

UNITED STATES DEPARTMENT OF DEFENSE

# Data Validation Guidelines Module 2: Data Validation Procedure for Metals by ICP-OES

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Environmental Data Quality Workgroup

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# Data Validation Guidelines

## Module 2

JORDAN.BRIAN.D.1141739820 Digitally signed by  
JORDAN.BRIAN.D.1141739820  
Date: 2020.05.18 15:21:13 -05'00'

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Brian Jordan Date  
Army Principal

GILLETTE.JOHN.S.1123328350 Digitally signed by  
GILLETTE.JOHN.S.1123328350  
Date: 2020.05.18 10:25:08 -05'00'

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Seb Gillette, Ph.D. Date  
Air Force Principal

ADELSON.JORDAN.M.1268 Digitally signed by  
ADELSON.JORDAN.M.1268693137  
693137 Date: 2020.05.18 16:42:19 -04'00'

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Jordan Adelson, Ph.D. Date  
Navy Principal, EDQW Chair

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## Module 2: Data Validation Procedure for Metals by ICP-OES (SW-846 6010)

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### 1.0 Purpose

This document provides guidance on the validation of metals analyzed via Inductively Coupled Plasma Optical Emission Spectroscopy (ICP-OES) by SW-846 Method 6010. The objective of this procedure is to provide the end user with a clear understanding of the quality and limitations of the data through documented validation procedures and to encourage consistency in the validation and reporting for metals data generated for Department of Defense (DoD) projects. The users of this document should apply these data validation procedures to definitive data only.

Project Specific requirements as identified in the Quality Assurance Project Plan (QAPP) should always supersede the requirements of this document.

This document assumes the user is familiar with data validation conventions and qualifiers used in the *DoD General Data Validation Guidelines* (Rev. 1 2019). This document is also not intended to obviate the need for professional judgment during the validation process.

This document references the *Uniform Federal Policy for Quality Assurance Project Plans (UFP-QAPP) Optimized Worksheets (March 2012)*. Other QAPP formats are equally acceptable, as determined by the project team.

### 2.0 Procedure

#### 2.1 Introduction

**This document was written with primary consideration to SW-846 method 6010D with Quality Control (QC) criteria identified in the DoD Quality Systems Manual (QSM) Version 5.3.** However, some projects require other revisions such as method 6010B and 6010C. Validation should proceed using the acceptance criteria for the method version specified in the QAPP. Appendix A summarizes the QC checks and the required frequency and acceptance criteria for method 6010D and the QSM Version 5.3 requirements.

**Note: Since Method 6010D does not recommend low-level quantitative Mercury by this method, this module does not include the possible analysis of Mercury by ICP-OES.**

#### 2.2 Deliverables

Laboratory data deliverables consist of a combination of forms and raw data. The manner in which laboratories label their forms is not dictated nor specified. **The labeling convention below is used for simplicity and does not include all possible forms available.**

- Cover Sheet
- Case Narrative
- Sample Receipt and Conditions Summary
- Sample Results Summary
- Linear Range Check Summary
- Laboratory Control Sample Recovery Summary

- Matrix Spike/Matrix Spike Duplicate/Laboratory Duplicate Summary
- Method/Calibration Blank Summary
- Interference Checks Summary
- Initial Calibration Summary
- Initial/Continuing Calibration Verification Summary
- Low Level Calibration Check Standard Summary
- Serial Dilutions Summary
- Post Digestion Spike Recovery Summary
- Sequence and Digestion Logs

## 2.3 Validation Stages

The types of laboratory data deliverables, staged data validation, and the relationship between the two are outlined in the *DoD General Data Validation Guidelines*.

**Stage 1** data validation consists of a review of sample results forms, associated sample receipt summaries (chain of custody), and field QC data.

**Stages 2A and 2B** data validation consist of review of summary forms only.

**Stages 3 and 4** data validation require review of both summary forms and all associated raw data.

Both the laboratory deliverable requirements and the level of validation should be specified in the QAPP or other planning documents. Data review guidelines and how they apply to the different validation stages are indicated in the following sections.

Note: Any required stage of validation that reveals significant deviations from project requirements may require a higher stage of validation to uncover the source. Data validators are encouraged to communicate with their points of contact identified in the project QAPP (such as the UFP-QAPP Worksheet #6) to resolve discrepancies.

## 3.0 Stage 1 Validation

The following documents should be reviewed to ensure that the analytical method protocols outlined in the QAPP were performed (*representativeness*); to verify sampling and reporting *completeness*; to evaluate the performance of field blanks; and to verify compliance with *project sensitivity needs*:

- Cover Sheet
- Table of Contents
- Case Narrative
- Sample results form or equivalent Laboratory Report
- Chain of Custody (CoC) forms, Laboratory Receipt Checklists, and other supporting records
- Field QC forms and supporting records

Stage 1 is the validation of investigative and field QC samples.

### 3.1 Sample Results

Examine the Laboratory Report sample results (can also be called Form I) and verify the following information, ensuring that:

- Holding times have been met, as applicable.
- All sample identification labels are unique, and match the chain of custody.
- All project analytes identified in the QAPP and listed on the chain of custody have been analyzed and are reported.
- All laboratory reported Limits of Detection (LODs) and Limits of Quantitation (LOQs) are equal to or less than QAPP required LODs/LOQs (before adjustment for sample-specific conditions, such as dilution or moisture content).
- All project required LODs have been met and are lower than the LOQs.
- All project required Detection Limits (DLs) have been met and are lower than the LODs.
- All project required LOQs have been met and those LOQs are less than the project required action levels.
- All reported units (e.g., mg/kg) are accurate and reflect the requirements of the project and that units are consistent with the type of sample matrix.
- All required field QC samples (such as equipment blanks, and field duplicates) have been included in the Laboratory Report at the frequency specified in the QAPP.
- Soil samples have been reported on a dry weight basis, unless specified by the QAPP to report on a wet weight basis.
- Each laboratory report has a case narrative that explains all non-conformities with the data.

#### *Evaluation of the Laboratory Report*

Any samples received for analysis that were not analyzed should be noted in the data validation report, along with the reason(s) for failure to analyze the samples, if the reason(s) can be determined; conversely, samples that were analyzed by ICP but were not requested should also be noted.

Analytes that have project action levels less than the laboratory's LOQ may reveal a severe deficiency in the data and a failure to meet project goals, and should be noted in the data validation report. Analytes that have LODs or LOQs (before adjusting for sample-specific factors) that differ substantially from those presented in the project QAPP may also have an impact on the ability to meet the project goals and should be noted in the validation report. Errors in reported units and case narrative non-conformities that call into question the quality of the data should also be discussed in the validation report.

Errors in quantitation limits or missing and misidentified samples may require higher than Stage 1 validation. Data validators are encouraged reach out to their point of contact identified in the project QAPP (such as the UFP-QAPP Worksheet #6) when preparing the validation report.

For sample results (assuming no other qualifications due to data quality issues):

Qualification of data is based upon the reporting requirements of the project QAPP.

If the project QAPP changes reporting requirements from that specified in the QSM by reporting data down to the Detection Limit (DL), then any detects between the DL and LOQ are qualified as estimated **J**. Values below the DL are considered non-detects and are qualified as **U** at the stated DL.

If the project QAPP changes reporting requirements from that specified in the QSM by reporting data down to the Limit of Detection (LOD), then any detects between the LOD and below the LOQ are qualified as **J** estimated. Values below the LOD are considered non-detects and are qualified as **U** at the stated LOD.

If the project QAPP changes reporting requirements from that specified in the QSM by reporting data down to the Limit of Quantitation (LOQ), then any detects below the LOQ are considered non-detects and are qualified as **U** at the stated LOQ.

### **3.2 Chain of Custody (CoC)**

Examine the CoC form (some information may be included on Laboratory Receipt Checklists) for legibility and check that all ICP analyses requested on the CoC have been performed by the laboratory. Ensure all required analytes have been reported. Ensure that the CoC Sample Identification on the laboratory sample results form (Form I [or equivalent]) matches the Sample Identification on the CoC. Ensure the CoC was signed and dated during transfers of custody. Read the laboratory case narrative for additional information.

#### *Evaluation of the CoC*

Any discrepancies in sample naming between the CoC and sample results form should be noted in the data validation report with the correct sample name being identified in the report and on the annotated Form I (if applicable), if the correct sample name can be determined. These corrections should also be verified in any associated electronic data deliverables (EDDs).

If the receiving laboratory transferred the samples to another laboratory for analysis, both the original CoCs and transfer CoCs should be present. If the transfer CoCs are not present or if there is missing information (such as location of the laboratory), it should be documented in the data validation report. Make note in the validation report when signatures of 'relinquish' and 'receipt' of custody were not present.

#### **3.2.1 Sample Preservation**

Evaluate sample handling, transport, and laboratory receipt from the CoC and laboratory receipt checklists to ensure that the samples have been properly preserved and handled. The project QAPP (such as UFP-QAPP Worksheet #19) should provide specific preservation requirements. The following are general guidance if project specifications were not stipulated.

#### Metals by ICP-OES

- Metals samples should have been submitted in polyethylene or glass bottles (aqueous) or jars (soil).
- Aqueous samples should have been preserved with nitric acid to a pH  $\leq$  2. Glass bottles (preserved with acid) may not be appropriate if analyzing for Silica. Aqueous samples do not require thermal preservation.

- Soil samples do not require thermal preservation, although it is recommended to minimize loss of volatile compounds in the solid matrix containing metals of interest.
- Aqueous samples for dissolved metals analyses should have been filtered and then preserved on site prior to shipment to the laboratory. If this was not possible, the laboratory should have received an unpreserved aliquot to filter as soon as possible, prior to acid preservation and storage until digestion and analysis.

#### *Evaluation of Preservation*

If the pH of aqueous samples is > 2 upon receipt, the laboratory may add nitric acid to the samples upon receipt (samples should then be held at least 16 hours prior to extraction). If the samples are preserved in the laboratory, no data qualifiers should be applied, but the actions should be noted in the data validation report.

If the pH of aqueous samples is > 2 upon receipt, and the laboratory did not preserve the samples, detects should be qualified as estimated with a negative bias **J-** and non-detects as **X**, exclusion of data recommended.

If dissolved metals were required by the QAPP and the samples were not filtered, then qualify all detects as **X**, exclusion of data recommended. Non-detects do not require qualification, but lack of sample filtration should be noted in the case narrative.

#### **3.2.2 Holding Times**

Holding times for metals are measured from the time of collection (as shown on the CoC) to the start time of acid digestion or sample analysis (as applicable). Holding times for metals in aqueous and soil samples is 6 months.

