MPCA) at 651-297-4068 (voice), 651-297-7709 (fax), or judy.crane@pca.state.mn.us (email).

The sediment quality database was developed in MS Access 2000 format, which is the database format that is available on the project compact disk (CD). The database is also available in MS Access '97 format upon request. The data compiled in the MS Access 2000 database can also be accessed in the National Oceanic and Atmospheric Administrations (NOAA's) St. Louis River Watershed database, which can be viewed using NOAA's free Query Manager software. Query Manager provides a menu of flexible, built-in database queries, and seamless linking to two different mapping applications (ArcView 3.2 and MARPLOT). The advantage that Query Manager offers is the easy-to-use user interface, which is suitable for users with little or no experience using database software. Although there is some loss of flexibility if complex data analyses are necessary, Query Manager offers a wide range of data queries and provides an excellent way for most users to view and query the data. Although the focus of this Help Section for Database Users is designed to help users with the MS Access 2000 version of the database, Chapter 7 provides an overview of the Query Manager software, as well as guidance for accessing and installing the following products available from NOAA's Office of Restoration and Response: Query Manager software, MARPLOT software, the St. Louis River Watershed database, and tools to link Query Manager and ArcView 3.2 [Coastal Protection and Restoration Division (CPRD) tools].

Users should note that this Help Section is not meant to replace formal training in the use of MS Access 2000 software. Microsoft™ provides formal training sessions, a detailed built-in Help section, as well as on-line technical support for MS Access 2000 users. Users should refer to these information sources for detailed guidance on the use of MS Access 2000 software.

Chapter 2. Summary of Phase II Updates

A total of 13 new data sets were added to the MS Access 2000 database as part of the Phase II updates. These data sets included sediment quality data from the federal navigation channels in the Duluth-Superior Harbor and from sites located along the Wisconsin side of the St. Louis River AOC. More details are provided regarding these additions in the accompanying Technical Documentation (Smorong *et al.* 2004a). The database design was nominally refined during the Phase II project. These refinements consisted primarily of modifications of the database field descriptions (modifications are indicated with an asterisk in Table 1). Few changes were made to the remainder of this Help Section, which was previously reported by Smorong and Crane (2003). As such, the GIS-based sediment quality database is not referred to as either Phase I or Phase II for the remainder of this Help Section, unless necessary.

Chapter 3. Database Design

This chapter is intended to provide database users with a description of the design of the GIS-based sediment quality database in MS Access 2000 format. As such, the design of the database is described in terms of the database components (outlines how they are organized), database structure (outlines the content and function of each database component), and database relationships (outlines the connections between database components).

3.1 Organization of Database Components

The GIS-based sediment quality database was designed primarily as a data storage system for sediment quality data collected from the St. Louis River AOC. In designing the database structure, the existing sediment quality data were examined to facilitate the identification of data types, key variables, and required database fields. Some of the factors that were considered during the design of the database included:

- The need to retrieve data by chemical class [e.g., metals, polycyclic aromatic hydrocarbons (PAHs), polychlorinated biphenyls (PCBs), organochlorine pesticides, etc.];
- The need for assuring compatibility with GIS (e.g., ESRI's ArcView 3.2 and ArcMap 8.3 software) to facilitate geographic interpretation of the underlying data;
- The requirement for incorporating ancillary information related to the data (e.g., laboratory qualifier codes, sample dates, sediment descriptions, bibliographic references); and,
- The need for subsequently expanding the database to include other data types (e.g., benthic invertebrate community data, etc.).

The design of the GIS-based sediment quality database was patterned after NOAA's Watershed databases, in which sediment chemistry, sediment toxicity, and tissue chemistry

data have been compiled, and which can be viewed using NOAA's Query Manager software. A key component of this design is that each sample is georeferenced to facilitate spatial analyses of the underlying data and presentation of the information on appropriate base maps (i.e., in ArcView 3.2 or ArcMap 8.3). In order to design a system that most directly met the needs of stakeholders, two meetings were held in October 2001 to garner stakeholder input on a number of data management issues. Stakeholders were asked to describe how they foresee using the database (i.e., the likely ways that they would extract and use the data contained in the database), to identify and prioritize sediment quality indicators, and to identify and prioritize GIS data sets of interest. A comprehensive account of the stakeholder meetings has been compiled in a summary report which is available on the MPCA's Contaminated Sediments Web page at <http://www.pca.state.mn.us/water/sediments/stlouis-stakeholdermtg> (MacDonald *et al.* 2001). Subsequently, the draft database design was reviewed and approved by the MPCA grant and project manager (i.e., Judy Crane) for the Phase I project. Updates to the Phase II database design were approved by the MPCA project manager (Judy Crane).

3.1.1 Description of Database Structure

A detailed description of the database components is provided in Table 1. This table includes a description of the structure and content of each database table, and a description of the information contained in each of the fields (columns) that comprise the tables. To view table descriptions in the database, select the Details icon in the Database window (far right icon in the Database window header), and the table description will be displayed in the Description column. To view field descriptions in the database, see the upper portion of the table in the Design view, or click on a cell in the field of interest and the field description will display in the Status Bar at the bottom of the screen.

3.1.2 Description of Database Relationships

The GIS-based sediment quality database is a relational database. This means that the database consists of several tables that can be linked together (i.e., relationships have been defined) to facilitate retrieval of the data in a wide variety of ways. The purpose of defining relationships is to coordinate the retrieval of information in the different tables. The main

advantage of a relational database is that queries, forms, and reports can be created to display information from several tables at once. A relationship works by matching data in key fields (usually a field with the same name in both tables), and these matching fields provide a unique identifier for each data record. Figure 1 shows the database relationships for the database. The key fields that are used to match the data in different tables, and thus provide a unique identifier, are the ***SITEID***, ***STUDYID***, ***STATIONID***, ***SAMPLEID***, ***FIELDREP***, ***LABREP***, and ***CHEMCODE*** fields. These fields were designed to be consistent with fields used in Query Manager.

Chapter 4. Retrieving Data from the GIS-Based Sediment Quality Database

4.1 Introduction

This chapter is intended to provide database users with instructions to support the retrieval of information from the GIS-based sediment quality database . To fulfill this objective, this chapter includes instructions for designing custom queries in MS Access 2000, along with a description of the different data treatment options that should be considered when designing queries.

4.2 Instructions for Designing Custom Queries

Queries are used to view, change, and analyze data in different ways. The power of queries lies in being able to bring together or perform an action on data from more than one table in the database. For example, it may be desirable to view the concentration of mercury for each sediment sample along with the location of each sample. To see this information, data needs to be extracted from two tables (i.e., **ptbl - CHEM** and **ptbl - SAMPLE**). The most common type of query is a *select query*. A *select query* retrieves data from one or more tables by using specified criteria and then displays it in the desired order (e.g., ascending order based on data value). For more information on queries refer to Appendix 1, which presents instructions for creating *select* queries. In addition, refer to the following topics in MS Access 2000 Help (search for “query” in the Answer Wizard or Index):

- Queries: What they are and how they work;
- Design a query;
- Ways to customize a query;
- Types of queries; and,
- Open or run a query.

Chapter 5. Options for Data Treatment

There are characteristics of sediment quality data (e.g., detection limits for chemical parameters) that require database users to make decisions regarding how to handle the information for the purposes of data analyses. It is essential that these decisions are understood and that the implications of these decisions are considered in the design of database queries (e.g., by including criteria to eliminate certain results and/or samples). This chapter reviews the alternatives that are available for treating undetected data, QA/QC samples, acid volatile sulfide (AVS) and simultaneously extracted metal (SEM) results, and non-numerical or zero results. In addition, there is a summary of the database development decisions that were made regarding how samples were categorized as surficial or sub-surface, how totals were calculated, and how chemical names were standardized.

5.1 Treatment of Undetected Data

Detection limits are estimates of concentrations at which one can be fairly certain that the compound is present. Concentrations below this limit may not be detected. Concentrations above this limit are almost certainly detected in the analysis. Analytical laboratories use several different kinds of detection limits. An instrument detection limit (IDL) is the lowest limit that the instrument can detect. It is determined on samples which have not gone through any sample preparation steps. A method detection limit (MDL) is similar to an IDL, but it is based on samples which have gone through the entire sample preparation scheme prior to analysis. A practical quantitation limit (PQL) is normally 3 to 10 times the MDL and is considered the lowest concentration that can be accurately measured, as opposed to just detected. Detection limits are actually determined by the analysis of low-level samples or blanks. This information gives the variation in instrument response at levels near the detection limit, from which 99% confidence limits are calculated from the standard deviation. The type of detection limit information included in the MS Access 2000 database varied depending on how the detection limits were reported in each study.

