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STL STANDARD OPERATING PROCEDURE

TITLE: ACID DIGESTION OF AQUEOUS SAMPLES BY SW846 AND MCAWW 200 SERIES METHODS

(SUPERSEDES: REVISION 0)

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1. **SCOPE AND APPLICATION**

1.1. This procedure describes the preparation of aqueous samples for the analysis of certain metals by Graphite Furnace Atomic Absorption (GFAA), Flame Atomic Absorption (FLAA) and Inductively Coupled Plasma Atomic Emission Spectroscopy (ICP) using the MCAWW 200 series methods (NPDES) and SW846 Methods 3005A, 3010A, 3020A and 7060A/7740 (RCRA).

1.2. The applicability of each of these preparation protocols to specific analytes is detailed in Tables I and II (Appendix A) and the applicable determinative methods are illustrated by Figures 6 and 7 (Section 17). Additional elements may be analyzed following digestion by these protocols provided that the method performance criteria specified in Section 13.0 of this SOP are met.

1.3. This SOP provides procedures applicable to the preparation of dissolved, suspended, total recoverable and total elements in ground water, aqueous samples, solids, sludges, wastes, sediments, air sampling media, biological tissue and leachates/extracts.

1.4. SW-846 Method 3005A is used to prepare surface and groundwater samples for total recoverable and dissolved metals determination by FLAA, ICP and GFAA (antimony only).

1.5. MCAWW Method 200.7 Section 9.4 is used to prepare surface water, domestic and industrial waste samples for total recoverable and dissolved metals determination by ICP.

1.6. SW-846 Method 3010A is used to prepare aqueous samples, EP and mobility-procedure extracts, and wastes that contain suspended solids for total metals analysis by FLAA or ICP.

1.7. MCAWW Method 200.7 Section 9.3 is used to prepare surface water and wastes that contain suspended solids for total metals analysis by ICP.

1.8. SW-846 Method 3020A is used to prepare aqueous samples, EP and mobility-procedure extracts, and wastes that contain suspended solids for total metals by GFAA.

1.9. MCAWW Method 200.0 Section 4.1.3 is used to prepare surface water and wastes that contain suspended solids for total metals analysis by GFAA.
1.10. MCAWW Method 200.0 Section 4.1.4 is used to surface water, domestic and industrial waste samples for total recoverable and dissolved metals determination by GFAA.

1.11. SW-846 Methods 7060A and 7740, respectively, contain the procedure for the preparation of aqueous samples for arsenic and selenium.

1.12. MCAWW Methods 206.2 and 270.2, respectively, contain the procedure for the preparation of aqueous samples for arsenic and selenium.

1.13. All matrices require digestion prior to analysis with the exception of analyses for dissolved metals in filtered and acidified aqueous samples. Although digestion is not specifically required by the method, some clients and regulators do require digestion of dissolved samples and this must be clarified before project initiation.

2. SUMMARY OF METHOD

2.1. Method 3005A / Method 200.7 Section 9.4 - Preparation for Total Recoverable or Dissolved Metals Analysis by FLAA or ICP Spectroscopy

A representative aliquot of sample is heated with nitric and hydrochloric acids and substantially reduced in volume. The digestate is filtered (if necessary) and diluted to volume.

2.2. Method 3010A / Method 200.7 Section 9.3 - Preparation for Total Metals Analysis by FLAA or ICP Spectroscopy

A representative aliquot of sample is refluxed with nitric acid. This step is repeated until the digestate is light in color or until its color has stabilized. After the digestate has been reduced to a low volume, it is refluxed with hydrochloric acid, filtered (if necessary) and brought up to volume.

2.3. Method 3020A / Method 200.0 Section 4.1.3 - Preparation for Total Metals for Analysis by GFAA Spectroscopy

A representative aliquot of sample is refluxed with nitric acid. This step is repeated until the digestate is light in color or until its color has stabilized. After the digestate has been reduced to a low volume, it is cooled, filtered (if necessary) and brought up to volume.
2.4. Methods 7060A/206.2 and Methods 7740/270.2 - Preparation for Arsenic/Selenium Analysis by GFAA

A representative aliquot of sample is heated with nitric acid and peroxide until the digestion is complete or until the volume is reduced by one-half. The sample is cooled, filtered (if necessary) and brought up to volume.

2.5. Method 200.0 Section 4.1.4 - Total Recoverable GFAA Preparation (NPDES)

A representative aliquot of sample is heated with nitric acid until the volume is reduced to 15 - 20 mL. The sample is cooled, filtered (if necessary) and brought up to volume.

3. DEFINITIONS

Additional definitions of terms used in this SOP may be found in the glossary of the QAMP.

3.1. Dissolved Metals: Those elements which pass through a 0.45 um membrane. (Sample is acidified after filtration).

3.2. Suspended Metals: Those elements which are retained by a 0.45 um membrane.

3.3. Total Metals: The concentration determined on an unfiltered sample following digestion.

3.4. Total Recoverable Metals: The concentration determined on an unfiltered sample following treatment with hot, dilute mineral acid.

4. INTERFERENCES

4.1. There are numerous routes by which samples may become contaminated. Potential sources of trace metals contamination include: metallic or metal-containing labware (e.g., talc gloves which contain high levels of zinc), containers, impure reagents, dirty glassware, improper sample transfers, dirty work areas, atmospheric inputs such as dirt and dust, etc. Be aware of potential sources of contamination and take appropriate measures to minimize or avoid them.

4.2. The entire work area, including the bench top and fume hood, should be thoroughly cleaned on a routine schedule in order to minimize the potential for environmental contamination. Refer to Appendix D for additional contamination control guidelines.
4.3. Boron and silica from the glassware will migrate into the sample solution during and following sample processing. For critical low level determinations of boron and silica, only quartz and/or plastic labware should be used.

4.4. Physical interference effects may contribute to inaccuracies in the determinations of trace elements. Oils, solvents and other matrices may not be digested using these methods if they are not soluble with acids. If physical interferences are present, they should be documented.

4.5. Visual interferences or anomalies (such as foaming, emulsions, precipitates, etc.) must be documented.

4.6. Allowing samples to boil or go dry during digestion may result in the loss of volatile metals. If this occurs the sample must be reprepared. Antimony is easily lost by volatilization from hydrochloric acid media.

4.7. Precipitation of silver chloride (AgCl) may occur when chloride ions and high concentrations of silver (i.e., greater than 1 mg/L) are present in the sample.

4.8. Specific analytical interferences are discussed in each of the determinative methods.

5. **SAFETY**

5.1. Procedures shall be carried out in a manner that protects the health and safety of all STL associates.

5.2. Eye protection that satisfies ANSI Z87.1 (as per the Chemical Hygiene Plan), laboratory coat, and appropriate gloves must be worn while samples, standards, solvents, and reagents are being handled. Disposable gloves that have been contaminated will be removed and discarded; other gloves will be cleaned immediately.

5.3. The health and safety hazards of many of the chemicals used in this procedure have not been fully defined. Additional health and safety information can be obtained from the Material Safety Data Sheets (MSDS) maintained in the laboratory. The following specific hazards are known:

5.3.1. The following materials are known to be **corrosive**:

    hydrochloric acid and nitric acid.
5.3.2. The following materials are known to be oxidizing agents:

- nitric acid and hydrogen peroxide.

5.3.3. All sample digestions, including cooling of digestates, must be carried out in a fume hood.

5.4. The acidification of samples containing reactive materials may result in the release of toxic gases, such as cyanides or sulfides. Acidification of samples should be done in a fume hood. The analyst should also be aware of the potential for a vigorous reaction.

5.5. Exposure to chemicals must be maintained as low as reasonably achievable. Therefore, unless they are known to be non-hazardous, all samples should be opened, transferred and prepared in a fume hood, or under other means of mechanical ventilation. Solvent and waste containers will be kept closed unless transfers are being made.

5.6. The preparation of standards and reagents will be conducted in a fume hood with the sash closed as far as the operation will permit or under other means of mechanical ventilation.

5.7. All work must be stopped in the event of a known or potential compromise to the health and safety of a STL associate. The situation must be reported immediately to a laboratory supervisor.

5.8. Always carry bulk concentrated acid bottles in appropriate impact proof containers.

5.9. Acid/peroxide spills must be neutralized immediately, flushed with water and cleaned up using appropriate spill kits.

5.10. Discard chipped or broken beakers to prevent injury. Chipped glassware may be fire polished as an alternative to disposal.

5.11. Any and all accidents and spills must be reported to the lab supervisor or EH&S coordinator.

6. **EQUIPMENT AND SUPPLIES**

6.1. Hot plate, digestion block or other adjustable heating source capable of maintaining a temperature of 90 - 95°C.
6.2. Thermometer that covers a temperature range of 0-200°C.

6.3. Griffin beakers of assorted sizes or equivalent.

6.4. Watch glasses, ribbed or equivalent.

6.5. Whatman No. 41 filter paper or equivalent.

6.6. Funnels or equivalent filtration apparatus.

6.7. Centrifugation equipment (if desired method of removing particulates is centrifugation).

6.8. Graduated cylinder or equivalent capable of measuring 50 mL within 3% accuracy.

6.9. Analytical balance capable of accurately weighing to the nearest 0.01 grams.

6.10. Repipetors or suitable reagent dispensers.

6.11. Calibrated automatic pipettes with corresponding pipet tips or Class A glass volumetric pipettes.


6.13. pH indicator strips (pH range 0 - 6).


7. REAGENTS AND STANDARDS

7.1. Reagent water must be produced by a Millipore DI system or equivalent. Reagent water must be free of the analytes of interest as demonstrated through the analysis of method blanks as defined in the determinative SOPs.

7.2. Laboratory Control Sample (LCS) and matrix spike (MS) solutions are purchased as custom STL solutions. All standards must be stored in FEP fluorocarbon or previously unused polyethylene or polypropylene bottles. Stock standard solutions must be replaced prior to the expiration date provided by the manufacturer. If no expiration date is provided, the stock solutions may be used for up to one year and must be replaced sooner if verification from an independent source indicates a problem.
7.3. Working ICP LCS/MS spike solution: The ICP LCS/MS working spike solution is provided directly by the vendor, no further standard preparation is necessary.

7.4. Working GFAA LCS/MS spike solution: Prepare the GFAA working LCS spike solution by diluting the custom stock solution (7.2) 100x. The working spike solution must be prepared in a matrix of 5% HNO₃. This acid (5 mL of concentrated HNO₃ per 100 mL) must be added to the volumetric flask before the addition of the stock standard aliquot. The working GFAA LCS solution must be made fresh every three months.

7.5. The TCLP MS working spike solution is provided directly by the vendor, no further standard preparation is necessary.

7.6. The LCS and MS samples must contain all the elements designated for analysis in each batch of samples. If a non-routine element is required that is not contained in the custom STL solution, the individual facility must purchase a solution from the designated vendor that will cover the additional analyte(s) of interest and provide for a final spike concentration that is appropriate to the determinative method.

7.7. Aqueous laboratory control samples (LCSW) and matrix spike samples are prepared as described in Sections 9.5 and 9.6. Refer to Tables II and III (Appendix A) for details regarding the stock, working standard and final digestate spike concentrations for ICP and GFAA LCS and matrix spike preparations.

7.8. Nitric acid (HNO₃), concentrated, trace metal grade or better.

7.9. Nitric acid, 1:1 - dilute concentrated HNO₃ with an equal volume of reagent water.

Note: When preparing diluted acids always add acid to water. If the water is added to the acid a violent reaction may occur.

7.10. Hydrochloric acid (HCl), concentrated, trace metal grade or better.

7.11. Hydrochloric acid, 1:1 - dilute concentrated HCl with an equal volume of reagent water.

Note: When preparing diluted acids always add acid to water. If the water is added to the acid a violent reaction may occur.