Based on input from the DoD Environmental Data Quality Workgroup (EDQW) holding time exceedances are calculated as follows:

Total holding time is based on the time-frame (i.e., months for metals) of the requirement. The following example gives guidance on how hold time exceedances are measured:

For a test with a recommended maximum holding time measured in **months**, the holding time is tracked by the **month**.

- An exceedance of holding time for a sample with a 6 month holding time will occur when the start of the 7th month is reached. Therefore, a sample with a 6 month holding time collected at 8:30 AM on April 4th must begin analyzed or digested before 12:00 AM (midnight) on September 30th, or an exceedance has occurred (October 1st).

#### *Evaluation of Holding Times*

If the holding time is exceeded, qualify all associated detects as estimated **J-** and all associated non-detects as estimated **UJ** and document that holding times were exceeded.

If holding times for metals are grossly exceeded (defined as more than 30 days beyond the 6 month holding time), detects should be qualified as estimated **J-** and non-detects as **X**, exclusion of data recommended.



### 3.3 Field QC

Field QC can consist of various blanks, field duplicates, and field replicates. The purpose of blanks is to identify potential cross-contamination at different stages of sampling and cleaning of equipment for reuse. Duplicates and replicates help a project identify reproducibility among samples at the project site.

#### 3.3.1 Field Blanks

Not every field blank type may be utilized during any given sampling event and there may be more blank types than described in this document. Field blanks may be varied throughout the sampling events of a project. The types of blanks and their collection frequency should be stipulated in the QAPP. Generally, the blanks are collected once a day or one per twenty field investigative samples, by each sampling team, and may be matrix dependent.

Below are the common types of field blanks for metals analysis.

An **equipment blank** (also called a **rinse or rinsate blank**) is an aliquot of reagent water subjected to all aspects of sample collection (usually poured over or through the sample collection device). Analytes detected in equipment blanks indicate the possibility of cross-contamination between samples due to improper equipment decontamination. Equipment blanks are usually collected at a frequency of one per twenty investigative samples (per matrix per sampling technique), or as specified in the QAPP.

A **source blank** may be collected from each source of water used during each sampling event. This type of field blank may be analyzed to assess whether the chemical nature of the water used in decontamination may have affected the analytical results of site samples. A source blank is usually collected once per source prior to sample collection.

#### *Evaluation of Field Blanks*

Determine which field blanks apply to samples in the sample delivery group (SDG) from the CoC or any QC sample associative listing. If the applicability of multiple field blanks cannot be determined, communicate with the point of contact identified in the project QAPP to inquire if applicability can be determined.

Note: SDGs can be called by different names such as SEDD Lab Reporting Batch, depending on the project.

Ensure that units are correct when applying field blank qualifications.

Note: it may not be appropriate to make a direct quantitative comparison for aqueous field blanks (such as equipment blanks reported as  $\mu\text{g/mL}$ ) to a solid parent sample (such as a soil sample reported as  $\text{mg/kg}$ ). At best, only a qualitative comparison can be made.

Professional judgment should be applied to any equipment blank result that was associated with a contaminated source water blank. Generally, when multiple blank type contaminations are present, the evaluation should not involve a 'hierarchy' of one blank type over another. Each blank is evaluated separately and independently. The final validated result should be assessed on the blank with the highest value (i.e., greatest effect on sample analyte concentration). For example, if both a source water blank and an

equipment blank were collected in the same batch and the source water blank was also used as the equipment blank water (and both were found contaminated), the sample results would be qualified based on the blank with the higher contaminant concentration.

The source blank water should be analyte free (undetected; less than Detection Limit) and provided with the sample bottle kit by the contracted laboratory performing the analysis. To ensure the origin of the water used, consult with the field sampling team leader (as some sampling teams may obtain their source water separately from the laboratory) via appropriate channels identified in the QAPP (such as UFP-QAPP Worksheet #6).

If analytes (as appropriate) are detected in the field blanks, the procedure for the qualification of associated sample results is summarized below.

Compare the results of each type of field blank with the associated sample results. The reviewer should note that the blank analyses may not involve the same units or volumes as the associated samples. These factors should be taken into consideration when applying the 5x criteria discussed below, such that a comparison of the total amount of contamination is actually made. Care should be taken to account for any dilution factors when doing comparisons between detects in the sample and the blank.

If an analyte is detected in the field blank (at any concentration), but not in the associated samples, no action is taken.

If field blanks were not collected at the proper frequency required by the QAPP, then use professional judgment to qualify the data, and make note of this in the data validation report.

If an analyte is detected in the field blank (at any concentration) and in the associated samples, the action taken depends on both the blank and sample concentrations (Table I).

If a field blank has a negative blank result with an absolute value greater than the Detection Limit (DL) or Limit of Detection (LOD) as defined in the QAPP, then it should be evaluated against sample results (Table II).

**Table I: Blank Qualifications**

| Row Number | Blank       | Sample                |   |                      |
|------------|-------------|-----------------------|---|----------------------|
|            | Result      | Result                | Validated Result                          | Validation Qualifier |
| 1          | ≤ DL or LOD | ≤ DL or LOD           | Report as required by QAPP (at DL or LOD) | None                 |
| 2          | > DL or LOD | ≤ DL or LOD           | Report at DL or LOD                       | U                    |
| 3          | > DL or LOD | > DL or LOD but ≤ LOQ | Report at LOQ                             | U                    |
| 4          | > DL or LOD | > LOQ but ≤ 5x blank  | Report at Sample Result                   | J+                   |
| 5          | > DL or LOD | > LOQ and > 5x blank  | Report at Sample Result                   | None                 |

**LOD** = Limit of Detection **LOQ** = Limit of Quantitation **DL** = Detection Limit

**Note:** The laboratory B qualifier is maintained and the validation qualifier is added in addition to the laboratory qualifier. The QAPP should specify reporting at either the DL or LOD.

**Table II: Negative Blank Qualifications**

| Row Number | Negative Blank Result     | Sample Result            | Validation Qualifier(s) |
|------------|---------------------------|--------------------------|-------------------------|
| 1          | DL or LOD <  blank  ≤ LOQ | < DL or LOD              | UJ                      |
| 2          | DL or LOD <  blank  ≤ LOQ | ≥ DL or LOD but ≤ LOQ    | J-                      |
| 3          | DL or LOD <  blank  ≤ LOQ | > LOQ but ≤ 5x  blank    | J-                      |
| 4          | DL or LOD <  blank  ≤ LOQ | > LOQ and ≥ 5x  blank    | None                    |
| 5          | blank  > LOQ              | < DL or LOD              | X                       |
| 6          | blank  > LOQ              | ≥ DL or LOD but ≤ 5x LOQ | X                       |
| 7          | blank  > LOQ              | > 5x LOQ                 | J-                      |

**LOD** = Limit of Detection **LOQ** = Limit of Quantitation **DL** = Detection Limit

### 3.3.2 Field Duplicates (can also be called replicates)

Field duplicates consist of either collocated or subsampled (split) samples. Field duplicates for groundwater and surface water samples are generally considered to be collocated samples. Soil duplicate samples may be split samples or collocated, as specified in the QAPP. Field duplicate results are an indication of both field and laboratory precision; the results may be used to evaluate the consistency of sampling practices. Field duplicates do not measure total precision and they may not account for sample-to-sample variability.

#### *Evaluation of Field Duplicates*

Check to ensure that field duplicates were collected and analyzed as specified in the QAPP. If the sampling frequency is less than the frequency stated in the QAPP, no qualification of the associated sample results is necessary, but the incident should be discussed in the data validation report.

Relative Percent Differences (RPDs) should be calculated when detected results are reported for the duplicate(s) and at least one of the results is greater or equal to the LOQ. For field duplicate results, if the RPDs or absolute differences are greater than those stated in the QAPP, qualify the associated sample results as estimated **J**, and any non-conformities should be noted in the data validation summary.

Professional judgment may be required in instances where the sample and field duplicate results are less than the LOQ or project Reporting Limits (RLs). RPD results can be elevated when low (e.g., < 5x the LOQ) or estimated concentrations in the samples and duplicates are reported. If one or both results in a duplicate pair are < 5x the LOQ, the

absolute difference between the two results can be used as an alternative acceptance criterion, if approved by the QAPP or project point of contact. When comparing a detected result with a non-detected result, the LOD should be used as the nominal value of the non-detected result.

Some sampling schemes (such as Incremental Sampling Methodology (ISM) if used to collect metals soil samples) require specific replicate calculations, which should be specified in the QAPP.

It should be noted that RPDs or absolute differences for field duplicates are generally not calculated or reported by the laboratory, and should be calculated by the validator.

There are instances where an RPD is not calculable (for example, when one result is a non-detect and the other is greater than the LOQ). In those cases, the RPDs are not calculated but the non-conformity should be noted in the data validation report. The reported concentrations should be carefully examined to determine what conditions would permit one result to be reported at or above the LOQ/Reporting Limit (RL) and the other to be reported below the LOQ/RL or as a non-detect.

The equation for RPD calculations is given in Appendix B.

#### **4.0 Stage 2A Validation**

**Note: Stage 2A includes all of Stage 1**

Stage 2A requires the review and qualification of the following summary documents.

- Method Blank Summary
- Laboratory Control Sample/Laboratory Control Sample Duplicate
- Matrix Spike/Matrix Spike Duplicate or Laboratory Duplicate
- Dilution Test or Post Digestion Spike Summary
- Serial Dilution Summary

Stage 2A is the validation of preparation batch specific QC data in addition to any sample specific parameters included in Stage 1.

Generally, a “preparation batch” of samples consists of up to twenty field samples (maximum) along with a method blank, laboratory duplicate or matrix spike/matrix spike duplicate, and laboratory control sample. They are meant to be analyzed together on a single instrument. While these samples would ideally be analyzed together on a single instrument, laboratories may choose to split up a batch over multiple instruments to maximize efficiency. In such cases, the validation report should clearly differentiate between preparation or digestion batches and analytical batches or sequences when discussing the QC associated with the samples. The use of multiple instrumentation should be noted in the data validation report.

#### **4.1 Method Blanks**

A method blank is used to identify contamination originating in the laboratory that may have a detrimental effect on project sample results. The validator should identify samples associated with each method blank using a method blank summary form (or equivalent). Verify that the method blank has been reported per batch.

Compare the results of each method blank with the associated sample results. The reviewer should note that the blank analyses may not involve the same weights, volumes, percent moistures, or dilution factors as the associated samples.

These factors should be taken into consideration when applying the 5x criteria (discussed in section 3.3.1), such that a comparison of the total amount of contamination is actually made. Care should be taken to factor in the percent moisture or dilution factor when doing comparisons between detects in the sample and the method blank.