A number of investigators have evaluated the implications of applying various procedures for estimating the concentrations of contaminants from undetected data (Gaskin *et al.* 1990; Porter and Ward 1991; El-Shaawari and Esterby 1992; Clarke and Brandon 1994; Clarke 1998). While there is no consensus on which data censoring method should be used in various applications, the simplest methods tend to be used most frequently, including deletion of undetected values or substitution of a constant, such as zero, the detection limit, or one-half the detection limit for the undetected values (USACE 1995).

To address the need for guidance on statistical treatment of undetected data, the United States Army Corps of Engineers (USACE 1995) conducted a simulation study to assess the performance of 10 methods for censoring data. The results of that investigation indicated that no single data censoring method works best in all situations. Accordingly, the USACE recommended a variety of methods depending on the proportion of the data that requires censoring, the distribution and variance of the data, and the type of data transformation to be applied. For data sets for which a low to moderate proportion of the data require censoring, substitution of the detection limit is generally the preferred method [i.e., to optimize statistical power and control Type I error (an error in a statistical test which occurs when a true hypothesis is rejected; a false negative in terms of the null hypothesis)]. However, as the proportion of the data that requires censoring and the coefficient of variation of the data increase, statistical power is better maintained by substituting of one-half the detection limit for the undetected data, particularly for lognormally distributed and transformed data. Substitution of zero or other constants was also recommended in several circumstances. Overall, it was concluded that simple substitution methods work best to maintain power and control error rates in statistical comparisons of chemical concentration data (USACE 1995; Clarke 1998).

The GIS-based sediment quality database was designed so that there are four data treatment options available to the database user for censoring undetected data:

1. Substituting undetected values with one-half the detection limit.

If this is the desired data treatment option, the data user must consider the **MESL_C_CALC** results field in the **ptbl - CHEM** table.

2. Deleting undetected values.

If this is the desired data treatment option, the data user must consider the **CONC** results field in the **ptbl - CHEM** table. In addition, the criteria **Like “NUM”** must be entered in the query design grid for the **MESL_QUAL_CALC** field (i.e., only detected results are included in the query results).

3. Substituting undetected values with the detection limit.

If this is the desired data treatment option, the data user must consider the **CONC** results field in the **ptbl - CHEM** table.

4. Excluding undetected values with high detection limits.

This data treatment option can be used in conjunction with the first and third data treatment options described above. The undetected values with high detection limits [i.e., detection limits greater than the Level II SQTs recommended for use in Minnesota (Crane *et al.* 2000; 2002)] have been identified in the **MESL_EXCLUDE HIGH ND** field. As such, the criteria **Not Like “X”** must be entered in the query design grid for this field to exclude these results.

5.2 Treatment of Data for Quality Assurance/Quality Control Samples

In a number of studies, additional samples were collected and/or analyzed as part of the quality assurance program (i.e., field replicates, analytical laboratory duplicates). The reasons QA/QC samples are analyzed along with environmental samples is to provide information on the accuracy and precision of the analytical results. Data for field replicate and analytical laboratory duplicate samples were included in the database, although other types of QA/QC samples were not included (e.g., method blanks, matrix blanks, trip blanks, etc.). Field replicate samples (i.e., two or more individual samples collected in close proximity to assess the degree of spatial variability in field samples) are identified in the **FIELDREP** field in the **ptbl - SAMPLE** and **ptbl - CHEM** tables. Analytical laboratory duplicates (i.e., one sample

that has been split into two sub-samples and analyzed by the laboratory to assess method precision) are identified in the **MESL_LABDUP_AVG** field in the **ptbl - CHEM** table. It should be noted that laboratory split samples are NOT entered as unique samples, rather, the results from the two sub-samples have been averaged to support subsequent data analysis.

5.3 Treatment of AVS and SEM Results

AVS is produced by bacterial breakdown of organic material in sediments, and it is represented as the sum of iron sulfide, manganese sulfide, and metal sulfides. AVS varies with temperature and depth (i.e., lowest in early spring, highest in summer, and increases with depth). SEM include cadmium, copper, lead, nickel, silver, and zinc that are extracted during the AVS procedure. Assuming that AVS binds a molar equivalent of SEM metal (Di Toro *et al.* 1990), the fraction of SEM metals in excess of AVS concentrations may be available for uptake by benthic biota. Most benthic organisms, including those used in toxicity tests, survive in sediments that have a thin oxidized surface layer and then an anoxic layer. The anoxic layer can have higher AVS concentrations that would reduce the metal activity to which these organisms are exposed (Di Toro *et al.* 1992). When SEM exceeds AVS by a factor of 5 (on a molar basis), a higher incidence of toxicity (80% to 90%) has been observed in freshwater and saltwater amphipod tests (USEPA 1997). Thus, $[SEM] - [AVS] \geq 5$ is a better predictor of sediment toxicity to amphipods.

In the MS Access 2000 database, the results for SEM have been reported in multiple units to support subsequent data analysis. Results for SEM in the **ptbl - CHEM** table are reported in units of parts per million (ppm). This facilitates including both SEM and total metals in data analysis (e.g., making comparisons to Level I and Level II SQTs). The **MESL_SEMQUAL** field is populated with “B” to indicate SEM results for samples that also have results for total metals. The purpose of this field is to exclude SEM results for samples that also have results for total metals (i.e., enter the criteria Not Like “B” to only include total metal results for samples that have results for total metals and SEM).

In addition, the SEM results have been included in the **ptbl - AVS and SEM** table in units of $\mu\text{mol/g}$. This table also includes AVS results reported in units of $\mu\text{mol/g}$ (note that AVS

is **NOT** included in the **ptbl - CHEM** table in units of ppm, as these data are not interpreted on this basis).

5.4 Non-Numerical and Zero Results

There are numerous sediment chemistry results in the database for which either non-numerical (e.g., not reported, missing, not quantified, etc.) or zero results were reported in the original data files. The data treatment decision for zero results was to assume these were undetected results (i.e., “ND” was substituted). The data treatment decision for non-numerical results was to include this information in the database, along with the capacity to exclude these results for the purpose of data analyses. The *MESL_QUAL_CALC* field in the **ptbl - CHEM** table allows the user to select criteria to ensure that only numerical results are considered. This field has been populated with “NUM” (detected), “U” (undetected), “UX” (undetected and detection limit not indicated), and “X” (no result). Therefore, when designing queries, enter NOT LIKE “*X*” criteria in the *MESL_QUAL_CALC* field to consider only numerical results [note the use of the asterisk (*) as it is a wild card]. However, when counting the sample number with undetected results, the criteria should be such that the “UX” results are included (i.e., NOT LIKE “X”, therefore not using the asterisk wild card).

5.5 Categorizing Samples as Surficial or Sub-surface

The **ptbl - SAMPLE** table includes the *MESL_SURF_SUB* field, which designates each sample as either a surficial or sub-surface sample. The criteria that have been used to categorize samples on this basis are consistent with the criteria used by NOAA for the St. Louis River Watershed database. Samples considered to be surficial have an upper sampling depth of zero and a lower sampling depth of less than or equal to 30 cm (i.e., 0 to ≤ 30 cm). Samples considered to be sub-surface have an upper sampling depth either greater than zero or a lower sampling depth of greater than 30 cm (i.e., >0 cm or ≥ 30 cm). For example, a

sample collected at 0-10 cm is categorized as a surficial sample, whereas samples collected at 0-40 cm or 5-10 cm are categorized as sub-surface samples.

5.6 Methods for Calculating Total Chemical Concentrations and Mean PEC-Qs

The **ptbl - CHEM** table includes calculated totals for PAHs, PCBs, and pesticides (i.e., chlordane and DDTs). In addition, mean probable effect concentration quotients (PEC-Qs) have been calculated and included in the **ptbl - CHEM** table, which allows for these results to be queried along with other chemistry results. In addition, the mean PEC-Qs are incorporated in the **ptbl - Mean PEC-Q** table, which includes more information about the PEC-Qs contributing to the mean. Mean PEC-Qs were calculated to provide an overall measure of chemical contamination and to support an evaluation of the combined effects of multiple contaminants in sediments. The mean PEC-Qs have been shown to provide a reliable basis for classifying sediments as toxic or not toxic in the St. Louis River AOC, in the larger geographic areas of the Great Lakes, and elsewhere in North America (Ingersoll *et al.* 2001; Crane *et al.* 2002).