7.12. 30% Hydrogen peroxide (H₂O₂), reagent grade.
8. **SAMPLE COLLECTION, PRESERVATION AND STORAGE**

8.1. Sample holding time for metals included under the scope of this SOP is 180 days from the date of collection to the date of analysis.

8.2. Aqueous samples are preserved with nitric acid to a pH of <2 and may be stored in either plastic or glass. If boron or silica are to be determined, plastic containers are preferred. Refrigeration is not required. Preservation must be verified prior to analysis.

8.3. For dissolved metals analysis, the samples should be filtered through a 0.45 um filter prior to preservation. Filtration must be done in the field or within 24 hours of collection.

**Note:** If a sample being analyzed for dissolved metals is found to contain sediment the analyst should contact their supervisor or group leader. The client should be notified of the problem to decide how to treat the sample.

9. **QUALITY CONTROL**

Table VI (Appendix A) provides a summary of quality control requirements including type, frequency, acceptance criteria and corrective action.

9.1. Initial Demonstration of Capability

Prior to analysis of any analyte using any method contained within this SOP the following requirements must be met:
9.1.1. Method Detection Limit (MDL) - An MDL must be determined for each analyte/matrix prior to the analysis of any samples. The MDL is determined using seven replicates of reagent water, spiked with all the analytes of interest, that have been carried through the entire analytical procedure. MDL’s must be redetermined on an annual basis in accordance with 40 CFR Part 136 Appendix B requirements as detailed in STL QA Policy QA-005. The spike level must be between the calculated MDL and 10X the MDL to be valid. The result of the MDL determination must be below the STL reporting limit.

9.1.2. Initial Demonstration Study - This requires the analysis of four QC check samples. The QC check sample is a well-characterized laboratory generated sample used to monitor method performance, which should contain all the analytes of interest. The results of the initial demonstration study must be acceptable before analysis of samples may begin. The results of the initial demonstration study may be used to extend a method for the analysis of other elements provided all acceptance criteria are met.

9.1.2.1. Four aliquots of the check sample (LCS) are prepared and analyzed using the procedures detailed in this SOP and the determinative SOPs.

9.1.2.2. Calculations and acceptance criteria for QC check samples are given in the determinative SOPs (CORP-MT-0001, CORP-MT-0003).

9.2. Preparation Batch - A group of up to 20 samples that are of the same matrix and are processed together using the same procedures and reagents. The preparation batch must contain a method blank, a LCS and a matrix spike/matrix spike duplicate. In some cases, at client request, it may be appropriate to process a matrix spike and sample duplicate in place of the MS/MSD. If clients specify specific samples for MS/MSD, the batch may contain multiple MS/MSD pairs.

9.3. Sample Count - Laboratory generated QC samples (method blanks, LCS) are not included in the sample count for determining the size of a preparation batch. MS/MSD are not included in the sample count unless there are multiple sets of MS/MSD per batch. In other words, the first MS/MSD are not counted; all additional MS and MSDs are counted as samples.

9.4. Method Blank (MB) - One method blank must be processed with each preparation batch. The method blank consists of reagent water containing all reagents specific to the method that is carried through the entire analytical procedure, including preparation and analysis. The method blank is used to identify any system and process interferences or contamination of the analytical system that may lead to the reporting of elevated analyte concentrations or false positive data. Criteria for the
acceptance of blanks are contained within the individual analytical method SOP’s. If the method blank does not meet the criteria contained within the analytical method SOPs, the blank and all associated samples in the batch must be redigested.

9.4.1. Aqueous method blanks are prepared by taking 50 mL or 50 g of reagent water through the appropriate procedure as described in Section 11.

9.4.2. TCLP method blanks are prepared by taking 50 mL or 50 g of leachate fluid through the appropriate procedure as described in Section 11.

9.5. Laboratory Control Sample (LCS) - One aqueous LCS must be processed with each preparation batch. The LCS must contain all analytes of interest and must be carried through the entire analytical procedure. The LCS is used to monitor the accuracy of the analytical process. On-going monitoring of the LCS results provides evidence that the laboratory is performing the method within acceptable accuracy and precision guidelines. Criteria for the acceptance of LCS results are contained within the individual analytical method SOP’s. Corrective action when LCS results fail to meet control limits will be repreparation and reanalysis of the batch. Refer to Section 7.3 and 7.4 for instructions on preparation of the aqueous LCS spike solution.

9.5.1. The aqueous LCS is prepared by spiking a 50 mL aliquot of reagent water with 0.5 mL of the working LCS/MS spike solution (7.3 or 7.4). The LCS is then processed through the appropriate procedure as described in Section 11.

9.6. Matrix Spike/Matrix Spike Duplicate (MS/MSD) - One MS/MSD pair must be processed for each preparation batch. A matrix spike (MS) is a field sample to which known concentrations of target analytes have been added. A matrix spike duplicate (MSD) is a second aliquot of the same sample (spiked identically as the MS) prepared and analyzed along with the sample and matrix spike. Some client specific data quality objectives (DQO’s) may require the use of sample duplicates in place of or in addition to MS/MSD’s. The MS/MSD results are used to determine the effect of a matrix on the precision and accuracy of the analytical process. Samples identified as field blanks cannot be used for MS/MSD analysis. If any analyte recovery or RPD falls outside the acceptance range, the recovery of that analyte must be in control for the LCS. If the recovery of the LCS is outside limits, corrective action must be taken. Corrective action will include repreparation and reanalysis of the batch. Corrective action when MS results fail to meet control limits does not include repreparation of samples unless the results indicate that a spiking error may have occurred.

9.6.1. The aqueous matrix spike sample is prepared by spiking a 50 mL aliquot of a sample with 0.5 mL of the working LCS/MS spike solution (7.3 or 7.4). The matrix spike sample is then processed as described in Section 11.
9.6.2. The TCLP matrix spike sample is prepared by spiking a 50 mL aliquot of a leachate with 0.5 mL of the working TCLP spike solution (7.5). The matrix spike sample is then processed as described in Section 11.

NOTE: The TCLP matrix spike must be added prior to preservation of the leachate.

9.6.3. If insufficient sample is available to process a MS/MSD, then a second LCS must be processed. The LCS pair is then evaluated according to the MS/MSD criteria.

9.7. Quality Assurance Summaries - Certain clients may require specific project or program QC which may supersede the SOP requirements. Quality Assurance Summaries (QAS) should be developed to address these requirements.

10. CALIBRATION AND STANDARDIZATION

10.1. Hotplate temperature must be verified daily for each hotplate used and must be recorded on either the metals preparation log or in a hotplate temperature logbook. The hotplate temperature should be verified by measuring the temperature of a beaker of reagent water placed on each hotplate.

11. PROCEDURE

11.1. One time procedural variations are allowed only if deemed necessary in the professional judgment of supervision to accommodate variation in sample matrix, radioactivity, chemistry, sample size, or other parameters. Any variation in procedure shall be completely documented using a Nonconformance Memo and is approved by a Technical Specialist and QA Manager. If contractually required, the client shall be notified. The Nonconformance Memo shall be filed in the project file.

11.2. Any unauthorized deviations from this procedure must also be documented as a nonconformance, with a cause and corrective action described.

11.3. All preparation procedures must be carried out in a properly functioning hood.

11.4. All samples are to be checked out of sample control with the chain of custody documentation filled out completely.

11.5. Proper sample identification is extremely important in any preparation procedure. Labeling of beakers and bottles must be done in a manner to ensure connection with the proper sample.
11.6. Samples are typically logged in as either waters or soils. Wastes such as organic liquids or sludges and tissues (animal/vegetable) are usually logged in with solid test codes. When initiating prep examine the sample to see if the sample matches the matrix designation. If the sample is logged in as aqueous but it appears more like a waste (biphasic, sludge like, organic liquid, lots of sediment etc.) contact the lab supervisor or project manager for further instructions. In some cases it may be more appropriate to process these samples as solids.

11.7. If possible prepare all the samples of a project at the same time to minimize the QC required and streamline the flow of the project through the lab and reporting group.

11.8. In most cases, both AA and ICP digests are required on each sample. It is recommended that both aliquots be measured out and processed at the same time.

11.9. Guidelines are provided in the appendices on procedures to minimize contamination of samples and standards.

11.10. The following procedure must be followed for all aqueous sample preparations:

11.10.1. Measure and record sample pH with pH paper on a separate aliquot of sample.

  **Note:** If the sample pH is > 2 pH units, the client must be notified of the anomaly.

  **Note:** If sample pH has already been verified and documented in sample receipt this step may be omitted.

11.10.2. Mix sample by shaking the container.

11.10.3. Measure and transfer 50 mL or 50 g of the sample into a beaker.

  **Note:** This SOP allows for samples to be weighed instead of measured volumetrically (See Section 17.1.1.2).

11.10.4. Measure two extra aliquots of sample selected for the MS/MSD analysis. Spike each aliquot with 0.5 mL of spiking solution (7.3 or 7.4).

11.10.5. Measure and transfer 50 mL of reagent water into a beaker for the method blank.

11.10.6. Measure and transfer 50 mL of reagent water into a beaker for the LCS and add 0.5 mL of spiking solution (7.3 or 7.4).
11.11. Proceed to the appropriate Section for the desired method as follows:

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<tr>
<td>Method 200.0 Section 4.1.4</td>
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</table>

11.12. **Method 3005A / Method 200.7 Section 9.4 - Preparation for Total Recoverable or Dissolved Metals Analysis by FLAA or ICP (See Figures 1, 6 and 7)**

11.12.1. To the sample beaker, add 1 mL of concentrated HNO₃ and 2.5 mL of concentrated HCl.


11.12.3. Heat at 90 - 95°C until volume is reduced to between 15 and 20 mL.

**NOTE: DO NOT ALLOW SAMPLE TO BOIL OR GO DRY.** Doing so will result in the loss of analyte and the sample must be reprepared.

11.12.4. Cool the beaker in a fume hood.

11.12.5. Wash down beaker walls and watch glass with reagent water.

11.12.6. Filter sample, if insoluble materials are present, though Whatman 41 filter paper that has been pre-rinsed with dilute nitric acid.

**Note:** If any samples in a preparation batch are filtered, the method blank and LCS associated with that batch must also be filtered.

**Note:** In place of filtering, the samples, after dilution and mixing, may be centrifuged or allowed to settle by gravity overnight to remove insoluble material.
11.12.7. Rinse beaker and filter paper with reagent water to ensure complete sample transfer.

11.12.8. Add 1.5 mL concentrated HNO₃ and adjust the final volume/mass to 50 mL or 50 g with reagent water. The sample is now ready for analysis.

11.13. Method 3010A / Method 200.7 Section 9.3 - Preparation for Total Metals Analysis by FLAA or ICP Spectroscopy (See Figures 2, 6 and 7)

11.13.1. To the sample beaker, add 1.5 mL of concentrated HNO₃.

11.13.2. Cover with ribbed watch glass.

11.13.3. Place beaker on hotplate (90-95 °C) and evaporate to low volume of 5 - 10 mL while ensuring that no portion of the bottom of the beaker is allowed to go dry.

**NOTE: DO NOT ALLOW SAMPLE TO BOIL OR GO DRY.** Doing so will result in the loss of analyte and the sample must be reprepared.

11.13.4. Cool the beaker in a fume hood.

11.13.5. Add another 1.5 mL portion of concentrated HNO₃ and re-cover the beaker.

11.13.6. Continue refluxing until the digestion is complete.

**Note:** Digestion is complete when the digestate is light in color or does not change in appearance. For most samples the addition of two nitric acid aliquots is sufficient, additional aliquots of nitric acid may be added if necessary.

11.13.7. Evaporate to low volume of 5 - 10 mL while ensuring that no portion of the bottom of the beaker is allowed to go dry.