#### *Evaluation of Method Blanks*

If no method blank was analyzed, qualify detects in samples with no associated method blank as **X**, exclusion of data recommended. Non-detects do not require qualification.

If gross contamination exists (defined as greater than a Project Action Limit) in the method blank(s), all analytes affected in all associated samples in the preparation batch should be qualified **X** due to interference. This should be noted in the data validation comments. An exception is that any sample result that is  $\geq 10x$  the concentration detected in a grossly contaminated method blank should be qualified as estimated **J**.

If an analyte is detected in the method blank, but not in the associated samples, no action is taken.

If an analyte is detected in the method blank and in the associated samples, the action taken depends on both the blank and sample concentrations. Tables I and II (Blank and Negative Blank Qualifications) and section 3.3.1 discussions on evaluations of results from the LOD to LOQ is also applicable to the method blank.

Additionally, there may be instances where little or no contamination was present in the associated method blanks, but qualification of the sample was deemed necessary. Contamination introduced through dilution water is one example. Although it is not always possible to determine, instances of this occurring can be detected when contaminants are found in the diluted sample result but are absent in the undiluted sample result. It may be impossible to verify this source of contamination. However, if the reviewer determines that the contamination is from a source other than the sample, the data should be qualified. In this case, the 5x rule does not apply. The sample value should be reported as a non-detect and the reason should be documented in the data validation report.

Multiple blank contaminations (such as a batch with field blanks and a method blank) does not establish a 'hierarchy' of one blank over another. Each blank must be evaluated individually. Blanks should not be qualified due to the results of other blanks.

#### **4.2 Laboratory Control Sample/Laboratory Control Sample Duplicate (LCS/LCSD)**

An LCS is an analyte free sample matrix spiked with known amounts of the analytes of interest and taken through all sample preparation, cleanup and analytical steps. LCSs establish the method precision and bias for a specific batch of samples. Analysis of LCSDs may be required by the QAPP, or may be used as an indication of batch precision in instances where matrix spike duplicate analysis is not possible (e.g., a limited volume of sample).

LCS and, if analyzed, LCSD recoveries should be within the QC limits specified in the QAPP or as listed in the QSM. If an LCSD was analyzed, the RPDs should be within the QC limits specified in the QAPP or as listed in the QSM.

An LCS is analyzed in every preparation batch of 20 environmental samples.

#### *Evaluation of LCS/LCSD*

Verify that an LCS was analyzed with each batch of samples.

Verify that results (from appropriate summary form), percent recoveries, RPDs (if applicable) and acceptance limits were reported for all target analytes.

If the spike percent recovery control criteria displayed in the deliverable are not the same range (i.e., outside or wider than) as those stipulated in the QAPP or the DoD QSM, reference the required control ranges for evaluation instead of the summarized ranges in the deliverable. The project team should be informed to implement changes to the current deliverables or those to be created in the future.

In-house control limits are acceptable for any analytes not specified in the QAPP or DoD QSM. No qualification is necessary for any reported in-house control limit that is within its control range.

If the LCS percent recoveries were greater than the upper control limit, qualify detects for the analyte in associated samples as estimated with a positive bias **J+**. Non-detects should not be qualified.

If the LCS percent recovery is less than the lower control limit but  $\geq 60\%$ , qualify associated detects as estimated with a negative bias **J-** and associated non-detects as **UJ**.

If the LCS percent recovery is  $< 60\%$ , qualify associated detects as estimated with a negative bias **J-** and associated non-detects as **X**, exclusion of data recommended.

If the LCS/LCSD was not spiked with all target analytes, notify the project team by following the notification protocols outlined in the QAPP (such as UFP-QAPP Worksheet #6) and qualify all detects and non-detects for those analytes not spiked as **X**, exclusion of data recommended.

If the LCS/LCSD RPDs were greater than the acceptance limits, qualify detects for the analyte in the associated sample(s) as estimated **J**. Non-detects should not be qualified.

If the project QAPP requires the use of a Standard Reference Material (SRM), the control limits specified by the SRM should be the basis for decisions regarding qualification of data. If the SRM falls outside of the specified control limits, the LCS evaluations as listed above should be used for the SRM, as applicable.

Professional judgment should be utilized in qualifying data for circumstances other than those listed above.

### **4.3 Matrix Spike/Matrix Spike Duplicate (MS/MSD) and Laboratory Duplicate (LD)**

MS/MSD and LD (sometimes referred to as a Sample Duplicate or Matrix Duplicate that is not spiked) data are used to determine the effect of the matrix on a method's recovery efficiency and precision for a specific sample matrix. LD analyses are also performed to demonstrate acceptable method precision by the laboratory at the time of analysis. If the project QAPP does not specify a statistical sampling design, each preparatory batch should have one site specific MS and either a LD or MSD. For sample designs that rely on Incremental Sampling Methodology (ISM), three or more replicates may be specified by the project QAPP.

Field blanks should not be used as the parent sample for the MS/MSD or LD analyses.

Note: If a field blank was used for the MS/MSD or LD, the information must be included in the data validation report, but the data should not be qualified. Sample matrix effects should not be observed with field blanks; therefore, no site-specific matrix effects can be determined from a field blank.

The MS and MSD should be spiked per QSM requirements with all target analytes. If the parent sample for the MS/MSD was from another site or project (for example, not enough sample collected, or multiple site samples analyzed within a single batch), the reason should be documented in the data validation report, and sample results should not be qualified due to any non-conformities noted in non-site-specific matrices.

#### *Evaluation of MS/MSD and LD*

Verify that MS/MSD analyses were performed at the specified frequency.

Verify that the MS/MSD were spiked with all target analytes, and that percent recoveries and RPDs were reported for all target analytes. If the MS/MSD was not spiked with all target analytes, notify the project team by following the notification protocols and qualify all detects and non-detects in the parent sample for those analytes not spiked as **X**, exclusion of data recommended.

If the parent sample concentration was > 4x the spike concentration, the MS and MSD percent recovery criteria do not apply. This should be noted in the data validation report.

If the MS/MSD or LD results do not meet the technical criteria, apply the action to all samples in the same preparation batch of the same matrix, if the samples are considered sufficiently similar. Exercise professional judgment in determining sample similarity when making use of all available data, including: samples of the same matrix from the same project site with similar analyte concentrations; site and sampling documentation (e.g., location and type of sample, descriptive data, soil classification); field test data; and laboratory data for other parameters. If no samples in the SDG are sufficiently similar to the parent sample, only the parent sample should be qualified. This should be noted in the data validation report.

Compare the RPD for each analyte with LCS control limits established by the QAPP or DoD QSM. If the spike percent recovery control criteria displayed in the deliverable are not the same range (i.e., outside or wider than) as those or stipulated in the QAPP or the DoD QSM, reference the required control ranges for evaluation instead of the summarized ranges in the deliverable. The project team should be informed to implement changes to the



current deliverables or those to be created in the future. Follow the notification protocols outlined in the QAPP (such as UFP-QAPP Worksheet #6).

For ISM samples, the RPDs of the replicates should be evaluated against the criteria established in the project QAPP. If the RPDs are greater than the acceptance limits, qualify detects for the analyte in the associated sample(s) as estimated **J**. Non-detects should be qualified as estimated **UJ**.

If the MS or MSD percent recoveries were greater than the upper control limit, qualify detects for the analyte in the associated parent sample as estimated **J+**. Non-detects should not be qualified.

If the MS or MSD percent recoveries were less than the lower acceptance limit but  $\geq 30\%$ , qualify detects for the analyte in the associated parent sample as estimated **J-** and non-detects as estimated **UJ**.

If the percent recoveries were  $< 30\%$ , qualify detects for the analyte in the associated parent sample as estimated **J-** and non-detects as **X**, exclusion data recommended.

If the MS/MSD or parent sample result/LD pair RPDs were greater than the acceptance limits ( $\leq 20\%$  per QSM), qualify detects for the analyte in the associated sample(s) as estimated **J**. Non-detects should be qualified as estimated **UJ**.

MS/MSD exceedances due to the presence of a target analyte in the parent sample at  $> 4x$  the spike concentration should not necessarily result in any qualifications. If a Post Digestion Spike (PDS) or Dilution Test (DT) was performed and can be evaluated for the analyte in question, use the PDS or DT results and apply professional judgment to determine if the associated data should be qualified. Document which target analytes were affected and the application of professional judgment in the data validation report.

Note: The above qualifiers assume no PDS or DT was performed for failed MS/MSD control limits. If a PDS or DT was performed, see section 4.4 for possible qualifiers.

#### **4.4 Dilution Test (DT) and Post Digestion Spike (PDS)**

If a MS or MSD failed the percent recovery acceptance limits for a target metal, a PDS or DT, also known as a serial dilution (SD), should be performed, preferably using the same sample used for MS/MSD analysis. This is the case even when the MS/MSD failed due to the parent sample being  $\geq 4x$  the spiking concentration.

A DT is performed when a MS or MSD fails the percent recovery acceptance limits and a target analyte concentration is within the calibration range of the instrument and is considered sufficiently high (minimally, a factor of 25x greater than the Lower Limit of Quantitation (LLOQ) for Method 6010D or 50x greater than the LOQ for the QSM). The DT is an analysis of a 5x dilution that should agree to within  $\pm 20\%$  of the original target analyte concentration ( $\pm 10\%$  for the QSM). If not, then a chemical or physical interference effect should be suspected. The MS is often a good choice of sample for the DT, since reasonable concentrations of the target analytes are present.

A PDS test is used when a MS fails control acceptance limits established by the QAPP or QSM. The test only needs to be performed for the specific target analytes that failed the original MS limits, and only if the spike concentration added was greater than the

concentration determined in the unspiked sample (Method 6010D). The spike addition should be based on the original concentration of each target analyte of interest in the sample. The recovery of the post-digestion MS should fall within a  $\pm 25\%$  (75-125%) acceptance range (or 80-120% if QSM requirements are applied), relative to the known true value.

#### *Evaluation of DT and PDS*

If a DT or PDS was not analyzed due to a MS or MSD failure, notify the project team by following the notification protocols outlined in the QAPP (such as UFP-QAPP Worksheet #6) and qualify all detects and non-detects for those failed analyte(s) as **X**, exclusion data recommended.

If the target analyte(s) had a sufficiently high concentration that a DT was run, then no qualification is required if the DT passes acceptance criteria. Make note of the DT in the data validation report.

If the DT exceeds  $\pm 20\%$  ( $\pm 10\%$  for QSM) then qualify the affected target analyte(s) in the parent sample as estimated **J**. Non-detects should not be qualified.

If a PDS was performed on a MS that failed acceptable limits:

If the MS percent recoveries failed low ( $<30\%$ ) but the PDS was acceptable (within  $\pm 25\%$ ) then qualify detects in the parent sample as estimated **J** and non-detects as estimated **UJ**. If the PDS also fails, then qualify detects as estimated **J-** and non-detects as **X**, exclusion of data recommended.

