

Method Detection Limit Verification

1.0 BACKGROUND

The EPA MDL procedure, CFR Chapter 40, Part 136 Appendix B, Revision 2, requires that all laboratories perform a minimum of two low-level verification spikes and two blanks every quarter in which samples are being analyzed to evaluate their Method Detection Limit (MDL) and Limit of Quantitation (LOQ) for each quality system matrix, technology, and analyte (where appropriate). The spike levels should be at or below the LOQ of the analyte for a given method and matrix.

In the 2016 TNI Standard, Volume 1, Module 2, Section 3.1 the LOD is defined as “The minimum result, which can be reliably discriminated from a blank with a predetermined confidence level”. Chapter 40, Part 136 Appendix B, Revision 2 (2017) of the Code of Federal Regulations, which has been adopted for use in Florida, sets ‘method detection limits’ as “... the minimum measured concentration of a substance that can be reported with 99% confidence that the measured concentration is distinguishable from method blank results”.

2.0 IMPLEMENTATION

The following interpretation of the requirements in the 2016 TNI Standard, Volume 1, Module 4, section 1.5.2 and additional guidance shall be implemented when verifying the MDL:

- 2.1 The MDL and PQL shall be verified every quarter (4 times a year) for each analyte/matrix/test combination and on each analytical instrument used to report results. A minimum of two blanks and two low-level spikes should be prepared in separate batches and analyzed in separate batches every quarter. This includes identical analytical instruments that have the same detection limits and are used to report the same suite of analytes such as multiple GC-MS instruments. All routine sample processing steps shall be performed when verifying the MDL.
- 2.2 Since the laboratory defines the MDL as the critical value where the false negative rate is 50%, it is not recommended to fortify the sample used for this verification at a level equal to the MDL (where 50% of the measurements will yield a non-detect). A spike that is at least twice the value of the MDL is recommended.
- 2.3 For MDL verifications carried out for tests on non-potable water, the low-level spike shall consist of fortified laboratory grade water. For tests on solids or biological materials, where no matrix sample is readily available, the low-level spike shall consist of fortified silica sand or Teflon or glass beads with sufficient solvent or laboratory grade water to support the fortification. For cases where a surrogate matrix which does not contribute analyte or contaminate the fortified

sample with the analyte of interest (at levels greater than the detection limit) cannot be found, the MDL verification shall be performed in the absence of a matrix (e.g., using laboratory grade water instead of another matrix or surrogate matrix).

- 2.4 MDL verifications shall consist of positive detection equal to or greater than the previously established method detection limits (i.e., a signal distinctly greater than the method blank) or qualitative identification of the analyte signal based on mass identification, retention time, or another unique characteristic deemed appropriate by the unit supervisor. In addition, where required by methods employing mass spectrometric detection (such as EPA 8260, 8270, etc.), the secondary identification ions must be present in the MDL verification at acceptable ratios defined by method or SOP relative to primary quantification ions.
- 2.5 All routine and quarterly MDL verification data must be uploaded to the LIMS QC Statistics Tracking module.
- 2.6 The spiking level should be re-evaluated at least once a year. Recalculate the MDL at least once every thirteen months using the procedure described in SOP LB-007. Include data generated within the last twenty-four months, but only data with the same spiking level and acceptable calibrations and batch QC.
- 2.7 If the verified (recalculated) MDL is within 0.5 to 2.0 times the existing MDL, and fewer than 3% of method blank results have values above the existing MDL, then the existing MDL may be left unchanged. Otherwise, adjust the MDL to the new verification MDL.
- 2.8 An initial MDL shall be determined for each analyte/matrix/test combination whenever a new instrument is set-up, any time an existing instrument undergoes major repairs or modifications and whenever major method modifications are implemented. The procedure for determining initial MDLs is documented in SOP LB-007.

3.0 REFERENCES

- 3.1 DEP SOP LB-007, *Procedure and Policy for Demonstration of Capability for Methods, Instruments and Laboratory Staff*.
- 3.2 2016 TNI Standard, Volume 1, Module 4, section 1.5.2.
- 3.3 EPA 821-R-16-006, *Definition and Procedure for the Determination of the Method Detection Limit, Revision 2, December 2016*.