11.13.8. Cool the beaker in a fume hood.

11.13.9. Add 5 mL of 1:1 HCl.

11.13.10. Cover and reflux for an additional 15 minutes to dissolve precipitate or residue.
11.13.11. Wash down beaker walls and watch glass with reagent water.

11.13.12. Filter sample, if insoluble materials are present, through Whatman 41 filter paper that has been pre-rinsed with dilute nitric acid (1%).

**Note:** If any samples in the QC batch are filtered the method blank and LCS associated with that batch must also be filtered.

**Note:** In place of filtering, the samples, after dilution and mixing, may be centrifuged or allowed to settle by gravity overnight to remove insoluble material.

11.13.13. Rinse beaker and filter paper with reagent water to ensure complete sample transfer.

11.13.14. Adjust final volume/mass to 50 mL or 50 g with reagent water. The sample is now ready for analysis.

11.14. **Method 3020A / Method 200.0 Section 4.1.3 - Preparation for Total Metals Analysis by GFAA** (See Figures 3, 6 and 7)

11.14.1. To the sample beaker, add 1.5 mL of concentrated HNO₃.


11.14.3. Place beaker on hotplate (90-95 °C) and evaporate to low volume of 5 - 10 mL while ensuring that no portion of the bottom of the beaker is allowed to go dry.

**NOTE: DO NOT ALLOW SAMPLE TO BOIL OR GO DRY.** Doing so will result in the loss of analyte and the sample must be reprepared.


11.14.5. Add another 1.5 mL portion of concentrated HNO₃.


11.14.7. Continue refluxing and adding acid as necessary until digestion is complete.
Note: Digestion is complete when the digestate is light in color or does not change in appearance. For most samples the addition of two nitric acid aliquots is sufficient, additional aliquots of nitric acid may be added if necessary.

11.14.8. After the digestion is complete evaporate to low volume of 5 - 10 mL while ensuring that no portion of the bottom of the beaker is allowed to go dry.


11.14.10. Add 10 mL of reagent water and mix sample.

11.14.11. Heat sample for 10 to 15 minutes more to dissolve any residue.


11.14.14. Filter sample, if insoluble materials are present, though Whatman 41 filter paper that has been pre-rinsed with dilute nitric acid (1%).

Note: If any samples in the QC batch are filtered the method blank and LCS associated with that batch must also be filtered.

Note: In place of filtering, the samples, after dilution and mixing, may be centrifuged or allowed to settle by gravity overnight to remove insoluble material.

11.14.15. Rinse beaker and filter paper with reagent water to ensure complete sample transfer.

11.14.16. Adjust final volume to 50 mL with reagent water. The sample is now ready for analysis.

11.15. Method 7060A/7740 and Method 206.2/270.2 - Preparation for Arsenic and Selenium Analysis by GFAA (See Figures 4, 6 and 7)

11.15.1. To the sample beaker, add 1 mL of 30 % H₂O₂ and 0.5 mL of concentrated HNO₃.

11.15.2. Heat, until the digestion is complete, at 90 - 95°C or until the volume has been reduced to slightly less than 25 mL.
11.15.3. Cool beaker.

11.15.4. Filter sample, if insoluble materials are present, through Whatman 41 filter paper that has been pre-rinsed with dilute nitric acid.

**Note:** If any samples in the QC batch are filtered the method blank and LCS associated with that batch must also be filtered.

**Note:** In place of filtering, the samples, after dilution and mixing, may be centrifuged or allowed to settle by gravity overnight to remove insoluble material.

11.15.5. Rinse beaker and filter paper with reagent water to ensure complete sample transfer.

11.15.6. Adjust final volume to 50 mL with reagent water. The sample is now ready for analysis.

11.16. **Method 200.0 Section 4.1.4 - Preparation for Total Recoverable GFAA Analyses. (See Figures 5 and 7)**

11.16.1. To the sample beaker, add 0.5 mL of concentrated HNO₃.

11.16.2. Heat, until the digestion is complete, at 90 - 95°C or until the volume has been reduced to 15 - 20 mL.

11.16.3. Cool beaker.

11.16.4. Filter sample, if insoluble materials are present, through Whatman 41 filter paper that has been pre-rinsed with dilute nitric acid.

**Note:** If any samples in the QC batch are filtered the method blank and LCS associated with that batch must also be filtered.

**Note:** In place of filtering, the samples, after dilution and mixing, may be centrifuged or allowed to settle by gravity overnight to remove insoluble material.

11.16.5. Rinse beaker and filter paper with reagent water to ensure complete sample transfer.
11.16.6. Adjust final volume to 50 mL with reagent water. The sample is now ready for analysis.

12. DATA ANALYSIS AND CALCULATIONS

Not Applicable.

13. METHOD PERFORMANCE

13.1. Method performance is determined by the analysis of matrix spike and matrix spike duplicate samples as well as method blanks and laboratory control samples. In general, the matrix spike recovery should fall within +/- 20% and the matrix spike duplicates should compare within 20% RPD. Method blanks must meet the criteria specified in determinative SOPs. The laboratory control samples should recover within 20% of the true value until in house control limits are established. Acceptance criteria are given in the determinative SOPs.

13.2. The initial demonstration study as detailed in Section 9.1.2 must be acceptable before the analysis of field samples under this SOP may begin. The results of the initial demonstration study may be used to extend a method for the analysis of other elements provided all acceptance criteria are met.

13.3. Training Qualification:

The group/team leader has the responsibility to ensure that this procedure is performed by an associate who has been properly trained in its use and has the required experience.

14. POLLUTION PREVENTION

14.1. This method allows for the proportional reduction of sample and reagent volumes to decrease waste generation.

15. WASTE MANAGEMENT

15.1. Waste generated in the procedure must be segregated and disposed according to the facility hazardous waste procedures. The facility EH & S coordinator should be contacted if additional information is required.

15.2. Standards should be purchased and prepared in volumes consistent with laboratory use to minimize the volume of expired standards to be disposed.
16. **REFERENCES**


16.5. QA-003, STL QC Program.

16.6. QA-004, Rounding and Significant Figures.

16.7. QA-005, Method Detection Limits.

17. **MISCELLANEOUS (TABLES, APPENDICES, ETC. . . )**

17.1. Modifications/Interpretations from reference methods.

17.1.1. Modifications applicable to SW-846 reference methods.

17.1.1.1. Chapter 1 of SW-846 states that the method blank should not contain any analyte of interest at or above the MDL. This SOP states that the method blank must not contain any analyte of interest at or above the reporting limit. Common lab contaminants are allowed up to two times the reporting limit in the blank following consultation with the client.

17.1.1.2. This SOP allows for aqueous samples to be weighed instead of measured volumetrically. This assumes the density of the sample is close to 1.0 g/mL. Samples with large amounts of sediment or suspended solids, sludges, non-aqueous liquids must be processed volumetrically. Weighing samples directly into the digestion vessel minimizes the potential for cross contamination, offers improved accuracy over the use of graduated cylinders (comparable
to volumetric flask accuracy), uses less glassware and is more efficient.

17.1.1.3. The referenced methods as well as Table 3-1 of SW-846 refer to the use of a 100 mL aliquot for digestion. This SOP requires the use of a 50 mL sample size to reduce waste generation. The use of reduced sample volumes are supported in EPA’s document “Response to Public Comments Background Document, Promulgation of the Second Update to SW-846, Third Edition” dated November 3, 1994. This document stated “...flexibility to alter digestion volumes is addressed and “allowed” by the table (3-1) and is also inherently allowed by specific digestion methods. Table 3-1 is only to be used as guidance when collecting samples...” EMSL-Ci has also taken the stance that “reduction in sample size and appropriate corresponding reduction in sample volume is not considered a significant change in the methodology.” Additionally, in written correspondence from the Office of Solid Waste, Olliver Fordham stated “As a “representative sample” can be assured, scaling causes no loss of precision and accuracy in the analysis.”

17.1.2. Modifications Specific to Method 3005A

17.1.2.1. In order to matrix match the digestate to the ICP calibration standards, Section 11.12.8 requires the addition of 1.5 mL of concentrated nitric acid to the digestate prior to dilution to final volume. This step ensures that bias due to differences in acid matrix will not be a factor in the analytical determination. Since this step is performed post-digestion it does not impact the digestion recoveries. This approach to matrix matching was discussed with Olliver Fordham of OSW who indicated that it was an acceptable practice.

17.1.3. Modifications Specific to Method 3010A

17.1.3.1. Section 11.13.7 of this SOP requires the sample be reduced to a volume of 5 - 10 mL. Section 7.2 of Method 3010A states the volume should be reduced to 3 mL but also states that no portion of the bottom of the beaker should go dry. The SOP required volume is a closer approximation of the volume required to provide an adequate covering of the beaker so as to prevent the loss of critical analytes through volatilization.
17.1.3.2. The scope of 3010A has been expanded to include silver based on comparison studies with 7760A. Method 3010A consistently demonstrated improved accuracy and precision over Method 7760A in the matrices tested (reagent water, surface water and TCLP leachate) up to a concentration of 1 ppm silver.

17.1.4. Modifications Specific to Method 3020A

17.1.4.1. Section 11.14.8 of this SOP requires the sample be reduced to a volume of 5 - 10 mL. Section 7.2 of Method 3010A states the volume should be reduced to 3 mL but also states that no portion of the bottom of the beaker should go dry. The SOP required volume is a closer approximation of the volume required to provide an adequate covering of the beaker so as to prevent the loss of critical analytes through volatilization.

17.1.5. Modifications Specific to Method 7060A/7740

17.1.5.1. Methods 7060A and 7740A incorporate the use of a two step dilution to accommodate the addition of a nickel nitrate modifier. This SOP performs the dilution directly in one step and omits the addition of the modifier. The modifier is added automatically at the instrument by direct injection into the furnace.

17.1.6. Modifications Specific to MCAWW Methods

It was determined by technical review that several of the MCAWW methods were equivalent to the SW-846 methods and therefore were combined under the scope of this SOP as described in Section 11.0. The nature of the differences were deemed insignificant in regards to the amount of acid added and the evaporative volume based on the flexibility allowed by the methods (i.e., add additional acid as required) and the subjective wording of the methods (i.e., evaporate to near dryness vs. an exact volume).

17.2. Modifications from previous SOP

None.

17.3. Facility Specific SOPs

Each facility shall attach a list of facility specific SOPs or approved attachments (if applicable) which are required to implement this SOP or which are used in conjunction with this SOP. If no facility specific SOPs or amendments are to be
attached, a statement must be attached specifying that there are none. Refer to the Appendices for any facility specific information required to support this SOP.