Mean PEC-Qs were calculated using the methods that were recommended by Ingersoll *et al.* (2001) and outlined in Crane *et al.* (2000; 2002). Generally, a PEC-Q was first determined for each metal for which a reliable PEC was available (i.e., arsenic, cadmium, chromium, copper, lead, nickel, and zinc). Then, an average PEC-Q for metals was calculated by summing the PEC-Qs of each metal and dividing by the number of metals that were included in the calculation. PEC-Qs were also calculated for total PAHs (based on a subset of the 13 parent low molecular weight and high molecular weight PAHs) and total PCBs. Finally, the mean of the average PEC-Q for metals, the PEC-Q for total PAHs, and the PEC-Q for total PCBs was determined for each sediment sample (termed the mean PEC-Q).

The CHEMCODEs that delineate calculated totals can be identified by referring to the records in the **lkp - CHEMDICT** table that have “TOTAL” entered in the **CHEMCLASS**

field. More details regarding the methods for calculating totals and mean PEC-Qs are included in the **ptbl - STUDYNOT** table.

5.7 Standardization of Chemical Names

One of the main challenges associated with compiling several data sets into a database so the data are comparable and can be combined for the purpose of data analyses is standardizing the chemical names used in the individual data sources. Many chemicals typically have several synonyms (e.g., “ammonia” is commonly referred to as ammonia-N, ammonia, ammonia-total, ammonia as N, NH₃-N, etc.). A problem arises because data from each study were generated at different analytical laboratories, each potentially using their own list of chemical names to report the results. There are a variety of systems that provide unique identifiers for chemical substances to provide an unambiguous way to identify a chemical substance when there are many possible systematic, generic, proprietary, or trivial names [e.g., Chemical Abstracts Service (CAS)].

In order to handle the issue of standardizing chemical names in the database, a standard chemical code (i.e., the **CHEMCODE** field) was assigned for each chemical name. Assigning a short code to represent long chemical names also reduces the chance of error in entering long chemical names that must be exact to ensure that related data tables are correctly connected. The **lkp - CHEMDICT** table lists each of the unique chemical codes (i.e., **CHEMCODE**) and provides the full chemical name and other associated information that is helpful in the evaluation of the data (e.g., chemical classification, standard units of measure, etc.). It is especially important to refer to the **CHEMNAME** field in the **lkp - CHEMDICT** table when interpreting particle size data, as the micron size applicable to that **CHEMCODE** is specified in this field.

Chapter 6. Linking the GIS-Based Sediment Quality Database with Accompanying ArcView 3.2 Projects and ArcMap 8.3 Map Documents

The watershed GIS data that were compiled in the accompanying ArcView 3.2 projects and ArcMap 8.3 map documents provide a basis for users to spatially view and interpret the data incorporated in the sediment quality database (see the Help Section for ArcView Users for further information; Smorong *et al.* 2004b). The GIS maps developed for this project contain three basemap options in which to overlay the locations of sediment sample locations contained in the database: digital orthographic aerial photographs, digital orthographic topographic maps, and basic line and polygon theme data. Ten maps were produced using the GIS applications, each containing GIS watershed data categorized by major theme (e.g., contaminated areas, ecological areas, hydrology, land use, etc.), as well as the three basemap options. The ten maps have been produced in ArcView 3.2 and ArcMap 8.3, with very few differences between the two versions (i.e., database users can use either one or the other GIS application, but do not need to access both versions).

In order to plot the data in GIS applications, the data must have associated geographical coordinates. The sediment quality database stores the UTM Zone 15 NAD 83 coordinates in the **ptbl - STATION** table. As such, the user must combine the data of interest (e.g., mercury concentrations, toxic/not toxic designations for 10-day *Hyaella azteca* survival toxicity tests) with these geographical coordinates. To accomplish this task, a query must be designed and saved (see Appendix 1 for instructions for designing custom queries in MS Access 2000).

Data can be imported into any of the GIS maps using ArcView's Structured Query Language (SQL) connection feature or ArcCatalog to establish a database connection to ArcMap. These features allow users to query a database and to store the returned records in the GIS application. Chapter 5 of the Help Section for ArcView Users provide instructions for linking the GIS applications with the MS Access 2000 version of the GIS-based sediment quality database (Smorong *et al.* 2004b). Query results obtained from NOAA's Query Manager software can also be seamlessly linked to the ArcView 3.2 projects. Users should

refer to Chapter 7 for additional information regarding the accessibility, installation, and set-up of the Query Manager software, as well as information on how to link the Query Manager and ArcView applications.

Chapter 7. Database and Mapping Applications Available from NOAA

7.1 Introduction

Protecting and restoring coastal watersheds involves understanding an array of complex environmental issues and synthesizing various kinds of information. The challenge of evaluating multiple environmental issues is made easier by combining scientific data and watershed characteristics into a GIS. NOAA's Coastal Protection and Restoration Division (CPRD) has developed numerous Watershed Database and Mapping Projects that combine a standard database structure (Query Manager) with a database-mapping application (MARPLOT) and GIS tools. Sediment chemistry concentrations, sediment toxicity and tissue data, natural resource data, and potential habitat restoration projects can be overlaid on a watershed's features and land uses, and displayed on maps at flexible spatial scales.

The data compiled in the MS Access 2000 database will be converted and included in NOAA's St. Louis River Watershed database, which can be viewed using NOAA's Query Manager software. Query Manager provides a menu of flexible, built-in database queries, and provides seamless linking to two different mapping applications (ArcView 3.x and MARPLOT). The advantage that Query Manager offers is the easy-to-use user interface, which is suitable for users with little or no experience using database software. Although there is some loss of flexibility if complex data analyses are necessary, Query Manager offers a wide range of data queries and provides an excellent way for most users to view and query the data.

7.2 Accessing NOAA's Applications

The following products are available for free from NOAA's Office of Restoration and Response Web page:

(<http://response.restoration.noaa.gov/cpr/watershed/watershedtools.html>):

- Query Manager 2.5 software (data delivery application that offers a menu of flexible, built-in database queries);
- MARPLOT 3.3 software (a mapping application that can be seamlessly linked to display query results from Query Manager software; this software is best used by people lacking ArcView 3.x);
- NOAA's CPRD Tools (a collection of GIS tools created for ArcView 3.x to assist with the development and analysis of spatial data); and,
- The St. Louis River Watershed database.

Each of these applications is accompanied by detailed instructions for installation and use. Users should note the following details regarding NOAA's St. Louis River Watershed database:

- The Watershed database will be updated in late 2004 or early 2005 to incorporate all of the data included in the Phase I MS Access 2000 sediment quality database for the St. Louis River AOC. Inclusion of updated versions of the GIS-based sediment quality database in future updates of NOAA's Watershed database for the St. Louis River will depend on NOAA's financial resources;
- The data sets that are unique to the NOAA St. Louis River Watershed database (i.e., not included in the Phase I MS Access 2000 database) have not undergone the same data evaluation procedures as the data sets incorporated in the Phase I MS Access 2000 database;
- Users should be aware that there are other minor differences between the NOAA Watershed database and the MS Access 2000 sediment quality database (e.g., any field prefixed by "MESL_" in the MS Access 2000 database is not included in NOAA's database, mean PEC-Qs are calculated using different methods, field replicate samples are treated differently, etc.); and,
- Under the "Data and Maps" section of NOAA's Web page, the date of the most recent database update is provided (users should check this web page periodically for future updates).

7.3 Converting the MS Access 2000 Database to Query Manager Format

Some users may wish to use the Query Manager software with the version of the data compiled in the MS Access 2000 database, particularly because the mean PEC-Qs (as described in Section 5.6) are included in the **ptbl - CHEM** table. If so, contact Judy Crane (MPCA) at 651-297-4068 (voice), 651-297-7709 (fax), or judy.crane@pca.state.mn.us (email) for a copy of the Query Manager-compatible database files prepared by MESL staff from the MS Access 2000 database. As a short-hand descriptor, this database will be referred to as the Phase II “MESL Query Manager” database to avoid confusion with NOAA’s Watershed database for the St. Louis River.