If the MS percent recoveries fail low but still are  $\geq 30\%$ , and the PDS was acceptable (within  $\pm 25\%$ ) then qualify detects in the parent sample as estimated **J** and non-detects as estimated **UJ**. If the PDS also fails, then qualify detects as estimated **J-** and the non-detects as estimated **UJ**.

If the MS percent recoveries fail high, and the PDS was acceptable (within  $\pm 25\%$ ), then qualify detects in the parent sample as estimated **J** and non-detects do not require qualification. If the PDS also fails, then qualify detects as estimated **J+** and the non-detects do not require qualification.

#### **4.4.1 Method of Standard Addition (MSA)**

A technique that is sometimes employed with known complex matrices is called the Method of Standard Addition (MSA). It is applied when a sample matrix cannot adequately match the standards matrix used in the initial calibration. The MSA helps resolve a sample matrix that may be enhancing or depressing an analyte signal, and thus providing information that can properly characterize a sample result.

The MSA is a technique that is not routinely used by a laboratory. The use of MSA should be specifically outlined in the project QAPP.

### *Evaluation of MSA*

If a MSA was required by a project QAPP and was not performed by the laboratory, notify the project team by following the notification protocols outlined in the QAPP (such as UFP-QAPP Worksheet #6) and qualify all detects and non-detects for those MSA metal analyte(s) as **X**, exclusion of data recommended.

The MSA technique helps compensate for any analyte enhancement/suppression due to matrix affects. As such, when properly utilized MSA data should not require any qualification. In those cases where MSA has been applied incorrectly, qualify all detects and non-detects for those MSA metal analyte(s) as **X**. In all cases, the use of MSA should be noted in the case narrative.

### **4.5 Sample Dilutions and Reanalysis**

Laboratories may dilute samples due to high analyte concentrations or reanalyze samples due to quality control non-conformities, and document both sets of results. Generally, the laboratory will only report one value for a given analyte in the official laboratory report (or equivalent form). In these instances, if the results of multiple analyses are reported for the same analyte for a sample, the validator should evaluate all available results to determine which is the appropriate final result. The validator should consider the application of appropriate qualifiers to the reported results within the scope of the project due to elevated LODs/LOQs or other quality control non-conformities. Qualifiers apply only to the reported results in the official laboratory report.

#### *Evaluation of Sample Dilutions and Reanalysis*

Results reported from dilutions leads to elevated LODs for non-detects. The validation report should indicate the reason for all reported dilutions (including cases where the laboratory did not perform an undiluted analysis) resulting in elevated sensitivity limits for non-detected results.

When reanalysis has occurred due to quality control non-conformities, the validator should ensure that the non-conformity was corrected during the reanalysis. If that is not the case, then the appropriate qualifier should be placed on the reported results.

In some cases, using professional judgment, the validator may determine that an alternate result was more appropriate than the one reported. In those cases, explain the rationale for accepting the alternate result in the data validation report.

In some cases, reanalysis may lead to exceedances of holding time. Use professional judgment to evaluate the results and apply the appropriate qualifiers (if required).

### **5.0 Stage 2B Validation**

**Note: Stage 2B includes all of Stage 1, and Stage 2A**

Stage 2B requires the review and qualification of the following summary documents.

- Sequence and preparation logs (including any Instrument Blanks)
- Initial Calibration Summary (any equivalent to include Initial Calibration, Linear Regression or RSE)
- Initial/Continuing Calibration Verification Summary (any equivalent to include Initial and Continuing Calibration Verifications)
- Initial/Continuing Calibration Blank Summary
- Low Level Calibration Verification Summary
- Interference Check Summary (any equivalent to include ICS or SIC)
- Internal Standard Summary (optional)

Stage 2B is the validation of instrument specific QC data.

### **5.1 Sequence and Preparation Logs**

Sequence logs are reviewed by the data validator to ensure all QC samples (both batch and instrument specific) had been analyzed within a specific batch, in the correct order. Preparation logs are reviewed by the data validator to ensure that samples had the proper extraction performed, within specified holding times. The logs themselves do not require validation. However, non-conformities uncovered in the review of the logs may point the validator to specific samples that require further review. Non-conformities uncovered in preparation or sequence logs should be noted in the data validation report.

Sequence logs are helpful in identifying when multiple instrumentation is used to analyze a batch of samples. For example, it is not uncommon to analyze a single batch of twenty samples at the same time on two or more different instruments. At a minimum, each instrument must be calibrated independently. Batch QC should be reviewed on each instrument, as appropriate. Non-conformities involving the use of multiple instruments should be noted in the data validation report.

Sequence logs are also helpful in identifying excess use of instrument blanks. A common problem is the use of multiple instrument blanks to 'clean' OES in order to achieve acceptable QC results but allowing sample throughput without benefit of blank runs. Such non-conformities should be noted in the validation report and associated data should be qualified as **X**.

### **5.2 Initial Calibration**

The objective of initial calibration is to ensure that the instrument is capable of producing acceptable qualitative and quantitative data. Initial calibration demonstrates that the instrument is capable of acceptable performance in the beginning of the analytical run and of producing an acceptable calibration curve.

OES instruments should be calibrated for all target analytes plus any required for interference correction. Single standard and multi-point calibration curves are both acceptable.

OES instruments must be calibrated daily, each time the instrument is set up, and after calibration verification failure.

Method 6010D: A calibration blank and at least one standard must be used in establishing the analytical (calibration) curve. If a multi-point curve is used at least three standards must be employed, and one of the standards must be at or below the Lower Limit of Quantitation (LLOQ).

The QSM requires a minimum of one high standard and a calibration blank. If a multi-point curve is used, at least three standards must be employed. The LOQ must lie at (lowest calibration point) or within a multi-point calibration curve.

Linearity for a multi-point curve is determined using linear regression analysis, inversely weighted linear regression analysis, or Relative Standard Error (RSE).

The multi-point curve correlation coefficient ( $r$ ) must be  $\geq 0.995$ , the coefficient of determination ( $r^2$ ) must be  $\geq 0.99$ , or the RSE must be  $\leq 20\%$ .

Note: Method 6010D establishes the Lower Limit of Quantitation (LLOQ) as spiked replicates at the lowest point of a multi-point calibration curve. The DoD QSM establishes the Limit of Quantitation (LOQ) as the lowest calibration point or within the calibration curve. If employing calibration with a calibration blank and a single high standard, the LOQ or LLOQ is determined by analyzing a series of low standards that establish precision and bias at the stated LOQ/LLOQ. The QAPP or other planning documents should verify which reporting convention is acceptable for the project.

#### *Evaluation of Initial Calibration*

If target analytes were not calibrated, qualify associated detects and non-detects as **X**, exclusion of data recommended. Samples should not have been run without a valid calibration in accordance with the DoD QSM.

Any manipulation of calibration points (such as 'dropping' calibration levels at the ends of a multi-point calibration curve) to achieve acceptance criteria should have a technical justification documented in the laboratory report. Use professional judgment to evaluate the data. If no technical justification is provided, then make note of this in the data validation report and qualify the associated data as **X**, exclusion of data recommended.

If Method 6010D is applied, the lowest calibration standard can establish the LLOQ. Detects below the LLOQ are qualified as estimated **J**.

If the LOQ (QSM criteria applied) is not the lowest calibration standard within the calibration curve (but is still within the calibration curve), then any detects that fall between the LOQ and the lowest calibration standard should be qualified as estimated **J**. Detects below the lowest calibration standard are also qualified as estimated **J**.

The lowest multi-point calibration standard should be at or below the LOQ (If QSM requirements are applied). If the LOQ is below the lowest calibration standard, then the LOQ has been reported in a manner that is inconsistent with QSM requirements. Qualify all associated detects as **X** and make note of this in the data validation report.

For multi-point curves:

If an insufficient number of calibration standards was used, qualify all associated detects estimated **J** and all associated non-detects estimated **UJ**.

If the  $r$  value is  $< 0.995$ ; the  $r^2$  value is  $< 0.99$ ; or the RSE is  $> 20\%$ , qualify all associated detects as estimated **J** and all non-detects as estimated **UJ**.

If the acceptance criteria are grossly exceeded (defined as  $r$  value  $< 0.95$ ;  $r^2$  value  $< 0.90$ ; or RSE  $> 30\%$ ) qualify associated detects as estimated **J** and non-detects as **X**, exclusion of data recommended.

For metals analysis, all target analyte results that are within the linear range but above the calibration range should first be diluted to within the calibration range before they are reported. As an alternative, a high level check standard to verify the linear range (within  $\pm 10\%$ ) can be used. If sample results are reported above the calibration range without dilution or analysis of a linear range check standard, qualify all detects as **X**, and make note of the lack of dilution(s) or linear range check standard(s) in the data validation report. Non-detects do not require qualification.

If dilution(s) were performed that were within the calibration range of the initial calibration, then qualification of the data is not necessary. Make note in the data validation report that dilution(s) were performed.

If linear range check standard(s) were within criteria, then qualification of the data is not necessary. Make note in the data validation report that linear range check standards were used.

When sample results are reported at more than one dilution due to analyte concentrations exceeding the calibration curve, the lowest LODs are generally used for the non-detects unless a QC criterion has been exceeded.

### **5.3 Initial (Secondary Source) and Continuing Calibration Verification**

The initial calibration curve should be verified with a standard that has been purchased or prepared from an independent source each time initial calibration is performed. This standard is called the secondary source or Initial Calibration Verification (ICV). The ICV should contain all of the metals that are reported. Note that multiple ICVs may be analyzed to encompass all of the target metals.

The ICV is associated with all sample results in the analytical batch. The percent recovery for each target analyte in the ICV must recover within  $\pm 10\%$  (90-110%).

Continuing calibration verification (CCVs) checks may be standards from the same source as the calibration standards. CCVs must be analyzed after every 10 field samples and at the end of the analytical run and must contain all target analytes. Field sample results are associated with bracketing CCVs, the CCV that ran before and the CCV that ran after each 10 field samples. The percent recovery for each target analyte in the CCV must recover within  $\pm 10\%$  (90-110%).

### *Evaluating the ICV and CCV*

Verify that the ICV was analyzed immediately following the initial calibration and contained all target analytes. Verify the CCVs were analyzed at the proper frequency (after every 10 field samples and at the end of the analytical run). Verify that the percent recoveries for all target analytes in both the ICV and bracketing CCVs were within 90-110% of the true values.

If the ICV (second source) has not been performed after an initial calibration or if samples have been analyzed prior to a valid ICV, qualify all associated data as **X**, exclusion of data recommended. No samples should have been analyzed in accordance with QSM requirements.