17.4. Documentation and Record Management

The preparation benchsheet should, at a minimum, include the following information:

- Preparation date, analyst name, matrix, prep type (ICP or GFAA), SOP reference.
- Sample ID, initial weight/volume and final weight/volume.
- Standards Documentation (source, lot, prep date, volume added).
- Analyst Signature.
- Reviewer’s Signature and date.
Verify sample pH < 2

Mix sample thoroughly

Aliquot 50 mL or 50 g of sample into beaker

Add 1 mL HNO3 and 2.5 mL HCl

Cover, Heat at 90 - 95 C

Reduce volume to 15 - 20 mL

Cool, filter if necessary

Dilute to 50 mL with reagent water

Proceed with analysis per determinative SOP.
Figure 2. Method 3010A / Method 200.7 Section 9.3 (Section 11.13)

- Verify sample pH < 2
- Mix sample thoroughly
- Aliquot 50 mL or 50 g of sample into beaker
- Add 1.5 mL HNO3
- Cover, Heat at 90 - 95 C
- Reduce volume to 15 - 20 mL
- Cool
- Add 1.5 mL HNO3
- Reflux, add acid as necessary
- Cool
- Add 5 mL 1:1 HCl
- Reflux 15 minutes
- Cool, filter if necessary
- Dilute to 50 mL with reagent water
- Proceed with analysis per determinative SOP.
Verify sample pH < 2
Mix sample thoroughly
Aliquot 50 mL or 50 g of sample into beaker
Add 1.5 mL HNO3
Cover, heat at 90 - 95°C
Reduce volume to 15 - 20 mL

Cool
Add 1.5 mL HNO3
Reflux, add acid as necessary
Cool
Add 10 mL reagent water
Reflux 15 minutes

Cool, filter if necessary
Dilute to 50 mL with reagent water
Proceed with analysis per determinative SOP.
Figure 4. Method 7060A/7740A and Method 206.2/270.2 (Section 11.15)

1. Verify sample pH < 2
2. Mix sample thoroughly
3. Aliquot 50 mL or 50 g of sample into beaker
4. Add 1 mL 30% H2O2 and 0.5 mL HNO3
5. Cover, heat at 90-95°C
6. Reduce volume to 25 mL
7. Cool, filter if necessary
8. Dilute to 50 mL with reagent water
9. Proceed with analysis per determinative SOP.
Figure 5. Method 200.0 Section 4.1.4 (Section 11.16)

1. Verify sample pH < 2
2. Mix sample thoroughly
3. Aliquot 50 mL or 50 g of sample into beaker
4. Add 0.5 mL HNO3
5. Cover, Heat at 90-95 C
6. Reduce volume to 15 - 20 mL
7. Cool, filter if necessary
8. Dilute to 50 mL with reagent water
9. Proceed with analysis per determinative SOP.
Figure 6. Overview of SW846 Aqueous Preparation Methods by Determinative Method

GFAA

3005A
Sb

7060A/7740
As, Se

3020A
Pb, Tl, Cd, Cr, Be, Co, Mo, V

ICP

3005A
TAL list + Mo

3010A
TAL list (except Sb) + Mo

FLAA

3005A
TAL list (except As, Se) + Mo

3010A
TAL list (except Sb, As, Se) + Mo

TAL list: Al, Sb, As, Ba, Be, Cd, Ca, Cr, Co, Cu, Fe, Pb, Mg, Mn, Ni, K, Se, Ag, Na, Tl, V, Zn
Figure 7. Overview of MCAWW Aqueous Preparation Methods By Determinative Technique

GFAA

206.2/270.2
As, Se

200.0 (4.1.3)
Pb, Tl, Sb, Ag, Ba, Be, Cd, Cr, Co, Cu, Mo, Mn, Ni, V

200.0 (4.1.4)
Pb, Tl, Sb, Ag, Ba, Be, Cd, Cr, Co, Cu, Mo, Mn, Ni, V

ICP

200.7 (9.3)
TAL list + B, Si, Mo

200.7 (9.4)
TAL list + B, Si, Mo

TAL list: Al, Sb, As, Ba, Be, Cd, Ca, Cr, Co, Cu, Fe, Pb, Mg, Mn, Ni, K, Se, Ag, Na, Tl, V, Zn
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X - Designates that the preparation method is approved for an element

**Note:** Additional elements may be analyzed following digestion by these protocols provided the method performance criteria specified in Section 13.0 of the SOP are met.
### TABLE II. Approved Preparation Method Analytes - NPDES

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<td>Sodium</td>
<td>Na</td>
<td>7440-23-5</td>
<td>X</td>
<td>X</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Thallium</td>
<td>Tl</td>
<td>7440-28-0</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td></td>
</tr>
<tr>
<td>Vanadium</td>
<td>V</td>
<td>7440-62-2</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td>X</td>
<td></td>
</tr>
<tr>
<td>Zinc</td>
<td>Zn</td>
<td>7440-66-6</td>
<td>X</td>
<td>X</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

X - Designates that the preparation method is approved for an element

**Note:** Additional elements may be analyzed following digestion by these protocols provided the method performance criteria specified in Section 13.0 of the SOP are met.
### TABLE III. ICP and FLAA Matrix Spike and Aqueous Laboratory Control Sample Levels

<table>
<thead>
<tr>
<th>ELEMENT</th>
<th>Working LCS/MS Standard (mg/L)</th>
<th>Aqueous LCS/MS Level * (ug/l)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Aluminum</td>
<td>200</td>
<td>2000</td>
</tr>
<tr>
<td>Antimony</td>
<td>50</td>
<td>500</td>
</tr>
<tr>
<td>Arsenic</td>
<td>200</td>
<td>2000</td>
</tr>
<tr>
<td>Barium</td>
<td>200</td>
<td>2000</td>
</tr>
<tr>
<td>Beryllium</td>
<td>5</td>
<td>50</td>
</tr>
<tr>
<td>Cadmium</td>
<td>5</td>
<td>50</td>
</tr>
<tr>
<td>Calcium</td>
<td>5000</td>
<td>50000</td>
</tr>
<tr>
<td>Chromium</td>
<td>20</td>
<td>200</td>
</tr>
<tr>
<td>Cobalt</td>
<td>50</td>
<td>500</td>
</tr>
<tr>
<td>Copper</td>
<td>25</td>
<td>250</td>
</tr>
<tr>
<td>Iron</td>
<td>100</td>
<td>1000</td>
</tr>
<tr>
<td>Lead</td>
<td>50</td>
<td>500</td>
</tr>
<tr>
<td>Lithium</td>
<td>100</td>
<td>1000</td>
</tr>
<tr>
<td>Magnesium</td>
<td>5000</td>
<td>50000</td>
</tr>
<tr>
<td>Manganese</td>
<td>50</td>
<td>500</td>
</tr>
<tr>
<td>Molybdenum</td>
<td>100</td>
<td>1000</td>
</tr>
<tr>
<td>Nickel</td>
<td>50</td>
<td>500</td>
</tr>
<tr>
<td>Phosphorous</td>
<td>1000</td>
<td>10000</td>
</tr>
<tr>
<td>Potassium</td>
<td>5000</td>
<td>50000</td>
</tr>
<tr>
<td>Selenium</td>
<td>200</td>
<td>2000</td>
</tr>
<tr>
<td>Silver</td>
<td>5</td>
<td>50</td>
</tr>
<tr>
<td>Sodium</td>
<td>5000</td>
<td>50000</td>
</tr>
<tr>
<td>Strontium</td>
<td>100</td>
<td>1000</td>
</tr>
<tr>
<td>Thallium</td>
<td>200</td>
<td>2000</td>
</tr>
<tr>
<td>Vanadium</td>
<td>50</td>
<td>500</td>
</tr>
<tr>
<td>Zinc</td>
<td>50</td>
<td>500</td>
</tr>
<tr>
<td>Boron</td>
<td>100</td>
<td>1000</td>
</tr>
<tr>
<td>Silica</td>
<td>1000</td>
<td>10000</td>
</tr>
<tr>
<td>Tin</td>
<td>200</td>
<td>2000</td>
</tr>
<tr>
<td>Titanium</td>
<td>100</td>
<td>1000</td>
</tr>
</tbody>
</table>

* Levels shown indicate the spike concentration in the final digestate of the aqueous LCS or matrix spike based on the addition of 0.5 mL working spike (7.3) to 50 mL of sample.

### TABLE IV. GFAA Matrix Spike and Aqueous LCS Spike Levels
### TABLE V. TCLP Reporting Limits, Regulatory Limits and Matrix Spike Levels

<table>
<thead>
<tr>
<th>ELEMENT</th>
<th>RL (ug/L)</th>
<th>Regulatory Limit (ug/L)</th>
<th>Spike Level (ug/L)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Arsenic</td>
<td>500</td>
<td>5000</td>
<td>5000</td>
</tr>
<tr>
<td>Barium</td>
<td>10000</td>
<td>100000</td>
<td>50000</td>
</tr>
<tr>
<td>Cadmium</td>
<td>100</td>
<td>1000</td>
<td>1000</td>
</tr>
<tr>
<td>Chromium</td>
<td>500</td>
<td>5000</td>
<td>5000</td>
</tr>
<tr>
<td>Lead</td>
<td>500</td>
<td>5000</td>
<td>5000</td>
</tr>
<tr>
<td>Selenium</td>
<td>250</td>
<td>1000</td>
<td>1000</td>
</tr>
<tr>
<td>Silver</td>
<td>500</td>
<td>5000</td>
<td>1000</td>
</tr>
</tbody>
</table>

* Levels shown indicate the spike concentration in the final digestate of the aqueous LCS or matrix spike based on the addition of 0.5 mL working spike (7.4) to 50 mL of sample.
### TABLE VI. Summary Of Quality Control Requirements

<table>
<thead>
<tr>
<th>QC PARAMETER</th>
<th>FREQUENCY</th>
<th>ACCEPTANCE CRITERIA</th>
<th>CORRECTIVE ACTION</th>
</tr>
</thead>
<tbody>
<tr>
<td>Method Blank</td>
<td>One per sample preparation batch of up to 20 samples.</td>
<td>Refer to determinative SOPs:</td>
<td>Redigest and reanalyze samples associated with the method blank.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>CORP-MT-0001</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>CORP-MT-0003</td>
<td></td>
</tr>
<tr>
<td>Laboratory Control Sample (LCS)</td>
<td>One per sample preparation batch of up to 20 samples.</td>
<td>Refer to determinative SOPs:</td>
<td>Redigest and reanalyze all samples associated with the LCS.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>CORP-MT-0001</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>CORP-MT-0003</td>
<td></td>
</tr>
<tr>
<td>Matrix Spike</td>
<td>One per sample preparation batch of up to 20 samples.</td>
<td>Refer to determinative SOPs:</td>
<td>Reprep not required unless preparation error suspected.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>CORP-MT-0001</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>CORP-MT-0003</td>
<td></td>
</tr>
<tr>
<td>Matrix Spike Duplicate</td>
<td>See Matrix Spike</td>
<td>Refer to determinative SOPs:</td>
<td>See Corrective Action for Matrix Spike.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>CORP-MT-0001</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>CORP-MT-0003</td>
<td></td>
</tr>
<tr>
<td>Prep Date</td>
<td>Analyst</td>
<td>SLICE</td>
<td>ID</td>
</tr>
<tr>
<td>-----------</td>
<td>---------</td>
<td>-------</td>
<td>----</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

SOP (circle)  
QC Batch:  
Method #:  

Matrix:  
Prep Type: (e.g.): ICP-GEA
APPENDIX C

CONTAMINATION CONTROL GUIDELINES
APPENDIX C. CONTAMINATION CONTROL GUIDELINES

The following procedures are strongly recommended to prevent contamination:

- All work areas used to prepare standards and spikes should be cleaned before and after each use.
- All glassware should be washed with detergent and tap water and rinsed with 1:1 nitric acid followed by deionized water.
- Proper laboratory housekeeping is essential in the reduction of contamination in the metals laboratory. All work areas must be kept scrupulously clean.
- Powdered or Latex Gloves must not be used in the metals laboratory since the powder contains silica and zinc, as well as other metallic analytes. Only vinyl or nitrile gloves should be used in the metals laboratory.
- Glassware should be periodically checked for cracks and etches and discarded if found. Etched glassware can cause cross contamination of any metallic analytes.
- Autosampler trays should be covered to reduce the possibility of contamination. Trace levels of elements being analyzed in the samples can be easily contaminated by dust particles in the laboratory.