Users can also make this conversion themselves by following the step by step instructions given below for converting the MS Access 2000 database from the project CD to a format compatible with Query Manager version 2.5:

1. Split the **ptbl - SAMPLE** table into two tables, one named “SAMPLE” which contains the surficial samples, and one named “SMPSEDSB” which contains the sub-surface samples (identified in the **MESL_SURF_SUB** field).
2. Split the **ptbl - CHEM** table into two tables, one named “CHEM” which contains the results for surficial samples, and one named “CHEMSB” which contains the results for sub-surface samples. This step will require the user to build a make-table query:
 - Add the **ptbl - CHEM** table and the **ptbl - SAMPLE** tables;
 - Change the Query Type to “Make-table” and enter the Table Name as CHEM;
 - Include all fields from the **ptbl - CHEM** table and the **MESL_SURF_SUB** field from the **ptbl - SAMPLE** table;
 - Click the check mark in the **MESL_SURF_SUB** field to uncheck it;
 - Enter “Surficial” as criteria in the **MESL_SURF_SUB** field;

- Run the query (click the button with the exclamation mark);
 - Change the criteria in the **MESL_SURF_SUB** field to “subsurface”;
 - Click on the Query Type button and change the Table Name to CHEMSB; and,
 - Run the query (click the button with the exclamation mark).
3. In the CHEM and CHEMSB tables, update the **MISSINGVAL** field to “-1” where there is an “X” in the **MESL_EXCLUDE HIGH ND** field. This will exclude the undetected results with detection limits greater than the Level II SQT when running queries in Query Manager.
4. In the CHEM and CHEMSB tables, update the **MISSINGVAL** field to “-1” where there is an “*X*” in the **MESL_QUAL_CALC** field. This will exclude the results that were flagged as unacceptable due to data quality concerns.
5. Export the following tables as database file (DBF) files to the C:\qm25win\stlouis\sl_data folder. Note that the table names must not include the “ptbl” prefix. Note that the following tables will be replaced if the St. Louis River Watershed database has previously been downloaded from NOAA’s web site.
- BIOSUMM
 - CHEM
 - CHEMSB (this is a table that has just been created)
 - CHEMTISS
 - QUALIFY
 - SAMPLE
 - SITE
 - SMPSEDSB (this is a table that has just been created)
 - SMPTISS
 - STATION
 - STUDY
 - STUDYNOT
 - STUDYREF

6. Export the following tables as DBF files to the C:\qm25win folder. Note that the table names must not include the “lqp” prefix. The following tables will be replaced in this step.
- CHEMDICT
 - SPECIES
 - SQC
 - SQCDICT
 - SQCPAIRS
 - TESTDICT
 - TISSTYPE
7. Proceed with using the Phase II “MESL Query Manager” database.

Although every effort has been made to ensure the database is compatible with NOAA’s Query Manager software, users should note that NOAA has not conducted a formal review of the MESL Query Manager database, nor the procedures for converting the MS Access 2000 database to a format compatible with Query Manager.

Chapter 8. Project Contact

For further information about the Phase II MS Access 2000 or Phase II “MESL Query Manager” sediment quality databases for the St. Louis River AOC, contact Judy Crane at:

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Users will be notified when additional phases of this project have been completed.

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Table 1. Detailed Description of Database Components (asterisks indicate changes from the Phase I database design).

| TABLE NAME / Field Name | Data Type | Field Size | TABLE DESCRIPTION / Field Description |
|-----------------------------------|-----------|------------|---|
| lkp - CHEMDICT¹ | NA | NA | Chemical dictionary (only included chemicals represented in the ptbl - CHEM table). |
| CHEMCODE | Text | 10 | Chemical code (defined in lkp - chemdict). |
| CHEMNAME | Text | 45 | Chemical name. |
| CHEMCLASS | Text | 8 | Chemical class.* |
| CATEGORY | Text | 8 | Query Manager field (not populated). |
| SUBCATGY | Text | 10 | Query Manager field (not populated). |
| CHEMTOTAL | Text | 10 | Query Manager field (not populated). |
| MOLWT | Number | 8 | Molecular weight of chemical. |
| CASNUM | Text | 24 | Chemical abstract services number. |
| UNITS | Text | 6 | Units of chemical concentration. |
| WA_UNITS | Text | 6 | Query Manager field (not populated). |
| EDITDATE2 | Text | 8 | Query Manager field (not populated). |
| EDITBY | Text | 15 | Query Manager field (not populated). |
| EDITDATE | Date/Time | 8 | Query Manager field (not populated). |
| MESL_synonym | Text | 150 | Synonym for chemical name. |
| MESL_syn2 | Text | 150 | Synonym for chemical name. |
| MESL_syn3 | Text | 150 | Synonym for chemical name. |
| lkp - QUALIFY | NA | NA | Lookup table for sediment chemistry qualifiers (QUALCODE). |
| SITEID | Text | 4 | Site ID code (from Query Manager). |
| STUDYID | Text | 2 | Study ID code. |
| QUALCODE | Text | 5 | Qualifier code for concentration value, modified to be compatible with Query Manager (all ND data has a "U" in this field). |
| QUALIFIERS | Text | 30 | Qualifier code for concentration value, as designated in report. |
| DESCRIPT | Text | 80 | Description of the meaning of the qualifier, as indicated in the original report or data file. |
| MESL_LIBNO | Text | 50 | MESL - library number. |

Table 1. Detailed Description of Database Components (asterisks indicate changes from the Phase I database design).

| TABLE NAME / Field Name | Data Type | Field Size | TABLE DESCRIPTION / Field Description |
|-------------------------|-----------|------------|---|
| lkp - SPECIES | NA | NA | Lookup table for tissue samples species type (SPP). |
| SPP | Text | 5 | Species code. |
| COMMONNAME | Text | 25 | Common name. |
| SCIENTIFIC | Text | 40 | Scientific name. |
| GROUP | Text | 25 | Query Manager field (not populated). |
| EDITDATE2 | Text | 8 | Query Manager field (not populated). |
| EDITBY | Text | 15 | Query Manager field (not populated). |
| EDITDATE | Date/Time | 8 | Query Manager field (not populated). |
| MESL_COMMENT | Text | 50 | MESL - comments. |
| lkp - SQC | NA | NA | Sediment Quality Criteria: Level I and Level II SQTs (Crane <i>et al.</i> 2000) and SQGs (WDNR 2003).* |
| SQCCODE | Text | 10 | Code for Sediment Quality Criteria (see lkp - Sqcdict for a description of the codes). |
| CHEMCODE | Text | 10 | Chemical code (defined in lkp - chemdict). |
| CONC | Number | 8 | Chemical concentration. |
| UNITS | Text | 6 | Units SQC is reported in.* |
| NORM | Text | 2 | Indicates measurement basis SQC are reported in. |
| MESL_chemical name | Text | 50 | MESL - chemical name. |
| MESL_units | Text | 50 | MESL - units. |
| MESL_comment | Text | 100 | MESL - comments. |
| lkp - Sqcdict | NA | NA | Lookup table for Sediment Quality Criteria references (SQCCODE). |
| SQCCODE | Text | 10 | Code for Sediment Quality Criteria. |
| SQCDESCR | Text | 90 | Description of the SQCCODE. |
| YEAR | Text | 4 | Year of publishing for study reporting the sediment quality criteria. |
| AUTHORS | Text | 160 | Authors for study reporting the sediment quality criteria. |
| TITLE | Text | 160 | Title of the study reporting the sediment quality criteria. |
| SOURCE | Text | 160 | Source (location) for study reporting the sediment quality criteria. |
| COMMENT | Text | 160 | Comments. |

Table 1. Detailed Description of Database Components (asterisks indicate changes from the Phase I database design).