If the CCV has not been analyzed (either continuing or end-of-run), qualify all associated data as **X**, exclusion of data recommended. No samples should have been analyzed without a valid CCV.

If any ICV percent recovery was > 110% or < 90% qualify all associated detects and non-detects as **X**, exclusion of data recommended.

If the CCV percent recovery was > 110% or < 90% qualify all associated detects and non-detects since the last acceptable CCV as **X**, exclusion of data recommended. This includes the end-of-run CCV.

### **5.4 Initial Calibration and Continuing Calibration Blank (ICB/CCB)**

Initial and Continuing Calibration Blanks (ICB and CCBs) are assessed to determine the existence and magnitude of contamination problems associated with sample extraction and analysis. If problems with any blank exist, all associated data must be carefully evaluated to determine whether there is any bias associated with the data, or if the problem is an isolated occurrence not affecting other data.

An ICB should be analyzed immediately after the ICV. ICBs apply to all samples in the associated analytical run. A CCB must be analyzed immediately after each CCV. The CCV/CCB analysis is considered a set. CCB non-conformances apply only to samples bracketed by the CCB. Each sample must have an associated ICB and bracketing CCBs.

ICB or CCB results indicate instrument-level contamination and should be compared to the raw values of the samples, if available. Since the raw data is usually not available in Stage 2B, the ICB and CCB detects should be converted to the reporting units for comparison, including percent moisture and dilutions in the conversion as applicable.

The laboratory acceptance criteria in the DoD QSM varies depending on the amount of contamination in the blank (and the associated sample result) and the type of corrective action required. For validation, all detects and negative results in the blanks are evaluated against the sample results.

#### *Evaluation of ICB and CCB*

Verify that ICB and CCBs were analyzed at the required frequencies.

The criteria outlined in Section 3.3.1 (Field Blanks) and summarized in Table I (Blank Qualifications) and Table II (Negative Blank Qualifications) are also applicable to ICB and CCB evaluations.

If the ICB was not analyzed, then all associated data (detects and non-detects) should be qualified as **X**, exclusion of data recommended. The QSM does not allow data to be reported without a valid ICB.

For ICBs that are qualified based on Table I or Table II, apply the action to all associated samples reported from the analytical sequence.

For CCBs that were not analyzed at the required frequencies, then all associated data that was not bracketed by valid CCBs should be qualified as **X**, exclusion of data recommended. The QSM does not allow data to be reported without valid CCBs that bracket all the samples.

For CCBs that are qualified based on Table I or Table II, apply the action to all associated samples analyzed between a previous acceptable analysis of the CCB and a subsequent acceptable analysis of the CCB in the analytical sequence.

Negative blank results with absolute values greater than the LOD should also be evaluated against sample results.

If an analyte is detected in a blank but the associated sample results are non-detects, the results should not be qualified.

If an analyte is detected in a blank and the associated sample results are > 5x the associated blank result(s) or non-detect, no data qualification is required.

### **5.5 Low Level Calibration Check Verification (LLCCV)**

The accuracy at the low end of the initial calibration curve must be verified with a standard or readback verification (containing all target analytes) at the LLOQ for Method 6010D (less than or equal to the LOQ for QSM). If a multiple-point calibration was performed, the lowest concentration standard may be re-quantitated against the calibration curve.

Alternately, if a single-point calibration was performed, a separate standard containing all target analytes must be analyzed prior to sample analysis. The LLCCV is analyzed daily, after calibration, and the percent recovery for each analyte in the LLCCV should be within  $\pm 20\%$  (80-120%) of the true value.

For Method 6010D requirements, a daily readback verification or standard is run at the mid-point of the Linear Range or middle calibration point for a multipoint calibration. All reported analytes should be within  $\pm 10\%$  of true value. The ICV/CCV (and qualification criteria listed in section 5.3) can be used as the mid-point verification.

#### *Evaluation of the LLCCV*

Verify that the lowest concentration standard was re-quantified against the appropriate calibration curve or that a separate standard containing all analytes at the LLOQ (less than or equal to the LOQ for QSM) was analyzed prior to sample analysis.



Verify that the percent recovery for each target analyte was 80-120%.

If a LLCCV was not analyzed, qualify associated detects as estimated **J** and non-detects as **X**, exclusion of the data recommended.

If the LLCCV percent recovery was > 120%, qualify associated detects less than the ICV or CCV concentration (whichever is lower) as estimated **J+**. Detects greater than the ICV or CCV concentration and non-detects should not be qualified if the ICV or CCV was acceptable.

If the LLCCV percent recovery was < 80%, but  $\geq$  50%, qualify associated detects less than the ICV or CCV concentration (whichever is lower) as estimated **J-** and non-detects as estimated **UJ**. Detects greater than the ICV or CCV concentration should not be qualified if the ICV or CCV was acceptable.

If the LLCCV percent recovery was < 50%, qualify associated detects less than the ICV or CCV concentration (whichever is lower) as estimated **J-** and non-detects as **X**, exclusion of data recommended. Detects greater than the ICV or CCV concentration should not be qualified if the ICV or CCV was acceptable.

## 5.6 Interference Check Solutions (ICS) or Spectral Interference Checks (SIC)

The Interference Check Solutions (ICs) or Spectral Interference Checks (SICs) verify that interference levels are corrected by the data system within appropriate limits. ICS or SIC analyses must be run after the initial calibration and prior to sample analysis.

- SIC applies to method 6010D
- ICS applies to previous versions of Method 6010 and QSM Version 5.3 requirements.

Note: The project QAPP should specify which criteria (Method 6010D or QSM Version 5.3) is applicable for the data that is being validated.

ICS analyses consist of the evaluation of two solutions. One solution (ICS-A) is composed of relatively high concentrations of common interfering analytes only and is evaluated to determine the effect of interferences below the calibration range (< 1/2 LOQ). The other solution (ICS-AB) is composed of the same high concentrations of interfering analytes and spiked with known concentrations of the target analytes. It is evaluated to determine the effect of interferences on detects within the working range of the instruments.

ICS-A and ICS-AB are analyzed daily after initial calibration, prior to sample analysis.

ICS-A: Absolute value of concentration for all non-spiked project analytes < 1/2 LOQ (unless there is a noted trace impurity on the Manufacturer's certificate of analysis from one of the spiked analytes);

ICS-AB: Within  $\pm$  20% of true value (not needed if instrument can read negative responses).

For SIC in Method 6010D, there are two types of solutions that are used. Individual element SIC solutions are performed when the instrument is initially setup, and periodically (at least once every 6 months) thereafter. The mixed element SIC solution is used daily to check

that the instrument is free from interference from elements typically observed in high concentrations and to check that any interference corrections applied are still valid. The SIC solutions must be used regardless of whether or not interelement corrections are applied. They evaluate both potential spectral interferences and the accuracy of any correction equations.

The daily mixed element SIC solution is used as an ongoing daily check of freedom from spectral interferences. The mixed element SIC solution is analyzed at least once per day, immediately after the initial calibration. The concentration measured for any unspiked target analytes must be less than  $\pm$  the LLOQ. If this criterion is not met then sample analysis may not proceed until the problem is corrected, or alternatively, the LLOQ may be raised to twice the concentration observed in the SIC solution.

#### *Evaluation of Daily ICS or SIC*

Verify that ICS-A, ICS-AB or SIC analyses were performed at the correct frequency for each sequence. For an interference check that does not meet criteria, apply the action to all sample results reported from the analytical sequence.

ICS and SIC analyses only apply to samples with interferent concentrations which are comparable to (within 10% of the concentration) or greater than their respective levels.

Results for the interfering analytes in the ICS-A, ICS-AB and SIC must fall within 20% of the true value. Results for the spiked target analytes in ICS-AB must fall within the control limits of  $\pm$  20% of the true value.

Results for the unspiked target analytes in ICS-A must be  $< 1/2$  LOQ (or less than  $\pm$  the LLOQ for the SIC). Some target analytes may be present as contaminants from the solution components rather than interferences. If it can be demonstrated that an analyte detect is the result of contamination via analysis by another analytical method or from the manufacturer's certificate of analysis, the result for that analyte must be the less than the LOD after the known contaminant true value is subtracted.

For samples with concentrations of interfering analytes which are comparable to (within 10% of the concentration) or greater than their respective levels in the interference check, recommended actions for ICS or SIC are summarized below.

If ICS or SIC analyses were not performed, qualify all applicable associated sample results as **X**, exclusion of data recommended.

If the result for an analyte not present in the ICS is  $\geq 1/2$  LOQ (greater than or equal to true value + the LLOQ for the SIC), qualify detects  $< 5x$  the unspiked element's concentration as estimated **J+**. Non-detects should not be qualified. Detects  $\geq 5x$  the unspiked element's concentration should not be qualified.

If the result for an analyte not present in the ICS is greater than or equal to the LOQ (greater than or equal to true value +  $2x$  the LLOQ for the SIC), this is considered a gross interelement correction failure. Qualify detects  $< 5x$  the unspiked element's concentration as **X**, exclusion of data recommended. Non-detects should not be qualified. Detects  $\geq 5x$  the unspiked element's concentration should be qualified as estimated **J+**.

If the result for an analyte not present in the ICS is negative with an absolute value  $\geq 1/2$  LOQ (greater than or equal to true value + the LLOQ for the SIC), qualify detects  $< 5x$  the unspiked element's concentration as estimated **J-** and non-detects as estimated **UJ**. Detects  $\geq 5x$  the unspiked element's concentration should not be qualified.

If the result for an analyte not present in the ICS is negative with an absolute value greater than or equal to the LOQ (greater than or equal to true value +  $2x$  the LLOQ for the SIC), this is considered a gross interelement correction failure. Qualify detects  $< 5x$  the unspiked element's concentration as estimated **J-** and non-detects as **X**, exclusion of data recommended. Detects  $\geq 5x$  the unspiked element's concentration should not be qualified.

If a ICS or SIC displays recovery for an analyte  $> 120\%$ , qualify positive sample results as estimated **J+**. Non- detects are not qualified.

If a ICS or SIC displays recovery for an analyte  $< 80\%$ , but  $\geq 50\%$ , qualify positive sample results as estimated **J-** and non-detects as estimated **UJ**.

If a ICS or SIC displays recovery results for an analyte are  $< 50\%$ , qualify positive sample results as estimated **J-** and non-detects as **X**, exclusion of data recommended.

The following Table III summarizes the evaluation of ICS or SIC analyses.