The following are helpful hints in the identification of the source of contaminants:

- Reagents or standards can contain contaminants or be contaminated with the improper use of a pipette.
- Improper cleaning of glassware can cause contamination.
- Separate glassware if an unusually high sample is analyzed and soak with sulfuric acid prior to routine cleaning.
LABORATORY-SPECIFIC

STL STANDARD OPERATING PROCEDURE

TITLE: PREPARATION AND ANALYSIS OF MERCURY IN AQUEOUS SAMPLES BY COLD VAPOR ATOMIC ABSORPTION, SW846 7470A AND MCAWW 245.1

(SUPERSEDES: CORP-MT-0005, REVISION 1)

Prepared by: 

Reviewed by: Technical Specialist

Approved by: Quality Assurance Manager

Approved by: Environmental Health and Safety Coordinator

Approved by: Laboratory Director

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1. **SCOPE AND APPLICATION**

1.1. This procedure describes the preparation and analysis of mercury (Hg, CAS # 7439-97-6) by Cold Vapor Atomic Absorption Spectroscopy (CVAA) using SW-846 Method 7470A and MCAWW Method 245.1. Both the water bath digestion and the autoclave digestion are available at the STL Pittsburgh facility, however the default practice is the autoclave digestion. Both are described in this SOP.

1.2. CVAA analysis provides for the determination of total mercury (organic and inorganic). The combination of the oxidants, potassium permanganate and potassium persulfate, has been found to give 100% recovery with both types of compounds. Detection limits, sensitivity and optimum concentration ranges for mercury analysis will vary with the matrices, instrumentation and volume of sample used.

1.3. Method 7470A is applicable to the preparation and analysis of mercury in ground water, aqueous samples, wastes, wipes, TCLP, EP and other leachates/extracts. Certain solid and sludge type wastes may also be analyzed, however Method 7471A (see CORP-MT-0007) is usually the method of choice. All matrices require sample preparation prior to analysis.

1.4. Method 245.1 is applicable to the determination of mercury in drinking, surface and saline waters, domestic and industrial wastes. All matrices require sample preparation prior to analysis.

1.5. The STL reporting limit for mercury in aqueous matrices is 0.0002 mg/L except for TCLP, SPLP or EPTOX leachates for which the reporting limit is 0.002 mg/L.

2. **SUMMARY OF METHOD**

2.1. This SOP describes a technique for the determination of mercury in solution. The procedure is a physical method based on the absorption of radiation at 253.7 nm by mercury vapor. A representative portion of the sample is digested in sulfuric and nitric acids. Organic mercury compounds are oxidized with potassium permanganate and potassium persulfate and the mercury reduced to its elemental state with stannous chloride and aerated from solution in a closed system. The mercury vapor passes through a cell positioned in the light path of an atomic absorption spectrophotometer. Absorbance is measured as a function of mercury concentration. Concentration of the analyte in the sample is determined by comparison of the sample absorbance to the calibration curve (absorbance vs. concentration).
3. DEFINITIONS

3.1. CRA: A standard which is run a part of the initial calibration, also known as "The Reporting Level Verification Standard"

3.2. Dissolved Metals: Those elements which pass through a 0.45 µm membrane. (Sample is acidified after filtration).

3.3. Suspended Metals: Those elements which are retained by a 0.45 µm membrane.

3.4. Total Metals: The concentration determined on an unfiltered sample following digestion.

4. INTERFERENCES

Chemical and physical interferences may be encountered when analyzing samples using this method.

4.1. Potassium permanganate which is used to breakdown organic mercury compounds also eliminates possible interferences from sulfide. Concentrations as high as 20 mg/L of sulfide as sodium sulfide do not interfere with the recovery of inorganic mercury from reagent water.

4.2. Copper has also been reported to interfere; however, copper concentrations as high as 10 mg/L had no effect on the recovery of mercury from spiked samples.

4.3. Chlorides can cause a positive interference. Seawaters, brines and industrial effluents high in chlorides require additional permanganate (as much as 25 mL) because, during the oxidation step, chlorides are converted to free chlorine, which also absorbs radiation at 253.7 nm. Care must be taken to ensure that free chlorine is absent before the mercury is reduced and swept into the cell. This is accomplished by adding excess hydroxylamine reagent (25 mL) and purging the sample head space before stannous chloride is added. Both inorganic and organic mercury spikes have been quantitatively recovered from seawater using this technique.

Note: Sufficient addition of permanganate is apparent when the purple color persists at least 15 minutes. Some samples may require dilution prior to digestion due to extremely high concentrations of chloride.
by subtracting the result of the sample run without the reducing reagent (stannous chloride) from that obtained with the reducing reagent.

4.5. Samples containing high concentrations of oxidizable organic materials, as evidenced by high COD levels, may not be completely oxidized by this procedure. When this occurs the recovery of mercury will be low. The problem can be eliminated by reducing the volume of original sample used.

4.6. The most common interference is laboratory contamination which may arise from impure reagents, dirty glassware, improper sample transfers, dirty work areas, etc. Be aware of potential sources of contamination and take appropriate measures to minimize or avoid them.

5. SAFETY

5.1. Procedures shall be carried out in a manner that protects the health and safety of all STL associates.

5.2. Eye protection that satisfies ANSI Z87.1 (as per the Chemical Hygiene Plan), laboratory coat, and appropriate gloves must be worn while samples, standards, solvents, and reagents are being handled. Disposable gloves that have been contaminated will be removed and discarded; other gloves will be cleaned immediately.

5.3. The health and safety hazards of many of the chemicals used in this procedure have not been fully defined. Additional health and safety information can be obtained from the Material Safety Data Sheets (MSDS) maintained in the laboratory. The following specific hazards are known:

5.3.1. The following materials are known to be corrosive:

hydrochloric acid, nitric acid and sulfuric acid.

5.3.2. The following materials are known to be oxidizing agents:

nitric acid, potassium permanganate, potassium persulfate and magnesium perchlorate.

5.3.3. Mercury is a highly toxic element that must be handled with care. The analyst must be aware of the handling and clean-up techniques before working with mercury. Since mercury vapor is toxic, precaution must be taken to avoid its inhalation, ingestion or absorption through skin. All lines should be checked for
leakage and the mercury vapor must be vented into a hood or passed through a mercury absorbing media such as:

5.3.3.1. Equal volumes of 0.1 M KMnO₄ and 10% H₂SO₄, or

5.3.3.2. Iodine, 0.25%, in a 3% KI solution.

5.3.4. Magnesium sulfate is known to be a reproductive toxin (mutagen).

5.4. Exposure to chemicals must be maintained as low as reasonably achievable. Therefore, unless they are known to be non-hazardous, all samples should be opened, transferred and prepared in a fume hood, or under other means of mechanical ventilation. Solvent and waste containers will be kept closed unless transfers are being made.

5.5. The preparation of standards and reagents will be conducted in a fume hood with the sash closed as far as the operation will permit.

5.6. All work must be stopped in the event of a known or potential compromise to the health and safety of a STL associate. The situation must be reported immediately to a laboratory supervisor.

5.7. Do not look directly into the beam of the Hg lamp. The UV light that these lamps radiate is harmful to the eyes.

5.8. Cylinders of compressed gas must be handled with caution, in accordance with local regulations. It is recommended that, wherever possible, cylinders be located outside the laboratory and the gas led to the instrument through approved lines.

5.9. The CVAA apparatus must be properly vented to remove potentially harmful fumes generated during sample analysis.

6. EQUIPMENT AND SUPPLIES

6.1. Temperature controlled water bath (capable of maintaining a temperature of 90-95 °C) or autoclave that is able to obtain conditions of 15 lbs., 120 °C for 15 minutes.

6.2. Atomic Absorption Spectrophotometer equipped with:

6.2.1. Absorption Cell with quartz end windows perpendicular to the longitudinal axis. Dimensions of the cell must result in sufficient sensitivity to meet the SOP defined reporting limit. The quartz windows must be maintained to provide
accurate measurements. Any scratches or fingerprints can alter the absorption of UV radiation.

6.2.2. Mercury specific hollow cathode lamp (HCL) or electrodeless discharge lamp (EDL).

6.2.3. Peristaltic pump which can deliver 1 L/min air.

6.2.4. Flowmeter capable of measuring an airflow of 1 L/min.

6.2.5. Recorder or Printer.

6.2.6. Aeration Tubing: A straight glass frit having a course porosity and Tygon tubing is used for the transfer of mercury vapor from the sample bottle to the absorption cell and return.

6.2.7. Drying device (a drying tube containing magnesium perchlorate or magnesium sulfate and/or a lamp with a 60 W bulb) to prevent condensation in cell. The lamp is positioned to shine on the absorption cell maintaining the air temperature in the cell about 10 °C above room temperature. Other drying devices that achieve the same purpose are also acceptable (i.e., Gortex filter).

**NOTE:** Instruments designed specifically for the measurement of mercury using the cold vapor technique may be substituted for the atomic absorption spectrophotometer.

6.3. BOD bottles or equivalent.

6.4. Nitrogen or argon gas supply, welding grade or equivalent.

6.5. Calibrated automatic pipettes or Class A glass volumetric pipettes.

6.6. Class A volumetric flasks.

6.7. Thermometer (capable of accurate readings at 95 °C).

6.8. Disposable cups or tubes.
7. REAGENTS AND STANDARDS

7.1. Reagent water must be produced by a Millipore DI system or equivalent. Reagent water must be free of the analytes of interest as demonstrated through the analysis of method blanks.

7.2. Stock (1000 ppm) mercury standards (in 10% HNO₃) are purchased as custom STL solutions. All standards must be stored in FEP fluorocarbon or previously unused polyethylene or polypropylene bottles. Stock standard solutions must be replaced prior to the expiration date provided by the manufacturer. If no expiration date is provided, the stock solutions may be used for up to one year and must be replaced sooner if verification from an independent source indicates a problem.

7.3. Intermediate mercury standard (10 ppm): Take 1 mL of the stock mercury standard (7.2) and dilute to 100 mL with reagent water. The intermediate standard must be made monthly and must be prepared in a matrix of 2% HNO₃. This acid (2 mL of concentrated HNO₃) must be added to the flask/bottle before the addition of the stock standard aliquot.

7.4. Working mercury standard (0.1 ppm): Take 1 mL of the intermediate mercury standard (7.3) and dilute to 100 mL with reagent water. The working mercury standard must be made daily and must be prepared in a matrix of 0.15% HNO₃. This acid (150 µL of concentrated HNO₃) must be added to the flask/bottle before the addition of the stock standard aliquot.

7.5. The calibration standards listed in Table I must be prepared fresh daily from the working standard (7.4) by transferring 0, 0.2, 0.5, 1.0, 2.0, 5.0 and 10.0 mL aliquots of the working mercury standard into 100 mL flasks and diluting to volume with reagent water.

Note: Alternate approaches to standard preparation may be taken and alternate volumes of standard may be prepared as long as the accuracy and final standard concentrations as detailed in Table I are maintained. For example, automated mercury systems do not require 100 mL of standard and therefore smaller volumes may be generated to reduce waste generation.

7.6. The initial calibration verification standard must be made from a different stock solution than that of the calibration standards.

7.7. Refer to Table I (Appendix A) for details regarding the working standard concentrations for calibration, calibration verification and spiking solutions. All standards must be processed through the entire analytical procedure including sample preparation.
7.8. Nitric acid (HNO₃), concentrated, trace metal grade or better.

Note: If a high reagent blank is obtained, it may be necessary to distill the nitric acid.

7.9. Sulfuric acid (H₂SO₄), concentrated, trace metal grade or better.

7.9.1. Sulfuric acid, 0.5 N: Dilute 14.0 mL of concentrated H₂SO₄ to 1 liter with reagent water.

7.10. Stannous sulfate solution: Add 25 g of stannous sulfate to 250 mL of 0.5 N sulfuric acid. This mixture is a suspension and should appear cloudy. This solution should be made daily and should be stirred continuously during use.

Note: Stannous chloride may be used in place of stannous sulfate. Prepare the stannous chloride solution according to the recommendations provided by the instrument manufacturer.