| TABLE NAME / Field Name | Data Type | Field Size | TABLE DESCRIPTION / Field Description |
|-------------------------|-----------|------------|---|
| lkp - SQCPAIRS | NA | NA | Lookup table for identifying Sediment Quality Criteria pairs. |
| PAIRNAME | Text | 50 | High and low Sediment Quality Criteria (how Query Manager will reference the pair). |
| SQCLOW | Text | 10 | SQCCODE of low Sediment Quality Criteria. |
| SQCHIGH | Text | 10 | SQCCODE of high Sediment Quality Criteria. |
| LOW NAME | Text | 8 | Low Sediment Quality Criteria (how Query Manager will reference the SQC). |
| HIGH NAME | Text | 8 | High Sediment Quality Criteria (how Query Manager will reference the SQC). |
| SORT ORDER | Number | 2 | Query Manager field (did not populate). |
| lkp - TESTDICT | NA | NA | Lookup table for toxicity test dictionary (TESTID). |
| TESTID | Text | 12 | Code describing the bioassay. |
| MEDIUM | Text | 15 | Medium used in toxicity test (e.g., bulk sediment or pore water). |
| MEDCODE | Text | 2 | Code used to indicate medium used in toxicity test. |
| GROUP | Text | 20 | Group of organism used in toxicity test (e.g., bacteria or amphipod). |
| ALTGROUP | Text | 20 | Group of organism used in toxicity test - alternate. |
| SPECIES | Text | 40 | Species used in toxicity test. |
| SPPCODE | Text | 3 | Code used to indicate species used in toxicity test. |
| LHS | Text | 10 | Life stage of organism used in toxicity test. |
| LHSCODE | Text | 1 | Code used to indicate life stage of organism used in toxicity test. |
| ENDPOINT | Text | 30 | Endpoint of toxicity test (e.g., growth or survival). |
| ENDCODE | Text | 2 | Code used to indicate endpoint of toxicity test. |
| DURATION | Text | 10 | Duration of toxicity test. |
| DURCODE | Text | 4 | Code used to indicate duration of toxicity test. |
| HABITAT | Text | 2 | Query Manager field (not populated). |
| EDITDATE | Date/Time | 8 | Query Manager field (not populated). |
| EDITDATE2 | Text | 8 | Query Manager field (not populated). |
| EDITBY | Text | 15 | Query Manager field (not populated). |
| lkp - TISSQUAL | NA | NA | Lookup table for tissue chemistry qualifiers (QUALCODE). |
| SITEID | Text | 4 | Site ID code (from Query Manager). |
| STUDYID | Text | 2 | Study ID code. |

Table 1. Detailed Description of Database Components (asterisks indicate changes from the Phase I database design).

| TABLE NAME / Field Name | Data Type | Field Size | TABLE DESCRIPTION / Field Description |
|-------------------------------|-----------|------------|---|
| lkp - TISSQUAL (cont.) | | | |
| QUALCODE | Text | 5 | Qualifier code for concentration value, modified to be compatible with Query Manager (all ND data has a "U" in this field). |
| QUALIFIERS | Text | 30 | Qualifier code for concentration value, as designated in report. |
| DESCRIPT | Text | 80 | Description of the meaning of the qualifier, as indicated in the original report or data file. |
| MESL_LIBNO | Text | 50 | MESL - library number. |
| lkp - TISSTYPE | | | |
| TISSCODE | NA | NA | Lookup table for tissue sample tissue types (TISSCODE). |
| DESCRIPT | Text | 6 | Tissue type code. |
| EDITDATE | Text | 50 | Description of tissue type. |
| EDITDATE2 | Date/Time | 8 | Query Manager field (not populated). |
| EDITBY | Text | 8 | Query Manager field (not populated). |
| | Text | 15 | Query Manager field (not populated). |
| ptbl - AVS_SEM | | | |
| | NA | NA | Sediment chemistry results for Acid Volatile Sulfides and Simultaneously Extracted Metals (units of $\mu\text{mol/g}$). |
| SITEID | Text | 4 | Site ID code (from Query Manager). |
| STUDYID | Text | 2 | Study ID code. |
| STATIONID | Text | 6 | Station ID code (this is the MESL_STATIONID, unless it exceeded 6 characters, then the QM_STATIONID was substituted). |
| SAMPLEID | Text | 2 | Sample ID code. |
| FIELDREP | Text | 2 | Identifies field replicate samples (samples collected in close proximity). |
| LABREP | Text | 2 | This field will not be populated (lab dups will be averaged and the MESL_LABREP_AVG field identifies these averaged results). |
| CHEMCODE | Text | 10 | Chemical code (defined in lkp - CHEMDICT). |
| QUALCODE | Text | 5 | Qualifier code for concentration value, modified to be compatible with Query Manager (all ND data has a "U" in this field). |
| CONC | Number | 8 | Chemical concentration (dry weight basis). |
| UNITS | Text | 6 | Units of chemical concentration. |

Table 1. Detailed Description of Database Components (asterisks indicate changes from the Phase I database design).

| TABLE NAME / Field Name | Data Type | Field Size | TABLE DESCRIPTION / Field Description |
|-------------------------------|-----------|------------|--|
| ptbl - AVS_SEM (cont.) | | | |
| MEASBASIS | Text | 2 | Measurement basis - dry weight (DW). |
| MISSINGVAL | Yes/No | 1 | 'Yes' indicates a missing value (i.e., not reported, not sampled, lost, etc; -9 entered in CONC field). |
| MESL_LIBNO | Text | 20 | MESL - library number. |
| MESL_STATIONID | Text | 50 | MESL - station ID (retains the Station ID code as it appears in the original datafiles and/or reports). |
| MESL_UNITS | Text | 7 | MESL - units of concentration value. |
| MESL_QUAL_CALC | Text | 10 | MESL - qualifier code for concentration value - used for calculation purposes (NUM - value, U - less than detect value; X - do not include in calculations). |
| MESL_semqual | Text | 50 | MESL - qualifier code to indicate whether to use SEM metal conc. or total metal conc. (B entered in this field indicates that both are measured, therefore do not use the SEM result). |
| MESL_C_TXT | Text | 50 | MESL - concentration value represented in a text field (nondetected results include a "<"). |
| MESL_LABDUP_AVG | Yes/No | 1 | MESL - indicates if results for laboratory duplicate samples were averaged.* |
| MESL_C_CALC | Number | 8 | MESL - concentration value represented in a number field (nondetected results included as 1/2 the detection limit). |
| MESL_EXCLUDE HIGH ND | Text | 50 | MESL - X entered in this field indicates a nondetected result with a detection limit greater than the Level II SQT. |
| MESL_comment | Text | 250 | MESL - comments. |
| ptbl - BIOSUMM | | | |
| | NA | NA | Sediment toxicity test and bioaccumulation test results. |
| SITEID | Text | 4 | Site ID code (from Query Manager). |
| STUDYID | Text | 2 | Study ID code. |
| STATIONID | Text | 6 | Station ID code (this is the MESL_STATIONID, unless it exceeded 6 characters, then the QM_STATIONID was substituted). |
| SAMPLEID | Text | 2 | Sample ID code. |
| FIELDREP | Text | 2 | Identifies field replicate samples (samples collected in close proximity). |
| TESTID | Text | 12 | Code describing the bioassay (see lkp - TESTDICT table for a description of the codes). |
| GROUP | Text | 2 | Query Manager field (did not populate). |
| SERIES | Text | 2 | Associates control sample results with test results. |
| EFFECTVAL | Number | 8 | Toxicity test result (e.g., percent survival). |

Table 1. Detailed Description of Database Components (asterisks indicate changes from the Phase I database design).

| TABLE NAME / Field Name | Data Type | Field Size | TABLE DESCRIPTION / Field Description |
|-------------------------------|-----------|------------|---|
| ptbl - BIOSUMM (cont.) | | | |
| SIGEFFECT | Yes/No | 1 | Toxic (-1) or Not toxic (0). |
| NEG | Yes/No | 1 | Negative control sample? Yes (-1) or No (0). |
| REF | Yes/No | 1 | Reference sample? Yes (-1) or No (0). |
| STAT | Yes/No | 1 | Identifies sample used to determine significance (T/NT) - ND results (i.e., growth endpoint not measured because of low survival) from SQT database added as NOT TOXIC. |
| SIG_ORIGIN | Text | 50 | Original significance designations - from QM database. |
| CTRLADJ | Number | 8 | Control adjusted result (test result/control result*100). |
| TOXCODE | Text | 1 | Query Manager field (did not populate). |
| MESL_LIBNO | Text | 50 | MESL - library number. |
| MESL_STATIONID | Text | 50 | MESL - Station ID (retains the Station ID code as it appears in the original datafiles and/or reports). |
| MESL_TOXIC | Text | 2 | Toxic (T), Not toxic (NT), or ND (growth endpoint not measured because of low survival). |
| MESL_comment | Text | 250 | MESL - comments. |
| ptbl - CHEM | | | |
| | NA | NA | Chemistry results for sediment samples.* |
| SITEID | Text | 4 | Site ID code (from Query Manager). |
| STUDYID | Text | 2 | Study ID code. |
| STATIONID | Text | 6 | Station ID code (this is the MESL_STATIONID, unless it exceeded 6 characters, then the QM_STATIONID was substituted). |
| SAMPLEID | Text | 2 | Sample ID code. |
| FIELDREP | Text | 2 | Identifies field replicate samples (samples collected in close proximity). |
| LABREP | Text | 2 | This field will not be populated (lab dups will be averaged and the MESL_LABREP_AVG field identifies these averaged results). |
| CHEMCODE | Text | 10 | Chemical code (defined in lkp - CHEMDICT). |
| QUALCODE | Text | 5 | Qualifier code for concentration value, modified to be compatible with Query Manager (all ND data has a "U" in this field). |
| CONC | Number | 8 | Chemical concentration (dry weight basis) |
| UNITS | Text | 6 | Units of chemical concentration. |