**Table III: ICS/SIC Qualifications**

| <b>Target Analyte</b>             | <b>Sample Result</b>      | <b>Qualifier(s)</b> |
|-----------------------------------|---------------------------|---------------------|
| %R > 120%                         | Non-detect                | None                |
| %R > 120%                         | Detect                    | J+                  |
| %R ≥ 50% < 80%                    | Non-detect                | UJ                  |
| %R ≥ 50% < 80%                    | Detect                    | J-                  |
| %R < 50%                          | Non-detect                | X                   |
| %R < 50%                          | Detect                    | J-                  |
| <b>Positive Unspiked Element</b>  |                           |                     |
| > LLOQ or ½ LOQ                   | Non-detect                | None                |
| > LLOQ or ½ LOQ                   | Detect < 5x<br>[unspiked] | J+                  |
| > LLOQ or ½ LOQ                   | Detect ≥ 5x<br>[unspiked] | None                |
| <b>Negative Unspiked Element</b>  |                           |                     |
| Absolute value ><br>LLOQ or ½ LOQ | Non-detect                | UJ                  |
| Absolute value ><br>LLOQ or ½ LOQ | Detect ≤ 5x<br> unspiked  | J-                  |
| Absolute value ><br>LLOQ or ½ LOQ | Detect > 5x<br> unspiked  | None                |

**Method 6010D SIC (LLOQ); DoD QSM ICS (1/2 LOQ)**

**5.7 (Optional) Internal Standards (IS)**

Internal standards (IS) performance criteria for Method 6010D is an option for difficult matrices and as an alternative to the Method of Standard Addition (MSA). The IS ensures that sensitivity and response are stable during every analytical run. Internal standards, if utilized, should be added to all calibration standards, blanks, instrument QC checks, samples, and batch QC.

The analysis of IS determines the existence and magnitude of instrument drift and physical interferences. For ICP-OES, Yttrium or Scandium is often used for this purpose. Other elements may need to be used as IS when samples contain significant native amounts of

the recommended IS (as indicated by high bias of IS recoveries). The IS element intensity is used to ratio the analyte intensity signals for both calibration and quantitation.

### *Evaluation of IS*

Internal Standard percent recoveries in the samples should be within 30-120% of the IS recoveries in the Initial Calibration Blank (ICB). Verify that internal standards were added to all analytical runs.

If the percent recovery is < 30% in samples, qualify detects as estimated **J+** and non-detects as estimated **UJ**.

Detects for analytes quantitated using an IS percent recovery > 120% should be qualified estimated **J-**. Non-detects should be qualified as estimated **UJ**.

If extremely low or high area counts are reported (<10% or > 150% of the area for associated standards), detects and non-detects should both be qualified **X**, exclusion of data recommended.

## **6.0 Stage 3 Validation**

**Note: Stage 3 validation includes all of Stage 1, Stage 2A and Stage 2B**

The following documents are used for a Stage 3 validation

- Raw Data (including any laboratory forms, instrument outputs, spreadsheets, or handwritten calculations necessary for recalculation and re-quantification)
- Standards Traceability forms and worksheets
- Optional Detection Limit studies (Instrument Detection Limits, Method Detection Limits)

Stage 3 validation includes the recalculation and re-quantification of selected samples, and method and instrument QC. The types of results that should be recalculated and re-quantified include target analytes, analytes with detects above the LOQ, and field QC samples (blanks and duplicates). For method QC results; spiked recoveries and method blanks should be considered. For instrument QC; calibrations (including regressions), calibration verifications, and internal standards should be recalculated and re-quantified. Some calculations may include the need to review standards preparation and serial dilutions.

### **6.1 Samples and Field QC Recalculations**

When choosing samples, field QC and analytes for recalculation and re-quantification, consideration should be given to the laboratory's batching scheme to ensure a representative subsample of recalculations is performed. Additionally, if priority contaminants or contaminants of concern are identified in the QAPP, those analytes should be selected for re-quantification and recalculation. Other circumstances that should be prioritized for recalculation and re-quantification are diluted samples, re-runs of samples due to QC failures, and field QC blank failures.

Recalculation and re-quantification should be performed on the designated percentage of the samples per SDG (or however defined in the QAPP, such as percentage of total project

samples) per analytical suite. As a minimum, it is recommended that 10% of the data should be recalculated and re-quantified unless specific instructions are given in the QAPP.

Sample recalculations should include the raw instrument result, re-quantified from the instrument response against the calibration function, and the final reported sample result, including any dilution, preparation factor, or percent moisture (if applicable). The equations in Appendix B can be used to calculate a sample result from the corresponding reported calibration or regression function, as appropriate.

Verify that one or more of the laboratory's reporting limits (such as limit of quantitation) are calculated correctly for the non-detects and reported accordingly. If a detection limit study was identified by the QAPP, recalculate one or more analyte detection limits.

Re-quantitate all detected target analytes in the 10% sample data chosen. For some samples, all results may be non-detects, therefore recalculation would not be necessary. Verify that sample-specific results have been adjusted correctly to reflect percent solids, original sample mass/volume, and any applicable dilutions.

Re-quantitate all detects found in the field QC blanks (such as field blanks or equipment blanks). Field QC sample duplicate recalculations should include re-quantification of the same detected analyte sample/duplicate pair and verification of the percent difference (%D), or relative percent difference (RPD), as reported.

When recalculations require rounding of data, that rounding should be completed only once at the end of all calculations to minimize rounding errors. Calculations should be rounded to the significant figures of the underlying criteria. For example, an LCS criteria of 90-107% would still be considered acceptable if the recalculation was 107.4%

#### *Evaluation of Sample and Field QC Recalculations*

If the laboratory's quantitation, or reporting limits (however defined) are calculated incorrectly, then continue to recalculate limits until it is determined that the problem is systemic (such as incorrect equations used) or isolated (such as a transcription or rounding errors).

For systemic (defined as widespread and major in nature) issues that cannot be corrected through a revised laboratory report, qualify all results as **X**, exclusion of data recommended.

For isolated cases, use professional judgment. It may be necessary to engage the point of contact as identified in the project QAPP to communicate with the laboratory, so they can provide revised (corrected) results. In all cases, if calculation errors affect project target analytes, the point of contact should be notified, and all affected results noted in the data validation report, including listing the calculation errors.

#### **6.2 Method QC Recalculations**

Re-quantification of batch QC sample results should use raw instrument response in tandem with the reported calibration slope; the preparation information; and percent moisture for solid samples to recreate the reported result.

### **6.2.1 Laboratory Control Sample/Laboratory Control Sample Duplicate (LCS/LCSD)**

To check that the spike percent recovery was calculated and reported correctly, using the equation in Appendix B, re-quantitate and then recalculate 10% of the target analytes as outlined in the UFP-QAPP Worksheet #12 or #15. Use a random 10% of the analytes in the LCS/LCSD if contaminants of concern have not been specifically identified. Recalculate RPDs (if applicable) from LCS/LCSD pairs that would result in the qualification of a sample.

### **6.2.2 Matrix Spike/Matrix Spike Duplicate (MS/MSD) or Laboratory Duplicate (LD)**

Re-quantitate 10% of the target analytes as listed in the UFP-QAPP Worksheet #12 or #15 for both the MS and the MSD (or LD). Use a random 10% of the analytes in the MS and MSD if contaminants of concern have not been identified. The RPDs of the recalculated MS/MSD pairs (or Sample Result/LD pairs) should be calculated from the MS/MSD concentrations, not from the recoveries.

### **6.2.3 Method Blank (MB)**

Method blank analytical results are assessed to determine the existence and magnitude of contamination problems associated with sample extraction (if applicable) and analysis. If problems with any method blank exist, all associated data should be carefully evaluated to determine whether there is any bias associated with the data, or if the problem is an isolated occurrence not affecting other data. Results may not be corrected by subtracting any blank values.

Re-quantitate one or more detects found in the method blank (if applicable) from the reported regression curve.

### **6.2.4 Dilution Test (DT) or Post Digestion Spike (PDS) and Serial Dilution (SD)**

To check that the spike percent recovery was calculated and reported correctly, re-quantitate and then recalculate 10% of the target analytes as outlined in the UFP-QAPP Worksheet #12 or #15 for at least one DT or PDS sample, if applicable. Use a random 10% of the analytes if contaminants of concern have not been specifically identified.

For any sample results (detects) that were in the linear range but required a serial dilution within the working calibration range to report the data, recalculate at least one target analyte from one or more samples that was diluted.

#### *Evaluation of all LCS, MS, MB, DT/PDS and SD Recalculations*

If transcription errors (or other minor issues such as rounding errors) are found in method QC results, use professional judgment to qualify the data. It may be necessary to engage the point of contact as identified in the UFP-QAPP to communicate with the laboratory, so they can provide revised (corrected) results. In all cases, if method QC calculation errors affect project target analytes, the point of contact should be notified, and all affected results noted in the data validation report, including listing the calculation errors.

For systemic (defined as widespread and major in nature) problems with LCS/LCSD calculations qualify all affected analytes in associated samples as **X**, exclusion of data recommended.

For systemic problems with method blanks, MS/MSD or LD, and DT/PDS calculations qualify all affected analyte detects in associated samples as estimated **J** and non-detects as estimated **UJ**.

### **6.3 Instrument QC Recalculations**

#### **6.3.1 Initial Calibration, Initial/Continuing Calibration Verification (ICV/CCV), and Low Level Calibration Check Verification (LLCCV)**

Initial calibration (ICAL) recalculations should use the raw instrument response for the target analytes and associated internal standards to recreate the calibration curve from the individual calibration standards. If multiple types of calibration curves are employed in an analytical suite, then one analyte per curve type should be recalculated.

Re-quantitate and recalculate the regression function (if used for multi-point calibration), slope, intercept, and (r) values reported for at least 10% of the target analytes per each internal standard (if used), preferably analytes of concern which were identified in the QAPP, per initial calibration curve type. Some OES instruments report  $r^2$  (coefficient of Determination) values instead of (r).

The laboratory may employ a linear or weighted linear least squares regression. The low standard should be recalculated using the calibration curve and evaluated. If the ICAL included refitting of the data back to the model (RSE), then recalculate 10% of the target analytes for the RSE in each ICAL.

Re-quantify and recalculate the ICV, CCV, and LLCCV result and %D for at least 10% of the target analytes for every ICV, CCV, and LLCCV bracketing reported results, proportionally selecting analytes based on calibration curve types used in each initial calibration.

#### **6.3.2 Initial Calibration Blank (ICB) and Continuing Calibration Blank (CCB)**

Verify each ICB and CCB bracketing reported results by comparing the summary form results to the raw data. Re-quantify one or more detects in the ICB and each CCB (if applicable) using the reported calibration curve.

#### **6.3.3 Interference Check Solutions (ICS) or Spectral Interference Checks (SIC)**

Verify the result and recalculate the percent recovery for at least 10% of the target analytes for every ICS or SIC bracketing reported results. Recalculate at least 10% of the reported concentrations of non-spiked metals in each ICS.