7.11. Sodium chloride-hydroxylamine hydrochloride solution: Add 12 g of sodium chloride and 12 g of hydroxylamine hydrochloride to every 100 mL of reagent water.

Note: Hydroxylamine sulfate may be used in place of hydroxylamine hydrochloride.

7.12. Potassium permanganate, 5% solution (w/v): Dissolve 5 g of potassium permanganate for every 100 mL of reagent water.

7.13. Potassium persulfate, 5% solution (w/v): Dissolve 5 g of potassium persulfate for every 100 mL of reagent water.

8. SAMPLE COLLECTION, PRESERVATION AND STORAGE

8.1. Sample holding time for mercury is 28 days from time of collection to the time of analysis.

8.2. Aqueous samples are preserved with nitric acid to a pH of <2 and may be stored in either plastic or glass. Refrigeration is not required. Preservation must be verified prior to analysis.

9. QUALITY CONTROL

Table II (Appendix A) provides a summary of quality control requirements including type, frequency, acceptance criteria and corrective action.

9.1. Initial Demonstration of Capability
Prior to the analysis of any analyte using 7470A or the 245.1, the following requirements must be met.

9.1.1. Method Detection Limit (MDL) - An MDL must be determined for each analyte/matrix prior to the analysis of any samples. The MDL is determined using seven replicates of reagent water, spiked with all the analytes of interest, that have been carried through the entire analytical procedure. MDLs must be redetermined on an annual basis in accordance with 40 CFR Part 136 Appendix B requirements. The spike level must be between the calculated MDL and 10X the MDL to be valid. The result of the MDL determination must be below the STL reporting limit.

9.1.2. Initial Demonstration Study - This requires the analysis of four QC check samples. The QC check sample is a well characterized laboratory generated sample used to monitor method performance. The results of the initial demonstration study must be acceptable before analysis of samples may begin.

9.1.2.1. Four aliquots of the check sample (LCS) are prepared and analyzed using the procedures detailed in this SOP and the determinative SOPs.

9.2. Preparation Batch - A group of up to 20 samples that are of the same matrix and are processed together using the same procedures and reagents. The preparation batch must contain a method blank, a LCS and a matrix spike/matrix spike duplicate. In some cases, at client request, it may be appropriate to process a matrix spike and sample duplicate in place of the MS/MSD. If clients specify specific samples for MS/MSD, the batch may contain multiple MS/MSD pairs.

9.3. Sample Count - Laboratory generated QC samples (method blanks, LCS) are not included in the sample count for determining the size of a preparation batch. MS/MSD are not included in the sample count unless there are multiple sets of MS/MSD per batch. In other words, the first MS/MSD are not counted; all additional MS and MSDs are counted as samples.

9.4. Method Blank (MB) - One method blank must be processed with each preparation batch. The method blank consists of reagent water containing all reagents specific to the method that is carried through the entire analytical procedure, including preparation and analysis. The method blank is used to identify any system and process interferences or contamination of the analytical system that may lead to the reporting of elevated analyte concentrations or false positive data. The method blank should not contain any analyte of interest at or above the reporting limit, or above 10% of either the measured concentration of that analyte in associated samples or the regulatory limit. See QA-003 for more detail on criteria and corrective actions. In addition, blank contamination should always be evaluated against project specific requirements.
• Repreparation and reanalysis of all samples associated with an unacceptable method blank is required when reportable concentrations are determined in the samples (see exception noted above).

• If there is no analyte greater than the RL in the samples associated with an unacceptable method blank, the data may be reported with qualifiers. Such action must be taken in consultation with the client and must be addressed in the project narrative.

• If the above criteria are not met and reanalysis is not possible, then the sample data must be qualified. This anomaly must be addressed in the project narrative and the client must be notified.

9.5. Laboratory Control Sample (LCS) - One aqueous LCS must be processed with each preparation batch. The LCS is used to monitor the accuracy of the analytical process. On-going monitoring of the LCS results provides evidence that the laboratory is performing the method within acceptable accuracy and precision guidelines. The LCS must be carried through the entire analytical procedure. The CCV results can be reported as LCS results since all CCVs (as well as all other standards) are processed through the sample preparation step with the field samples. No more than 20 samples can be associated with one CCV used for the purpose of reporting LCS data.

• If the LCS is outside established control limits the system is out of control and corrective action must occur. Until in-house control limits are established, a control limit of 80 - 120% recovery must be applied.

• In the instance where the LCS recovery is > 120% and the sample results are < RL, the data may be reported with qualifiers. Such action must be taken in consultation with the client and must be addressed in the case narrative.

• In the event that an MS/MSD analysis is not possible, a Laboratory Control Sample Duplicate (LCSD) must be analyzed. The RPD of the LCS and LCSD must be compared to the matrix spike RPD limits.

• Corrective action will be repreparation and reanalysis of the batch unless the client agrees that other corrective action is acceptable.

9.6. Matrix Spike/Matrix Spike Duplicate (MS/MSD) - One MS/MSD pair must be processed for each preparation batch. A matrix spike (MS) is a field sample to which known concentrations of target analytes have been added. A matrix spike duplicate (MSD) is a second aliquot of the same sample (spiked identically as the MS) prepared and analyzed along with the sample and matrix spike. Some client specific data quality objectives (DQO's) may require the use of sample duplicates in place of or in addition to
MS/MSD’s. The MS/MSD results are used to determine the effect of a matrix on the precision and accuracy of the analytical process. Due to the potential variability of the matrix of each sample, these results may have immediate bearing only on the specific sample spiked. Samples identified as field blanks cannot be used for MS/MSD analysis. Spiking levels are provided in Table I (Appendix A).

- If analyte recovery or RPD falls outside the acceptance range, the recovery of that analyte must be in control for the LCS. Until in-house control limits are established, a control limit of 75 - 125 % recovery and 20% RPD must be applied to the MS/MSD. If the LCS recovery is within limits, then the laboratory operation is in control and the results may be accepted. If the recovery of the LCS is outside limits, corrective action must be taken. Corrective action will include repreparation and reanalysis of the batch. MS/MSD results which fall outside the control limits must be addressed in the narrative.

- If the native analyte concentration in the MS/MSD exceeds 4 times the spike level for that analyte, the recovery data are reported as NC (i.e., not calculated). If the reporting software does not have the ability to report NC then the actual recovery must be reported and narrated as follows: “Results outside of limits do not necessarily reflect poor method performance in the matrix due to high analyte concentrations in the sample relative to the spike level.”

- If an MS/MSD is not possible due to limited sample volume, then a laboratory control sample duplicate (LCSD) should be analyzed. The RPD of the LCS and LCSD must be compared to the matrix spike RPD limits.

9.7. Initial Calibration Verification (ICV/ICB) - Calibration accuracy is verified by analyzing a second source standard (ICV). The ICV result must fall within 20% of the true value for that solution. An ICB is analyzed immediately following the ICV to monitor low level accuracy and system cleanliness. The ICB result must fall within +/- the reporting limit (RL) from zero. If either the ICV or ICB fail to meet criteria, the analysis should be terminated, the problem corrected and the instrument recalibrated. (See Section 11.2.11 and Section 11.2.12 for required run sequence). If the cause of the ICV or ICB failure was not directly instrument related the corrective action will include repreparation of the associated samples.

9.8. Continuing Calibration Verification (CCV/CCB) - Calibration accuracy is monitored throughout the analytical run through the analysis of a known standard after every 10 samples. The CCV must be a mid-range standard at a concentration other than that of the ICV. The CCV result must fall within 20% of the true value for that solution. A CCB is analyzed immediately following each CCV. (See Section 11.2.11 and 11.2.12 for required run sequence.) The CCB result must fall within +/- RL from zero. Each CCV and CCB analyzed must reflect the conditions of analysis of all associated samples.
Sample results may only be reported when bracketed by valid ICV/CCV and ICB/CCB pairs. If a mid-run CCV or CCB fails, the analysis must be terminated, the problem corrected, the instrument recalibrated, the calibration verified and the affected samples reanalyzed. If the cause of the CCV or CCB failure was not directly instrument related the corrective action will include repreparation of the associated samples.

9.9. Method of Standard Addition (MSA) - This technique involves adding known amounts of standard to one or more aliquots of the sample prior to preparation. This technique compensates for a sample interferent that may enhance or depress the analyte signal, thus producing a different slope from that of the calibration standards. It will not correct for additive interferences which cause a baseline shift. Refer to Section 11.2.13 for additional information on when full 4 point MSA is required as well as Appendix C for specific MSA requirements.

10. CALIBRATION AND STANDARDIZATION

10.1. Calibration standards must be processed through the preparation procedure as described in Section 11.1.

10.2. Due to the differences in preparation protocols separate calibration and calibration verification standards must be prepared for aqueous and solid matrices.

10.3. Calibration must be performed daily (every 24 hours) and each time the instrument is set up. The instrument calibration date and time must be included in the raw data.

10.4. Set up the instrument with the operating parameters recommended by the manufacturer. Allow the instrument to become thermally stable before beginning calibration (approximately 30 minutes of warm-up is required). Refer to the facility specific instrument SOP and CVAA instrument manual for detailed setup and operation protocols.

10.5. Calibrate the instrument according to instrument manufacturer’s instructions, using a minimum of five standards and a blank. One standard must be at the STL reporting limit. Analyze standards in ascending order beginning with the blank. Refer to Section 7.5 and Table I for additional information on preparing calibration standards and calibration levels.

10.6. The calibration curve must have a correlation coefficient of ≥0.995 or the instrument shall be stopped and recalibrated prior to running samples. Sample results can not be reported from a curve with an unacceptable correlation coefficient.
10.7. Refer to Section 9.0 for calibration verification procedures, acceptance criteria and corrective actions.

11. PROCEDURE

11.1. Sample Preparation:

11.1.1. All calibration and calibration verification standards (ICV, ICB, CCV, CCB) are processed through the digestion procedure as well as the field samples. *An exception to this is for Method 245.1 samples. The calibration curve samples are not heated.*

11.1.2. Transfer 100 mL of well mixed sample or standard to a clean sample digestion bottle.

*Note:* Reduced sample volumes can be used as long as a representative sample can be obtained and the reagent levels are adjusted to maintain the same sample to reagent ratio. All samples and standards must be processed similarly.

11.1.3. Add 5 mL of concentrated H₂SO₄ and 2.5 mL of concentrated HNO₃ mixing after each addition.

*Note:* All spiking should be done after the initial addition of acids.

11.1.4. Add 15 mL of potassium permanganate solution. For samples high in organic materials or chlorides, additional permanganate may be added. Shake and add additional portions of permanganate solution until a purple color persists for at least 15 minutes. If after the addition of up to 25 mL additional permanganate the color does not persist, sample dilution prior to reanalysis may be required.

*Note:* When performing analyses using automated vs. manual techniques the sample dilution resultant from the addition of more than the original aliquot of permanganate solution must be compensated for by the addition of the same volume of permanganate to all associated samples, standards, and QC samples (e.g. LCS and blank) in the run. In instances, where this is not feasible, the addition of excess reagent can be addressed through mathematical correction of the results to account for the resultant dilution effect.
be addressed through mathematical correction of the results to account for the resultant dilution effect.

11.1.5. Add 8 mL of potassium persulfate solution and heat for two hours in a water bath at 90 - 95 °C.

NOTE: Alternatively, for RCRA analyses using 7470A, samples may be digested using an autoclave for 15 minutes at 120 °C and 15 lbs.

11.1.6. Cool samples.

11.2. Sample Analysis:

11.2.1. Because of differences between various makes and models of CVAA instrumentation, no detailed operating instructions can be provided. Refer to the facility specific instrument operating SOP and the CVAA instrument manual for detailed setup and operation protocols.