Table 1. Detailed Description of Database Components (asterisks indicate changes from the Phase I database design).

| TABLE NAME / Field Name | Data Type | Field Size | TABLE DESCRIPTION / Field Description |
|----------------------------|-----------|------------|--|
| ptbl - CHEM (cont.) | | | |
| MEASBASIS | Text | 2 | Measurement basis - dry weight (DW). |
| MISSINGVAL | Yes/No | 1 | 'Yes' indicates a missing value (i.e., not reported, not sampled, lost, etc; -9 entered in CONC field). |
| MESL_LIBNO | Text | 20 | MESL - library number. |
| MESL_STATIONID | Text | 50 | MESL - station ID (retains the Station ID code as it appears in the original datafiles and/or reports). |
| MESL_UNITS | Text | 7 | MESL - units of concentration value. |
| MESL_QUAL_CALC | Text | 10 | MESL - qualifier code for concentration value - used for calculation purposes (NUM - value, U - less than detect value; X - do not include in calculations). |
| MESL_semqual | Text | 50 | MESL - qualifier code to indicate whether to use SEM metal conc. or total metal conc. (B entered in this field indicates that both are measured, therefore do not use the SEM result). |
| MESL_C_TXT | Text | 50 | MESL - concentration value represented in a text field (nondetected results include a "<"). |
| MESL_LABDUP_AVG | Yes/No | 1 | MESL - indicates if results for laboratory duplicate samples were averaged.* |
| MESL_C_CALC | Number | 8 | MESL - concentration value represented in a number field (nondetected results included as 1/2 the detection limit). |
| MESL_EXCLUDE HIGH ND | Text | 50 | MESL - X entered in this field indicates a nondetected result with a detection limit greater than the Level II SQT. |
| MESL_comment | Text | 250 | MESL - comments. |
| ptbl - CHEMTISS | | | |
| | NA | NA | Chemistry results for tissue samples.* |
| SITEID | Text | 4 | Site ID code (from Query Manager). |
| STUDYID | Text | 2 | Study ID code. |
| STATIONID | Text | 10 | Station ID code (this is the MESL_STATIONID, unless it exceeded 6 characters, then the QM_STATIONID was substituted). |
| SAMPLEID | Text | 2 | Sample ID code. |
| FIELDREP | Text | 2 | Identifies field replicate samples (samples collected in close proximity). |
| LABREP | Text | 2 | Not populated (lab dups will be averaged and the MESL_LABREP_AVG field identifies these averaged results). |
| CHEMCODE | Text | 10 | Chemical code (defined in lkp - chemdict) |

Table 1. Detailed Description of Database Components (asterisks indicate changes from the Phase I database design).

| TABLE NAME / Field Name | Data Type | Field Size | TABLE DESCRIPTION / Field Description |
|--------------------------------|-----------|------------|---|
| ptbl - CHEMTISS (cont.) | | | |
| CONC | Number | 8 | Chemical concentration (wet weight basis). |
| QUALCODE | Text | 5 | Qualifier code for concentration value, as designated in report (see lkp_TISSQUAL table for a description of the codes). |
| UNITS | Text | 6 | Units of chemical concentration. |
| MEASBASIS | Text | 2 | Measurement basis - wet weight (WW). |
| MISSINGVAL | Yes/No | 1 | 'Yes' indicates a missing value (i.e., not reported, not sampled, lost, etc; -9 entered in CONC field). |
| MESL_LIBNO | Text | 20 | MESL - library number. |
| MESL_STATIONID | Text | 50 | MESL - Station ID (retains the Station ID code as it appears in the original datafiles and/or reports). |
| MESL_UNITS | Text | 7 | MESL - units of concentration value. |
| MESL_QUAL_CALC | Text | 10 | MESL - qualifier code for concentration value - used for calculation purposes (NUM - value, U - less than detect value; X - do not include in calculations; UX - less than MDL, detection limit not known). |
| MESL_comment | Text | 250 | MESL - comments. |
| MESL_CONC_TXT | Text | 50 | MESL - concentration value represented in a text field (nondetected results include a "<").* |
| MESL_LABREP_AVG | Yes/No | 1 | MESL - indicates if results for laboratory duplicate samples were averaged.* |
| ptbl - Mean PEC-Q | | | |
| | NA | NA | Mean Probable Effect Concentration-Quotients (Mean PEC-Q). NOT A QUERY MANAGER TABLE. |
| SITEID | Text | 4 | Site ID code (from Query Manager). |
| STUDYID | Text | 2 | Study ID code. |
| STATIONID | Text | 6 | Station ID code (this is the MESL_STATIONID, unless it exceeded 6 characters, then the QM_STATIONID was substituted). |
| SAMPLEID | Text | 2 | Sample ID code. |
| FIELDREP | Text | 2 | Identifies field replicate samples (samples collected in close proximity). |
| LABREP | Text | 2 | This field will not be populated (lab dups will be averaged and the MESL_LABREP_AVG field identifies these averaged results). |

Table 1. Detailed Description of Database Components (asterisks indicate changes from the Phase I database design).

| TABLE NAME / Field Name | Data Type | Field Size | TABLE DESCRIPTION / Field Description |
|----------------------------------|-----------|------------|---|
| ptbl - Mean PEC-Q (cont.) | | | |
| PECQ met | Number | 8 | PEC quotient for metals. |
| PECQ pah | Number | 8 | PEC quotient for PAHs. |
| PECQ pcb | Number | 8 | PEC quotient for PCBs. |
| MeanPECQ | Number | 8 | Mean PEC quotient (as calculated). |
| MeanPECQ (3 sf) | Number | 8 | Mean PEC quotient (3 significant figures). |
| ptbl - SAMPLE | NA | NA | Sediment sample information.* |
| SITEID | Text | 4 | Site ID code (from Query Manager). |
| STUDYID | Text | 2 | Study ID code. |
| STATIONID | Text | 6 | Station ID code (this is the MESL_STATIONID, unless it exceeded 6 characters, then the QM_STATIONID was substituted). |
| SAMPLEID | Text | 2 | Sample ID code. |
| FIELDREP | Text | 2 | Identifies field replicate samples (samples collected in close proximity). |
| LABREP | Text | 2 | This field will not be populated (lab dups will be averaged and the MESL_LABREP_AVG field identifies these averaged results). |
| UDEPTH | Number | 8 | Upper sampling depth (cm). |
| LDEPTH | Double | 8 | Lower sampling depth (cm). |
| SAMPDATE | Text | 8 | Sample date (YYYYMMDD). |
| SAMPTIME | Text | 5 | Sample time. |
| TOC | Number | 8 | Total organic carbon (%). |
| PCTFINES | Number | 8 | Percent fines (sand + clay), the micron diameter used to define PCTFINES is <53 µm, unless otherwise noted in the MESL_comments field.* |
| UAN_PW | Number | 8 | Unionized ammonia in pore water. |
| H2S_PW | Number | 8 | Hydrogen sulfide in pore water. |
| EXSAMPID | Text | 15 | Original station ID reported in study or data file. |
| MESL_LIBNO | Text | 50 | MESL - library number. |
| MESL_STATIONID | Text | 50 | MESL - station ID (retains the Station ID code as it appears in the original datafiles and/or reports). |
| MESL_Mean PEC-Q | Number | 8 | MESL - Mean PEC-Q (3 significant figures). |

Table 1. Detailed Description of Database Components (asterisks indicate changes from the Phase I database design).