#### **6.3.4 (Optional) Internal Standards (IS)**

If Internal Standards were used, the analyte quantitation should be evaluated for all detects by evaluating the raw data. Analyte concentrations should be calculated based on the IS associated with that analyte. Quantitation should be based on the atomic emission line (wavelength) in the analytical method (or laboratory SOP listed in the QAPP) for both the IS and target analytes. The analyte quantitation should be based on the regression function from the appropriate ICAL.



Recalculate IS percent recoveries reported from the raw data for at least 10% of the samples per SDG), and verify internal standard results for samples that were qualified due to out-of-control internal standard results.

If errors are discovered, request revisions of the laboratory report per the QAPP point of contact.

#### *Evaluation of all ICAL, ICV/CCV/LLCCV, ICB/CCB, ICS/SIC, and IS Recalculations*

If the files provided do not match the quantitation report, the regression function reported is likely to be from another initial calibration and the laboratory report should be revised. The point of contact (UFP-QAPP Worksheet #6) should be reached to get a revised (corrected) report from the laboratory. For calculation errors for any type of regression equations that cannot be corrected in a revised report, qualify all the data as **X**, exclusion of data recommended.

The reprocessed low standard of a regression curve should be within 20% of the true value. If the recalculated concentration is not within 20% of the true value, qualify detects (at the LOQ and above) for the affected analytes as estimated **J** and non-detects as estimated **UJ** in the associated samples. If the recalculation shows gross error (> 30%), then non-detects should be qualified as **X**, exclusion of data recommended.

Qualify all associated data as **X** if the corresponding ICV/CCV %D has been calculated incorrectly by the laboratory and cannot be corrected in a revised laboratory report.

Qualify all associated analyte detects as estimated **J** and non-detects as estimated **UJ** if the corresponding LLCCV %D has been calculated incorrectly by the laboratory and cannot be corrected in a revised laboratory report or the corresponding true value cannot be determined.

Qualify all affected analyte detects in associated samples as estimated **J** and non-detects as estimated **UJ** if the corresponding ICB/CCB detects have been calculated incorrectly and cannot be corrected in a revised laboratory report.

Qualify all associated analyte detects as estimated **J** and non-detects as estimated **UJ** if the corresponding ICS/SIC interference checks have been calculated incorrectly and cannot be corrected in a revised laboratory report.

Qualify all data as **X** if the corresponding IS has been calculated incorrectly (or if the IS has been assigned to the wrong analyte) by the laboratory and cannot be corrected in a revised laboratory report.

In all cases where instrument QC are calculated incorrectly, the UFP-QAPP point of contact should be notified and noted in the validation report.

#### **6.4 Standards Traceability**

Evaluate the calibration standards used for the analytes of concern. From the Certificate of Analysis (however named), verify that the “true values” of each analyte of concern were correctly applied to create the calibration curve, and that all analytes of concern were in the calibration mix.

All initial instrument calibrations should be verified with a standard obtained from a second manufacturer prior to analyzing any samples. From the standard Certificate of Analysis, verify that a second source was used for the Initial Calibration Verification (ICV). The use of a standard from a second lot obtained from the same manufacturer (independently prepared from different source materials) is acceptable for use as a second source standard.

Check that the stock standards were diluted properly into working standards by recalculating the dilutions of one or more calibration standards. Recalculate one or more method QC sample dilutions (such as LCS or MS/MSD) from the stock to the working standard.

Note: It is not the role of the data validator to evaluate the Certificate of Analysis for compliance with the *ISO-17034 Standard*, but to verify that stock and working standards were correctly applied in the creation of calibration curves.

#### *Evaluation of Standards*

Professional judgment should be used when evaluating errors in standards preparation. The point of contact identified in the project QAPP (UFP-QAPP Worksheet #6) should be reached to get a revised (corrected) report from the laboratory. Issues {that does not affect the results of any target analytes} should be noted in the data validation report.

For systemic (widespread) issues that cannot be corrected by the laboratory, or issues that affect the results of target analytes, the data should be qualified as **X**, exclusion of data recommended.

For ICV standards that were not verified to be from a second source, qualify all affected data as **X**, exclusion of data recommended. No samples should have been run without a valid second source standard (per QSM requirements).

For expired standards, per QSM requirements, a laboratory cannot use a standard beyond its expiration date. All associated data should be qualified as **X** if expired standards were used. The expiration date of any working standard is based on the expiration date of the primary or stock standard.

### **6.5 Detection/Quantitation Limit Studies (Optional)**

In some cases, a project QAPP may specify the review and validation of a detection/quantitation limit study. This could include studies such as Method Detection Limits (MDLs), quarterly LOD verifications, or LOQ verifications. The project QAPP should specify the criteria for evaluating the study. As a minimum, at least 10% of the raw data in the study should be recalculated.

#### *Evaluation of Detection Limit Studies*

The criteria for evaluating a detection/quantitation limit study should be listed in the project QAPP. The following guidance should be enacted if the QAPP does not specify the evaluation criteria.

If transcription errors (or other minor issues such as rounding errors) are found in detection/quantitation limit studies, use professional judgment to qualify the data. It may be necessary to engage the point of contact as identified in the project QAPP to communicate with the laboratory, so they can provide revised (corrected) results. In all cases, if calculation errors affect project detection or quantitation limits, the point of contact should be notified, and all affected results noted in the data validation report, including listing the calculation errors.

When calculation errors are uncovered that cannot be corrected by the laboratory and that affect detection/quantitation results, consideration should be given to qualify the study as **X**, exclusion of data recommended.

## 7.0 Stage 4 Validation

**Note: Stage 4 validation includes all of Stage 1, Stage 2A, Stage 2B and Stage 3**

Raw Data (including any instrument outputs, raw interference/background corrections data files, Spectra)

Stage 4 is a qualitative/quantitative review of detected and non-detected analytes from the instrument outputs (raw data files). Quantitation reports, instrument background corrections, and interference corrections data files are required to perform the review of the instrument outputs.

The application of qualitative criteria for metals analysis requires professional judgment. It is up to the reviewer's discretion to obtain additional information from the laboratory through the QAPP point of contact, if necessary.

## 7.1 Raw Sample Results

For ICP-OES, confirm that reported results for 10% of the QC standards (including instrument and method QC) and 10% of the positive field sample result concentrations (detects at or above the LOQ) are the average of at least three readings (of a single injection). Verify that RSDs of these reported averages are < 5%. If the standards display high RSDs, this indicates a serious instrument problem that could impact field sample results. Qualify associated positive field sample results as estimated **J** and non-detects as **X**, exclusion of data recommended.

If the raw data result from a detect in an isolated field sample displays a high RSD the data validator should qualify all the associated detects as estimated **J**.

If, in the professional judgment of the validator, there are instances of unwarranted manipulation of data then those cases should be reported to the project team as soon as practical (UFP-QAPP Worksheet #6).

The following are some (non-inclusive) instances of data manipulation that should be reported to the project team:

- Manipulation of hold time data
- Analyzing samples known to be lacking acid preservation
- Excessive use of instrument rinse blanks to 'pass' QC criteria

- More than three readings of a single injection (for select samples and not the entire batch)
- Running multiple CCVs and using the last one that passes criteria
- Running ICVs known to be from the same source as the calibration standards

## 7.2 Spectral Interferences

Interelement effects must be evaluated for each individual instrument. Intensities will vary not only with optical resolution, but also with operating conditions (such as power and argon flow rates). The laboratory is required to document for each wavelength (analyte) the suspected interferences.

Depending on the type of instrument used, review the raw spectra files to qualitatively verify the absence of spectral interference. Some sequential instruments verify the absence of spectral interference by scanning over a range (0.5 nm), centered on the wavelength of interest. Samples that show an elevated background emission across the range may be background-corrected by applying a correction factor, or an alternate wavelength that does not exhibit a background shift or spectral overlap may also be used.

For multipoint calibration methods that employ whole spectral regions, spectral interferences are handled by including spectra of the interfering elements in the calibration algorithm. The interferences listed in Method 6010D (Table I) may be employed by the data validator to review the types of inter-element interferences encountered for each wavelength (analyte).

The interelement correction equations themselves are complex and instrument specific, but the accuracy of any interelement corrections are verified through the analysis of the SIC for Method 6010D. The mixed element SIC is used daily and is reviewed in Stage 2B. However, if spectral interference is observed the data validator should review the individual (single) element SIC (analyzed by the laboratory every 6 months and used to set the interelement corrections). The absolute value of the concentration observed for any unspiked analyte in the single element SIC check must be  $< 2 \times \text{LLOQ}$  for Method 6010D ( $< 1/2 \text{ LOQ}$  for QSM ICS-A).

When uncorrected, interelement interferences may produce false positive results. However, overcorrection can cause a negative bias. If there is any indication of spectral interference from the single element SIC raw data, the validator should note this in the data validation report and qualify all associated target analytes as estimated **J** for detects and estimated **UJ** for non-detects. If, in the professional judgment of the validator, the spectral interference calls into question the validity of the data, the QAPP point of contact should be notified via UFP-QAPP Worksheet #6.

For methods other than 6010D, the ICS-A specifies an absolute value of the concentration for all non-spiked analytes as  $< 1/2 \text{ LOQ}$ .

## 7.3 Instrument Detection Limits (IDLs) and Linear Ranges (LR)

ICP-OES instruments should have a Signal to Noise (S/N) ratio of at least 3:1. Qualitatively review instrument outputs to verify that S/N is appropriate to produce quantitative data. Instrument detection limits (IDLs) are useful means to evaluate excess instrument noise level.

Use professional judgment to review and qualify sample data that is reported with a low S/N ratio. Low ratios give evidence that background corrections were not applied or applied incorrectly. If, in the professional judgment of the validator, background interference calls into question the validity of the data, the QAPP point of contact should be notified via UFP-QAPP Worksheet #6.

The Linear Range (LR) establishes the highest concentration that may be reported without diluting the sample. Following calibration, the laboratory may choose to analyze a standard at a higher concentration than the high standard in the calibration. The standard must recover within 10% of the true value, and if successful, establishes the LR. The linear range standards must be analyzed in the same instrument run as the calibration they are associated with. If the project team is using Method 6010D requirements (as specified in the project QAPP), then qualify all detects above the working calibration range as estimated **J** as long as the LR is not exceeded. All detects above the working calibration range should be noted in the validation report.