11.2.2. All labs are required to detail the conditions/programs utilized for each instrument within the facility specific instrument operation SOP.

11.2.3. When ready to begin analysis, add 6 mL of sodium chloride-hydroxylamine hydrochloride solution to the samples to reduce the excess permanganate (the permanganate has been reduced when no purple color remains). Add this solution in 6 mL increments until the permanganate is completely reduced.

11.2.4. Manual determination:
11.2.4.1. Treating each sample individually, purge the head space of the sample bottle for at least one minute.

11.2.4.2. Add 5 mL of stannous chloride solution and immediately attach the bottle to the aeration apparatus.

11.2.4.3. Allow the sample to stand quietly without manual agitation while the sample is aerated (1 L/min flow). Monitor the sample absorbance during aeration. When the absorbance reaches a maximum and the signal levels off, open the bypass valve and continue aeration until the absorbance returns to its baseline level. Close the bypass valve and remove the aeration device.

11.2.4.4. Place the aeration device into 100 mL of 1% HNO₃ and allow to bubble rinse until the next sample is analyzed.

11.2.5. Automated determination: Follow instructions provided by instrument manufacturer.

11.2.6. Perform a linear regression analysis of the calibration standards by plotting maximum response of the standards vs. concentration of mercury. Determine the mercury concentration in the samples from the linear regression fit of the calibration curve. Calibration using computer or calculation based regression curve fitting techniques on concentration/response data is acceptable.

11.2.7. All measurements must fall within the defined calibration range to be valid. Dilute and reanalyze all samples for analytes that exceed the highest calibration standard.

11.2.8. If the sample results are negative and the absolute value of the negative result is greater than the reporting limit, the sample must be diluted and reanalyzed.

11.2.9. The samples must be allowed to cool to room temperature prior to analysis or a decrease in the response signal can occur.
11.2.10. Baseline correction is acceptable as long as it is performed after every sample or after the CCV and CCB; resloping is acceptable as long as it is immediately preceded and followed by a compliant CCV and CCB.

11.2.11. The following analytical sequence must be used with 7470A and 245.1:

Instrument Calibration  
ICV  
ICB  
Maximum 10 samples  
CCV  
CCB  
Repeat sequence of 10 samples between CCV/CCB pairs as required to complete run  
CCV  
CCB

Refer to Quality Control Section 9.0 and Table II (Appendix A) for quality control criteria to apply to Methods 7470A and 245.1.

Note: Samples include the method blank, LCS, MS, MSD, duplicate, field samples and sample dilutions.

11.2.12. The following run sequence is consistent with 7470A, CLP and 245.1 and may be used as an alternate to the sequence in 11.2.11. This run sequence is recommended if multiple method requirements must be accommodated in one analytical run:

Instrument Calibration  
ICV  
ICB  
CRA* (Reporting Level Verification Standard)  
CCV  
CCB  
10 samples  
CCV  
CCB  
Repeat sequence of 10 samples between CCV/CCB pairs as required to complete run.  
CCV  
CCB
Refer to the appropriate CLP SOPs (CORP-MT-0006) for quality control requirements for QC samples.

* Refer to the CLP SOPs for information on the CRA.

11.2.13. For TCLP samples, full four point MSA will be required if all of the following conditions are met:

1) recovery of the analyte in the matrix spike is not at least 50%,
2) the concentration of the analyte does not exceed the regulatory level, and,
3) the concentration of the analyte is within 20% of the regulatory level.

The reporting and matrix spike levels for TCLP analyses are detailed in Table I (Appendix A). Appendix E provides guidance on performing MSA analyses. For TCLP mercury determinations, MSA spikes must be added prior to sample preparation.

11.3. To facilitate the early identification of QC failures and samples requiring rerun it is strongly recommended that sample data be reviewed periodically throughout the run.

11.4. Guidelines are provided in the appendices on procedures to minimize contamination of samples and standards, preventive maintenance and troubleshooting.

11.5. One time procedural variations are allowed only if deemed necessary in the professional judgment of supervision to accommodate variation in sample matrix, radioactivity, chemistry, sample size, or other parameters. Any variation in procedure shall be completely documented using a Nonconformance Memo and is approved by a Technical Specialist and QA Manager. If contractually required, the client shall be notified. The Nonconformance Memo shall be filed in the project file.

11.6. Any unauthorized deviations from this procedure must also be documented as a nonconformance, with a cause and corrective action described.

12. **DATA ANALYSIS AND CALCULATIONS**

12.1. ICV percent recoveries are calculated according to the equation:

\[
\% R = 100 \left( \frac{\text{Found(ICV)}}{\text{True(ICV)}} \right)
\]

12.2. CCV percent recoveries are calculated according to the equation:
\[
\% R = 100 \left( \frac{\text{Found(CCV)}}{\text{True(CCV)}} \right)
\]

12.3. Matrix spike recoveries are calculated according to the following equation:

\[
\% R = 100 \left( \frac{\text{SSR} - \text{SR}}{\text{SA}} \right)
\]

Where:
- SSR = Spike Sample Result
- SR = Sample Result
- SA = Spike Added

12.4. The relative percent difference (RPD) of matrix spike/matrix spike duplicates or sample duplicates are calculated according to the following equations:

\[
\text{RPD} = 100 \left( \frac{|\text{MSD} - \text{MS}|}{\frac{\text{MSD} + \text{MS}}{2}} \right)
\]

Where:
- MS = determined spiked sample concentration
- MSD = determined matrix spike duplicate concentration

\[
\text{RPD} = 100 \left( \frac{|\text{DU1} - \text{DU2}|}{\frac{\text{DU1} + \text{DU2}}{2}} \right)
\]

Where:
- DU1 = Sample result
- DU2 = Sample duplicate result

12.5. The final concentration for an aqueous sample is calculated as follows:

\[
\text{mg/L} = C \times D
\]

Where:
- C = Concentration (mg/L) from instrument readout
- D = Instrument dilution factor
12.6. The LCS percent recovery is calculated according to the following equation:

\[
% R = 100 \left( \frac{\text{Found}(\text{LCS})}{\text{True}(\text{LCS})} \right)
\]

12.7. Appropriate factors must be applied to sample values if dilutions are performed.

12.8. Sample results should be reported with up to three significant figures in accordance with the STL significant figure policy.

13. **METHOD PERFORMANCE**

13.1. The CCV will be varied periodically to demonstrate verification of linearity of the curve.

13.2. Each laboratory must have initial demonstration of performance data on file for each analyte of interest as described in Section 9.0.

13.3. Method performance is determined by the analysis of method blanks, laboratory control samples, matrix spike and matrix spike duplicate samples. The matrix spike recovery should fall within +/- 25 % and the matrix spike duplicates should compare within 20% RPD. The method blanks must meet the criteria in Section 9.3. The laboratory control sample should recover within 20% of the true value until in house limits are established.

13.4. Training Qualification:

The group/team leader has the responsibility to ensure that this procedure is performed by an associate who has been properly trained in its use and has the required experience.

14. **POLLUTION PREVENTION**

14.1. This method allows for the proportional reduction of sample and reagent volumes to decrease waste generation.


16.4. QA-003, STL QC Program.

16.5. QA-004, Rounding and Significant Figures.

16.6. QA-005, Method Detection Limits.

17. MISCELLANEOUS (TABLES, APPENDICES, ETC. . . )

17.1. Modifications/Interpretations from reference method.

17.1.1. Modifications from both 7470A and 245.1.

17.1.1.1. The 200 series methods and Chapter 1 of SW846 specify the use of reagent water with a purity equivalent to ASTM Type II water. This SOP specifies the use of a Millipore DI system or equivalent to produce reagent water. This SOP requires that reagent water must be free of the analytes of interest as demonstrated through the analysis of method blanks.

17.1.1.2. This SOP allows for the use of reduced sample volumes to decrease waste generation. Reagent levels are adjusted to maintain the same ratios as stated in the source methods. According to a letter from Robert Booth of EPA EMSL-Cim to David Payne of EPA Region V, “Reduction in sample size and appropriate corresponding reduction in sample volume is not considered a significant change in the methodology.”

17.1.1.3. The alternate run sequence presented in Section 11.2.12 is consistent with method requirements. An additional QC analysis (CRA) was added to accommodate the CLP protocol requirements.

17.1.2. Modifications from Method 7470A

17.1.2.1. Chapter 1 of SW-846 states that the method blank should not contain any analyte of interest at or above the MDL. This SOP states that the method blank must not contain any analyte of interest at or above the reporting limit.
17.1.2.2. Documentation is on file from EPA’s Office of Solid Waste (Olliver Fordham 11/28/95) regarding the acceptance of the autoclave as an equivalent heating device to the water bath. In his letter, Mr. Fordham cited the CLP water protocol 245.1 CLP-M and therefore the operating parameters from that method were adopted for 7470A (15 minutes at 120 °C and 15 lbs.).

17.1.2.3. Method 7470A does not state control criteria within the text of the method. The QC section of 7470A refers the analyst to Section 8.0 of Method 7000A, the generic atomic absorption method, which discusses flame and furnace methods. The ICV criteria stated in Method 7000A is ±10%. This SOP requires ICV control limits of ±20% based on the fact that the mercury ICV, unlike the ICV for the flame and furnace analytes, is digested and therefore is equivalent to a LCS. The CLP protocol 245.1 CLP-M recognizes this factor and requires control limits of ±20%.

17.1.3. Modifications from 245.1

17.1.3.1. Method 245.1 Section 9.3 states concentrations should be reported as follows: Between 1 and 10 μg/L, one decimal; above 10 μg/L, to the nearest whole number. STL reports all Hg results under this SOP to two significant figures.

17.2. Documentation and Record Management

The following documentation comprises a complete CVAA raw data package:

- Raw data (direct instrument printout)
- Run log printout from instrument software where this option is available or manually generated run log. (A bench sheet may be substituted for the run log as long as it contains an accurate representation of the analytical sequence).
- Data review checklist - See Appendix B
- Standards Documentation (source, lot, date).
- Copy of digestion log.
- Non-conformance summary (if applicable).
Figure 1. Aqueous Sample Preparation - Mercury

Aliquot samples and standards to prep bottles

Add conc. H$_2$SO$_4$ and HNO$_3$

Add KMNO$_4$

Add Potassium Persulfate

Heat for 2 hrs. at 95 C or for 15 min at 120 C and 15 lbs. (7470A)

Cool
Figure 2. CVAA Mercury Analysis

1. Set up instrument
2. Reduce Excess Permanganate
3. Construct calibration curve
   - Recalibrate
   - Yes: ICV, ICB in control?
   - No: Failure instrument related?
   - Yes: Reprep and rerun samples.
   - No: Run samples
5. Results < high std.?
   - Yes: CCV, CCB in control?
   - No: Dilute and rerun samples.
   - Yes: Report results
   - No: Failure instrument related?
   - Yes: Recalibrate and rerun samples.
   - No: Reprep and rerun samples.
APPENDIX A

TABLES
### TABLE I. MERCURY REPORTING LIMITS, CALIBRATION STANDARD*, QC STANDARD AND SPIKING LEVELS (MG/L)

<table>
<thead>
<tr>
<th>Standard Aqueous RL</th>
<th>0.0002</th>
</tr>
</thead>
<tbody>
<tr>
<td>TCLP RL</td>
<td>0.002</td>
</tr>
<tr>
<td>Std 0</td>
<td>0</td>
</tr>
<tr>
<td>Std 1</td>
<td>0.0002</td>
</tr>
<tr>
<td>Std 2</td>
<td>0.0005</td>
</tr>
<tr>
<td>Std 3</td>
<td>0.001</td>
</tr>
<tr>
<td>Std 4</td>
<td>0.002</td>
</tr>
<tr>
<td>Std 5</td>
<td>0.005</td>
</tr>
<tr>
<td>Std 6 **</td>
<td>0.010</td>
</tr>
<tr>
<td>ICV</td>
<td>0.001 or 0.0025 ***</td>
</tr>
<tr>
<td>LCS/CCV</td>
<td>0.0025 or 0.005 ***</td>
</tr>
<tr>
<td>Aqueous MS</td>
<td>0.001</td>
</tr>
<tr>
<td>TCLP MS</td>
<td>0.005</td>
</tr>
</tbody>
</table>

* SOP specified calibration levels must be used unless prevented by the instrument configuration or client specific requirements. Deviations from specified calibration levels must be documented in the facility specific instrument operation SOP and must be approved by the facility technical manager and Quality Assurance Manager.