| TABLE NAME / Field Name | Data Type | Field Size | TABLE DESCRIPTION / Field Description |
|------------------------------|-----------|------------|---|
| ptbl - SAMPLE (cont.) | | | |
| MESL_MATCH | Text | 50 | MESL - indicates if the sample has matching sediment chemistry and toxicity data. |
| MESL_WATERDEPTH | Text | 50 | MESL - water depth at the point of sediment sampling (m). NR = not reported.* |
| MESL_SOFTDEPTH | Text | 50 | MESL - soft sediment depth (m). NR = not reported.* |
| MESL_SEDDISC | Text | 255 | MESL - sediment description (have included the sediment description if this data was available electronically). NA = not available.* |
| MESL_SURF_SUB | Text | 50 | MESL - indicates if the sample is designated as surficial or sub-surface, according to NOAA's Query Manager rules. NA indicates that the sampling depth was not specified.* |
| MESL_comments | Text | 255 | MESL - comments. |
| ptbl - SITE | NA | NA | Query Manager table. |
| SITEID | Text | 4 | Site ID code (from Query Manager). |
| SITENAME | Text | 40 | |
| EPAREGION | Number | 2 | |
| COUNTY | Text | 25 | |
| STATE | Text | 2 | |
| CERCLIS | Text | 12 | |
| REACH | Text | 8 | |
| REACHSEG | Text | 11 | |
| LATITUDE | Number | 8 | |
| LONGITUDE | Number | 8 | |
| WATERSHED | Text | 20 | |
| ptbl - SMPTISS | NA | NA | Tissue sample information. |
| SITEID | Text | 4 | Site ID code (from Query Manager). |
| STUDYID | Text | 2 | Study ID code. |
| STATIONID | Text | 10 | Station ID code (this is the MESL_STATIONID, unless it exceeded 6 characters, then the QM_STATIONID was substituted). |

Table 1. Detailed Description of Database Components (asterisks indicate changes from the Phase I database design).

| TABLE NAME / Field Name | Data Type | Field Size | TABLE DESCRIPTION / Field Description |
|-------------------------------|-----------|------------|---|
| ptbl - SMPTISS (cont.) | | | |
| SAMPLEID | Text | 2 | Sample ID code. |
| FIELDREP | Text | 2 | Identifies field replicate samples (samples collected in close proximity). |
| LABREP | Text | 2 | Not populated (lab dups will be averaged and the MESL_LABREP_AVG field identifies these averaged results). |
| SAMPDATE | Text | 8 | Date sample collected (YYYYMMDD). |
| SAMPTIME | Text | 5 | Time sample collected. |
| SPECIES | Text | 5 | Species from which the tissue sample was collected (see the lkp_SPECIES table for a description of the codes). |
| SPP | Text | 5 | Species code (see the lkp_SPECIES table for a description of the codes). |
| TISSUE | Text | 30 | Tissue type analyzed (see the lkp_TISSTYPE table for a description of the codes). |
| TISSCODE | Text | 6 | Tissue type code (see the lkp_TISSTYPE table for a description of the codes). |
| LIFESTAGE | Text | 1 | Lifestage of the organism at the time of sampling (not populated). |
| NOINCOMP | Number | 2 | Number of individuals in a composite sample. |
| LENGTH | Number | 8 | Length (cm) of individual organisms collected for tissue analysis. |
| WEIGHT | Number | 8 | Weight of individual organisms collected for tissue analysis (not populated). |
| SEX | Text | 1 | Sex of individual organisms collected for tissue analysis (not populated). |
| AGE | Number | 2 | Age of individual organisms collected for tissue analysis (not populated). |
| PCTLIPID | Number | 8 | Percent lipids (%). |
| EXSAMPID | Text | 15 | Query Manager field (not populated). |
| MESL_LIBNO | Text | 50 | MESL - library number. |
| MESL_STATIONID | Text | 50 | MESL - Station ID (retains the Station ID code as it appears in the original datafiles and/or reports). |
| MESL_COMMENT | Text | 255 | MESL - comments. |
| ptbl - STATION | | | |
| SITEID | Text | 4 | Site ID code (from Query Manager). |
| STUDYID | Text | 2 | Study ID code. |
| STATIONID | Text | 6 | Station ID code (this is the MESL_STATIONID, unless it exceeded 6 characters, then the QM_STATIONID was substituted). |

Table 1. Detailed Description of Database Components (asterisks indicate changes from the Phase I database design).

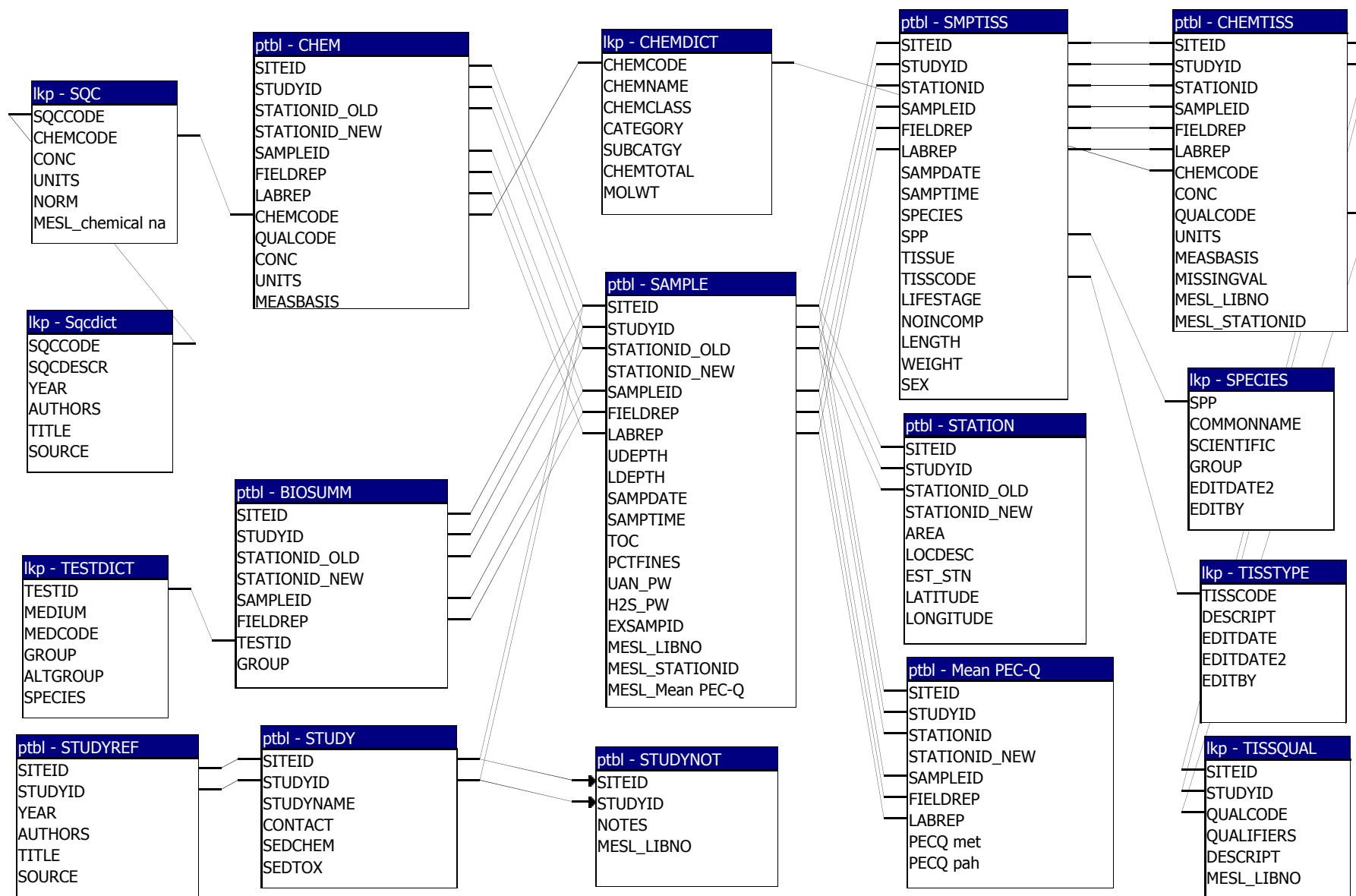
| TABLE NAME / Field Name | Data Type | Field Size | TABLE DESCRIPTION / Field Description |
|-------------------------------|-----------|------------|---|
| ptbl - STATION (cont.) | | | |
| AREA | Text | 50 | Waterbody (corresponds with 'DB_AREA' theme in GIS projects). |
| LOCDESC | Text | 50 | Reach (corresponds with 'Location Description' theme in GIS projects). |
| EST_STN | Text | 50 | Code indicating how the geographic coordinates were obtained (R = reported; P = plotted in GIS based on a map from the report; E = estimated using site descriptions from report; U = unknown). |
| LATITUDE | Number | 8 | Geographical coordinates (decimal degrees). |
| LONGITUDE | Number | 8 | Geographical coordinates (decimal degrees). |
| X-coord | Text | 50 | Geographical coordinates (UTM Zone 15 NAD83 datum). |
| Y-coord | Text | 50 | Geographical coordinates (UTM Zone 15 NAD83 datum). |
| MESL_LIBNO | Text | 50 | MESL - library number. |
| MESL_EST_STN | Text | 50 | MESL - description of how the geographic coordinates were obtained.* |
| MESL_CORELENGTH | Text | 50 | MESL - core length (units are in meters). Note that this field has only been populated when the information has been readily available (electronic format). |
| MESL_Habitat class | Text | 50 | MESL - relevant to REMAP studies only (STUDYID 04 & 06). Codes: 1 = Shallow area; 2 = Channel; 3 = Reservoir. |
| MESL_LOCDESC2 | Text | 50 | MESL - additional station location descriptions. |
| MESL_COMMENTS | Text | 150 | MESL - comments. |
| ptbl - STUDY | | | |
| | NA | NA | Study names and the types of data associated with each study. |
| SITEID | Text | 4 | Site ID code (from Query Manager). |
| STUDYID | Text | 2 | Study ID code. |
| STUDYNAME | Text | 40 | Study name. |
| CONTACT | Text | 40 | Contact person/agency. |
| SEDCHM | Yes/No | 1 | Indicates if the study has surficial sediment chemistry data incorporated in the database. |
| SEDTOX | Yes/No | 1 | Indicates if the study has sediment toxicity data incorporated in the database. |
| SUBSURF | Yes/No | 1 | Indicates if the study has sub-surface sediment chemistry data incorporated in the database. |
| LABACCUM | Yes/No | 1 | Indicates if the study has bioaccumulation test data incorporated in the database. |