The LR is also called the Linear Dynamic Range (LDR) in the QSM. If the QAPP uses DoD QSM requirements, then all detects above the working calibration range (but within the LDR) should be diluted to within the working calibration range. If the samples were not diluted, then all detects above the working calibration range are qualified as estimated **J**. If a high level check standard was run above the working calibration range and above the highest sample result (and agrees within  $\pm 10\%$ ), then qualification is not necessary. All detects above the working calibration range should be noted in the validation report.

## Appendix A: Method QC Tables

Note: The following Table is based on the QSM 5.3 Standard, with Method 6010D for comparison. The Table does not include all the QC elements from the method or as listed in this guidance document.

| QC Check  | 6010D Frequency and Acceptance Criteria  | QSM Version 5.3 Frequency and Acceptance Criteria   |
|---|--|---|
| <p><b>Linear Dynamic Range (LDR) or high-level check standard</b></p> | <p><b>Linear Range:</b> At initial set up and Daily with a high standard, if samples are reported above the Calibration Range</p> <p>The high check standard establishes the Linear Range.</p> <p>Within <math>\pm 10\%</math> of true value.</p>  | <p><b>Linear Dynamic Range:</b> At initial set up and checked every 6 months with a high standard at the upper limit of the range.</p> <p>Within <math>\pm 10\%</math> of true value.</p> |
| <p><b>Initial calibration (ICAL) for all analytes</b></p>             | <p>At instrument set-up and Daily, prior to sample analysis.</p> <p>Minimum one high standard and a Calibration Blank.</p> <p>If a multi-point calibration is used:</p> <p>Minimum of 3 standards, the low standard must be <math>\leq</math> LLOQ</p> <p>Each analyte should meet one of the linear regression options below:</p> <p><b>Coefficient of Determination (COD) <math>r^2 \geq 0.990</math></b></p> <p><b>Correlation Coefficient (r) <math>\geq 0.995</math></b></p> <p>or</p> <p><b>RSE for each analyte <math>\leq 20\%</math>.</b></p> | <p>Daily ICAL prior to sample analysis.</p> <p>Minimum one high standard and a Calibration Blank.</p> <p>If more than one calibration standard is used, <math>r^2 \geq 0.99</math>.</p>   |

| QC Check  | 6010D Frequency and Acceptance Criteria   | QSM Version 5.3 Frequency and Acceptance Criteria  |
|---|---|--|
| <b>Initial Calibration Verification (ICV)</b>       | <p>Once after each ICAL, analysis of a second source standard prior to sample analysis.</p> <p>All reported analytes within <math>\pm 10\%</math> of true value.</p>  | <p>Once after each ICAL, analysis of a second source standard prior to sample analysis.</p> <p>All reported analytes within <math>\pm 10\%</math> of true value.</p> |
| <b>Continuing Calibration Verification (CCV)</b>    | <p>After every 10 field samples and at the end of the analysis sequence.</p> <p>All reported analytes within <math>\pm 10\%</math> of the true value.</p>   | <p>After every 10 field samples and at the end of the analysis sequence.</p> <p>All reported analytes within <math>\pm 10\%</math> of the true value.</p>            |
| <b>Low-Level Calibration Check Standard (LLCCV)</b> | <p>Daily; A low level check standard or readback verification at the LLOQ for single point calibrations; or</p> <p>at the lowest calibration standard for a multi-point calibration.</p> <p>All reported analytes within <math>\pm 20\%</math> of true value.</p> <p>Daily; A mid-level check standard (readback verification) at the mid-point of the Linear Range for single point calibrations; or</p> <p>at the middle calibration point for multi-point calibrations.</p> <p>All reported analytes within <math>\pm 10\%</math> of true value.</p> | <p>Daily; LLCCV should be <math>\leq</math> LOQ.</p> <p>All reported analytes within <math>\pm 20\%</math> of true value.</p>  |

| QC Check  | 6010D Frequency and Acceptance Criteria  | QSM Version 5.3 Frequency and Acceptance Criteria  |
|---|--|--|
| <b>Initial and Continuing Calibration Blank (ICB/CCB)</b>                             | <p>Immediately after the ICV and immediately after every CCV.</p> <p>Target analytes must be &lt; ½ LLOQ for the ICB and &lt; LLOQ for the CCBs.</p>   | <p>Immediately after the ICV and immediately after every CCV.</p> <p>The absolute values of all analytes must be &lt; ½ LOQ or &lt; 1/10<sup>th</sup> the amount measured in any sample.</p>                                 |
| <b>Internal standards (IS)</b>  | <p>Optional, if spectral interferences are noted.</p> <p>Added to every calibration standard, field sample, blank, instrument and method QC sample.</p>  | Not specified in QSM Version 5.3   |
| <b>Method Blank (MB)</b>  | <p>One per preparatory batch.</p> <p>No target analytes in the method blank should be ≥ ½ LLOQ unless the sample concentration is &gt; 10x the blank contamination.</p>  | <p>One per preparatory batch.</p> <p>The absolute values of all analytes must be &lt; ½ LOQ or &lt; 1/10<sup>th</sup> the amount measured in any sample or 1/10<sup>th</sup> the regulatory limit, whichever is greater.</p> |
| <p><b>Interference Check Solutions (ICS) or Spectral Interference Check (SIC)</b></p> | <p><b>Single Element SIC</b></p> <p>At setup and every 6 months.</p> <p>Absolute value of the concentration observed for any unspiked analyte in the SIC &lt; 2x LLOQ.</p> <p><b>Mixed Element SIC</b></p> <p>Daily, After ICAL and prior to sample analysis.</p> <p>A mixed element check SIC that all target analytes must be &lt; ± the LLOQ.</p> | <p>After ICAL and prior to sample analysis.</p> <p><b>ICS-A:</b> Absolute value of concentration for all non-spiked project analytes &lt; 1/2 LOQ;</p> <p><b>ICS-AB:</b> Within ± 20% of true value.</p>                     |



| QC Check   | 6010D Frequency and Acceptance Criteria   | QSM Version 5.3 Frequency and Acceptance Criteria  |
|--|---|--|
| <p><b>Laboratory Control Sample (LCS)/Laboratory Control Sample Duplicate (LCSD);</b></p> <p><b>Matrix Spike (MS); Matrix Spike Duplicate (MSD)</b></p> <p><b>Matrix Duplicate (MD), also called Laboratory Duplicate (LD)</b></p> | <p>One each <b>LCS</b> (or <b>LCS/LCSD</b> pair) and one <b>MS/MSD</b> pair (or <b>Sample result/LD</b> pair) per preparatory batch.</p> <p>For <b>LCS</b>, use <math>\pm 20\%</math> for recovery until historical limits can be generated.</p> <p>For <b>MS</b>, use <math>\pm 25\%</math> for recovery until historical limits can be generated.</p> <p><b>MSD (LD) or LCSD:</b> RPD of all analytes <math>\leq 20\%</math> (between <b>MS/MSD</b> or <b>Sample result/LD, LCS/LCSD</b> pair).</p> | <p>One each <b>LCS</b> and <b>MS/MSD</b> pair per preparatory batch or use <b>Sample Result/MD</b> pair.</p> <p><b>LCS:</b> A laboratory must use the QSM Appendix C LCS Limits for batch control if project limits are not specified.</p> <p>If the analyte(s) are not listed, use in-house LCS limits if project limits are not specified.</p> <p><b>MS:</b> A laboratory should use the QSM Appendix C LCS limits as a basis of comparison for the <b>MS/MSD</b>.</p> <p><b>MSD or MD:</b> RPD of all analytes <math>\leq 20\%</math> (between <b>MS/MSD</b> or <b>Sample result/MD</b>).</p> |
| <p><b>Dilution Test (DT)</b></p>   | <p>Once per batch if a target analyte(s) concentration is within the Linear Range of the instrument and sufficiently high (minimally, 25x greater than the LLOQ). This sample can be the MS.</p> <p>1:5 dilution should agree to within <math>\pm 20\%</math> of the original determination.</p>  | <p>One per preparatory batch if MS or MSD fails. Only applicable for samples with concentrations <math>&gt; 50x</math> LOQ (prior to dilution).</p> <p>Five-fold dilution must agree within <math>\pm 10\%</math> of the original measurement.</p>   |

| QC Check                                 | 6010D Frequency and Acceptance Criteria  | QSM Version 5.3 Frequency and Acceptance Criteria   |
|--|--|---|
| <p><b>Post Digestion Spike (PDS)</b></p> | <p>Once per batch. If a high concentration sample is not available for performing the DT, then</p> <p>a PDS on the MS should be performed. The test only needs to be performed for the specific elements that failed original MS limits, and only if the spike concentration added was greater than the concentration determined in the unspiked sample.</p> <p>The recovery of the post-digestion MS should fall within a <math>\pm 25\%</math> acceptance range, relative to the known true value, or otherwise within the laboratory derived acceptance limits.</p> | <p>Perform if MS/MSD fails.</p> <p>One per preparatory batch (using the same sample as used for the MS/MSD if possible).</p> <p>Applies for samples with concentrations &lt; 50x LOQ (prior to dilution).</p> <p>Recovery within 80-120%.</p> |

## Appendix B: Formulas used in Stages 3 and 4 Data Validation

### Multi-point Calibration

Linear Regression:  $y = mx + b$

(Only if internal Standard is used):

$$C_s = \frac{\left(\frac{A_s}{A_{IS}} - b\right) * C_{IS}}{m}$$

Where:

$C_s$  = Concentration, Sample

$A_s$  = Area (element wavelength intensity), Sample

$A_{IS}$  = Area(element wavelength intensity), Internal standard

$C_{IS}$  = Concentration, Internal Standard

$b$  = Intercept

$m$  = Slope

**LCS Percent Recovery:**

$$\text{Percent Recovery} = \frac{C_s}{C_K} \times 100$$

Where:

$C_s$  = Concentration, Reported

$C_K$  = Concentration, Known

**MS or MSD Percent Recovery:**

$$\text{Percent Recovery} = \frac{C_M - C_S}{C_K} \times 100$$

Where:

$C_M$  = Concentration, MS or MSD

$C_S$  = Concentration, Sample

$C_K$  = Concentration, Known

**Field Duplicate, MS/MSD or LCS/LCSD Duplicate Relative Percent Difference (RPD):**

$$RPD = \frac{|C_s - C_d|}{(C_s + C_d)/2} \times 100$$

Where:

$C_s$  = Concentration, Sample

$C_d$  = Concentration, Duplicate

**Calculation of sample amounts:**

$$\% \text{ Dry Weight} = \frac{\text{g of dry sample}}{\text{g of sample}} \times 100$$

$$\text{Concentration}_{DW} = \frac{C \times V}{W \times S}$$

where:

Concentration on a dry weight basis:

C = Digest concentration (mg/kg)

V = Final volume after sample preparation (L)

W = Wet sample mass (kg)

S = % Solids/100 = % dry weight/100