** Optional standard which may be used to extend the calibration range as allowed by the instrument configuration. If the instrument configuration prevents the use of 6 standards, the 2 ppb standard may be eliminated in favor of the 10 ppb standard.

*** Concentration level dependent on high calibration standard used. CCV must be 50% of high standard concentration and ICV must be 20-25% of high standard concentration.
### TABLE II. Summary Of Quality Control Requirements

<table>
<thead>
<tr>
<th>QC PARAMETER</th>
<th>FREQUENCY *</th>
<th>ACCEPTANCE CRITERIA</th>
<th>CORRECTIVE ACTION</th>
</tr>
</thead>
<tbody>
<tr>
<td>ICV</td>
<td>Beginning of every analytical run.</td>
<td>80 - 120% recovery.</td>
<td>Terminate analysis; Correct the problem; Recalibrate or reprep batch (see Section 9.7).</td>
</tr>
<tr>
<td>ICB</td>
<td>Beginning of every analytical run, immediately following the ICV.</td>
<td>The result must be within +/- RL from zero.</td>
<td>Terminate analysis; Correct the problem; Recalibrate or reprep batch (see Section 9.7).</td>
</tr>
<tr>
<td>CCV</td>
<td>Every 10 samples and at the end of the run.</td>
<td>80 - 120% recovery.</td>
<td>Terminate analysis; Correct the problem; Recalibrate and rerun all samples not bracketed by acceptable CCV or reprep batch (see Section 9.8).</td>
</tr>
<tr>
<td>CCB</td>
<td>Immediately following each CCV.</td>
<td>The result must be within +/- RL from zero.</td>
<td>Terminate analysis; Correct the problem; Recalibrate and rerun all samples not bracketed by acceptable CCB or reprep batch (see Section 9.8).</td>
</tr>
<tr>
<td>Method Blank</td>
<td>One per sample preparation batch of up to 20 samples.</td>
<td>The result must be less than or equal to the RL. Sample results greater than 20x the blank concentration are acceptable. Samples for which the contaminant is &lt; RL do not require redigestion (See Section 9.4).</td>
<td>Redigest and reanalyze samples. Note exceptions under criteria section. See Section 9.4 for additional requirements.</td>
</tr>
</tbody>
</table>

*See Sections 11.2.11 and 11.2.12 for exact run sequence to be followed.*
<table>
<thead>
<tr>
<th>QC PARAMETER</th>
<th>FREQUENCY</th>
<th>ACCEPTANCE CRITERIA</th>
<th>CORRECTIVE ACTION</th>
</tr>
</thead>
<tbody>
<tr>
<td>Laboratory Control Sample (LCS)</td>
<td>One per sample preparation batch of up to 20 samples.</td>
<td>Aqueous LCS must be within 80 - 120% recovery or in-house control limits.</td>
<td>Terminate analysis; Correct the problem; Redigest and reanalyze all samples associated with the LCS (see Section 9.5).</td>
</tr>
<tr>
<td>Matrix Spike</td>
<td>One per sample preparation batch of up to 20 samples.</td>
<td>75 - 125 % recovery or in-house control limits. If the MS/MSD is out for an analyte, it must be in control in the LCS.</td>
<td>In the absence of client specific requirements, flag the data; no flag required if the sample level is &gt; 4x the spike added. (see Section 9.6) For TCLP see Section 11.2.13</td>
</tr>
<tr>
<td>Matrix Spike Duplicate</td>
<td>See Matrix Spike</td>
<td>75 - 125 % recovery or in-house control limits; RPD ≤ 20%. (See MS)</td>
<td>See Corrective Action for Matrix Spike.</td>
</tr>
</tbody>
</table>
APPENDIX B
STL Hg DATA REVIEW CHECKLIST
# STL Hg Data Review Checklist

## Run/Project Information

Run Date: ___________  Analyst: ___________________  Instrument: ___________

Prep Batches Run: ___________________

Circle Methods used:
- 7470A / 245.1 : CORP-MT-0005 Rev 1
- CLP - AQC : CORP-MT-0006 Rev 0
- 7471 / 245.5 : CORP-MT-0007 Rev 1
- CLP - SOL : CORP-MT-0008 Rev 0

## Review Items

### A. Calibration/Instrument Run QC

<table>
<thead>
<tr>
<th></th>
<th>Yes</th>
<th>No</th>
<th>N/A</th>
<th>2nd Level</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. Instrument calibrated per manufacturer’s instructions and at SOP specified levels?</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>2. ICV/CCV analyzed at appropriate frequency and within control limits?</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>3. ICB/CCB analyzed at appropriate frequency and within +/- RL or +/- CRDL (CLP)?</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>4. CRA run (CLP only)?</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

### B. Sample Results

1. Were samples with concentrations > the high calibration standard diluted and reanalyzed?
2. All reported results bracketed by in control QC?
3. Sample analyses done within holding time?

### C. Preparation/Matrix QC

1. LCS done per prep batch and within QC limits?
2. Method blank done per prep batch and < RL or CRDL (CLP)?
3. MS run at required frequency and within limits?
4. MSD or DU run at required frequency and RPD within SOP limits?

### D. Other

1. Are all nonconformances documented appropriately?
2. Current IDL/MDL data on file?
3. Calculations and Transcriptions checked for error?
4. All client/ project specific requirements met?
5. Date of analysis verified as correct?

## Analyst: ___________________  Date: ___________________

## Comments:

__________________________________________________________________________

__________________________________________________________________________

__________________________________________________________________________

__________________________________________________________________________

2nd Level Reviewer: ___________________  Date: ___________________
APPENDIX C

MSA GUIDANCE
APPENDIX C. MSA GUIDANCE

Method of Standard Addition

Four equal volume aliquots of sample are measured and known amounts of standards are added to three aliquots. The fourth aliquot is the unknown and no standard is added to it. The concentration of standard added to the first aliquot should be 50% of the expected concentration. The concentration of standard added to the second aliquot should be 100% of the expected concentration and the concentration of standard added to the third aliquot should be 150% of the expected concentration. The volume of the unspiked and spiked aliquots should be the same (i.e., the volume of the spike added should be negligible in relation to the volume of sample).

To determine the concentration of analyte in the sample, the absorbance (or response) of each solution is determined and a linear regression performed. On the vertical axis the absorbance (or response) is plotted versus the concentrations of the standards on the horizontal axis using 0 as the concentration of the unspiked aliquot. An example plot is shown in Figure 1. When the resulting line is extrapolated back to zero absorbance, the point of intersection of the horizontal axis is the concentration of the unknown. Calculate the correlation coefficient (r) and the x-intercept (where y=0) of the curve. The concentration in the digestate is equal to the negative x-intercept.

Figure 1
For the method of standard additions to be correctly applied, the following limitations must be taken into consideration.

The plot of the sample and standards must be linear over the concentration range of concern. For best results, the slope of the curve should be similar to that of a plot of the aqueous standard curve.

The effect of the interference should not vary as the ratio of the standard added to the sample matrix changes.
APPENDIX D

TROUBLESHOOTING GUIDE
## APPENDIX D. TROUBLESHOOTING GUIDE

<table>
<thead>
<tr>
<th>Problem</th>
<th>Possible Cause</th>
</tr>
</thead>
</table>
| Poor or No Absorbance or Sensitivity Check failed | Incorrect wavelength  
Dirty windows  
Window loose  
Etched or dirty optics  
Wrong lamp  
Bad lamp  
Not enough or no sample introduced  
Empty sample cup  
Incorrectly made standards  
Gas leak  
EDL power supply set on “Continuous” |
| Erratic Readings                             | Source lamp not aligned properly  
Lamp not prewarmed  
Injection tip partially clogged  
Contaminated reagents  
Contaminated glassware  
Drying tube saturated  
Bad lamp  
Injection tip hitting outside of tube  
Injection tip coated or not set properly  
Leak in sample tubing  
Power fluctuations  
Air bubbles in tubing |
| EDL Won’t Light                             | Lamp cable not plugged in  
Lamp power set at 0  
Lamp is dead  
Power supply fuse is blown  
Short in cord |
| Standards reading twice or half normal absorbance or concentration | Incorrect standard used  
Incorrect dilution performed  
Dirty cell |
| Background Correction Light Blinking        | Background screen or attenuator faulty |
APPENDIX E

CONTAMINATION CONTROL GUIDELINES
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The following procedures are strongly recommended to prevent contamination:

All work areas used to prepare standards and spikes should be cleaned before and after each use.

All glassware should be washed with detergent and tap water and rinsed with 1:1 nitric acid followed by deionized water.

Proper laboratory housekeeping is essential in the reduction of contamination in the metals laboratory. All work areas must be kept scrupulously clean.

Powdered or Latex Gloves must not be used in the metals laboratory since the powder contains silica and zinc, as well as other metallic analytes. Only vinyl or nitrile gloves should be used in the metals laboratory.

Glassware should be periodically checked for cracks and etches and discarded if found. Etched glassware can cause cross contamination of any metallic analytes.

Autosampler trays should be covered to reduce the possibility of contamination. Trace levels of elements being analyzed in the samples can be easily contaminated by dust particles in the laboratory.

The following are helpful hints in the identification of the source of contaminants:

Reagents or standards can contain contaminants or be contaminated with the improper use of a pipette.

Improper cleaning of glassware can cause contamination.

Separate glassware if an unusually high sample is analyzed and soak with sulfuric acid prior to routine cleaning.
APPENDIX F

PREVENTIVE MAINTENANCE
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A maintenance log is used to record when maintenance is performed on instruments. When an instrument problem occurs indicate the date, time and instrument number, then identify the problem and corrective action in the maintenance log.

The following procedures are required to ensure that the instrument is fully operational.

**Cold Vapor Atomic Absorption (Leeman PS 200)**

<table>
<thead>
<tr>
<th>Daily</th>
<th>Semi-annually</th>
<th>Annually</th>
</tr>
</thead>
<tbody>
<tr>
<td>Clean lens.</td>
<td>Check Hg lamp intensity.</td>
<td>Change Hg lamp.</td>
</tr>
<tr>
<td>Check aperture.</td>
<td>Check argon flow.</td>
<td>Check liquid/gas separator.</td>
</tr>
<tr>
<td>Check tubing.</td>
<td>Check drain.</td>
<td></td>
</tr>
<tr>
<td>Replace drying tube.</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**Cold Vapor Atomic Absorption (PE 5000)**

<table>
<thead>
<tr>
<th>Daily</th>
<th>Monthly</th>
</tr>
</thead>
<tbody>
<tr>
<td>Clean aspirator by flushing with DI water.</td>
<td>Clean cell in aqua regia.</td>
</tr>
<tr>
<td>Check tubing and replace if needed.</td>
<td>Clean aspirator in aqua regia.</td>
</tr>
<tr>
<td>Clean windows with methanol.</td>
<td></td>
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<tr>
<td>Change silica gel in drying tube.</td>
<td></td>
</tr>
<tr>
<td>Check argon gas supply.</td>
<td></td>
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<tr>
<td>Adjust lamp.</td>
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