Table 1. Detailed Description of Database Components (asterisks indicate changes from the Phase I database design).

| TABLE NAME / Field Name | Data Type | Field Size | TABLE DESCRIPTION / Field Description |
|-----------------------------|-----------|------------|--|
| ptbl - STUDY (cont.) | | | |
| TISSCHEM | Yes/No | 1 | Indicates if the study has tissue chemistry data incorporated in the database. |
| Location/Sampling Year | Text | 40 | Location and sampling year. |
| MESL_LIBNO | Text | 50 | MESL - library number. |
| ptbl - STUDYNOT | | | |
| | NA | NA | Study notes. |
| SITEID | Text | 4 | Site ID code (from Query Manager). |
| STUDYID | Text | 2 | Study ID code. |
| NOTES | Memo | - | Notes. |
| MESL_LIBNO | Text | 50 | MESL - library number. |
| ptbl - STUDYREF | | | |
| | NA | NA | Bibliographic references for each study. |
| SITEID | Text | 4 | Site ID code (from Query Manager). |
| STUDYID | Text | 2 | Study ID code. |
| YEAR | Text | 4 | Publish year for report. |
| AUTHORS | Text | 160 | Authors of the report. |
| TITLE | Text | 160 | Title of the report. |
| SOURCE | Text | 160 | Source (locations). |
| STUDYCOMM | Text | 160 | Comments. |
| MESL_LIBNO | Text | 50 | MESL - library number. |

¹Note that the following fields have been removed from this version of the database: MESL_casno and MESL_update.

Figure 1. Diagram showing the relationships between database components.



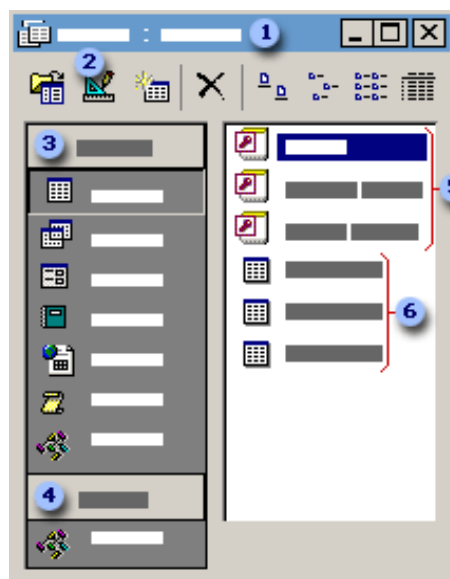
Appendix 1. Instructions for Creating Select Queries in Microsoft Access

A select query retrieves data from one or more tables by using specified criteria and then displays it in the desired order (e.g., ascending order). The following instructions outline the basic steps required to create a select query in query Design View.

Step 1. Under **Objects** (on the left-hand side of the DATABASE window), click **Queries**.

The database window looks like this:

The Database window is the command center of your Access file. From here, you can create and use any object in your Access database or Access project.



1. The title bar of the Database window shows the name and file format of the database.
2. On the Database window toolbar, use the Open button to work with existing objects, use the Design button to modify existing objects, or use the New button to create new objects.
3. Under Objects, click one of the object types, such as Tables or Forms, to show the list of objects of that type.
4. A list of groups of database objects appears under Groups. You can add objects of different types to a group, which consists of shortcuts to the database objects that belong to it.
5. You can use the new object shortcuts at the top of the object list to create new database objects.
6. The list of database objects changes according to which object type you have clicked under Objects.

Step 2. Click **New** (top of the DATABASE window).

Step 3. In the NEW QUERY window select **Design View**, then **OK**.

Step 4. Select the data you want to work with by adding the tables or queries that contain the data of interest.

Step 5. The query is completed by filling in the design grid:

The design grid looks like this:

1. Add or remove tables, queries, and fields

2. Calculate amounts

3. Limit results using criteria

4. Sort records

Step 5a. To add a field to the design grid, drag the field from the field list to a column in the design grid (**1.**), or double-click the field name in the field list (to remove a field from the design grid, click the column selector to highlight the column, and then press the **DELETE** key);

Step 5b. To sort records in the query results, click in the **Sort** cell (**4.**) for the field you want to sort, click the arrow, and then select a sort order (e.g., sort Chemical name in ascending order);

1. If you specify a sort order for more than one field, Microsoft Access sorts the leftmost field first, so you should arrange the fields you want to sort from left to right in the design grid.

2. Sort by ascending or descending order, or remove a sort.

Step 5c. To limit the records that you see in the query's results, specify criteria in the **Criteria** row (**3.**) for one or more fields (e.g., to include only samples collected in Lac de Gras enter “*Lac de Gras*”); and,

| | | |
|-----------|-------------------------------------|--------------------------|
| Field: | LastName | OrderDate |
| Table: | Employees | Orders |
| Total: | Group By | Where |
| Sort: | | |
| Show: | <input checked="" type="checkbox"/> | <input type="checkbox"/> |
| Criteria: | Between #6/1/01# And #6/15/01# | |

1. To limit the records in the query's results, enter criteria in one or more fields.

2. Use the Or row for alternative criteria in the same field.

3. Enter criteria for different fields. For example, for orders between 6/1/01 and 6/15/01 ...

4. ... calculate total order amounts, but display only those that are more than \$100,000.

| | | | |
|-----------|--------------------------------|-----|--------------------------------|
| Criteria: | Between #6/1/01# And #6/15/01# | or: | Between #6/1/01# And #6/15/01# |
|-----------|--------------------------------|-----|--------------------------------|

| | | | |
|-----------|-------------------------------------|--------------------------------|-------------------------------------|
| Field: | LastName | OrderDate | Subtotal |
| Table: | Employees | Orders | Order Subtotals |
| Total: | Group By | Where | Sum |
| Sort: | | | Descending |
| Show: | <input checked="" type="checkbox"/> | <input type="checkbox"/> | <input checked="" type="checkbox"/> |
| Criteria: | | Between #6/1/01# And #6/15/01# | > 100000 |

Step 5d. To perform calculations on the values in a field, click **Totals** on the tool bar (Sum icon; or select **Totals** from the **View** menu) to display the **Total** row in the design grid, then select a function by using the drop-down list [e.g., sum, average, etc.(**2.**)].

| | | |
|--------|-------------------------------------|----------|
| Field: | OrderID | Subtotal |
| Total: | Count | Sum |
| | | Group By |
| | <input checked="" type="checkbox"/> | Sum |
| | | Avg |

1. Use an aggregate function, such as Sum or Avg, to calculate one amount for all the records in each field in the design grid.

2. Use Group By to calculate separate amounts for groups of records in a field.

| | | |
|--------|------------|----------|
| Field: | EmployeeID | Subtotal |
| Total: | Group By | Sum |

Step 6. Run the query by clicking the **Run** button (exclamation point icon) on the toolbar.