Total (Extracellular and Intracellular) Microcystins – ADDA by ELISA Analytical Methodology

Quick Reference	Standard/Reagent	Requirements
	Liquid Disinfectant	Manufacturer's Recommendations
Standard/Reagent Storage	Analysis Kit	Manufacturer's Recommendations
	Sodium Thiosulfate	Manufacturer's Recommendations
	0.10 N Sodium Hydroxide (NaOH) Solution	Manufacturer's Recommendations
	0.10 N Hydrochloric Acid (HCl) Solution	Manufacturer's Recommendations
Standard/Reagent	Standard/Reagent	Expiration
	Liquid Disinfectant	Manufacturer's Expiration Date
	Analysis Kit	1 Year After Opening/ Manufacturer's Expiration Date
Expiration	Sodium Thiosulfate	1 Year After Opening/ Manufacturer's Expiration Date
	0.10 N Sodium Hydroxide (NaOH) Solution	1 Year After Opening/ Manufacturer's Expiration Date
	0.10 N Hydrochloric Acid (HCI) Solution	1 Year After Opening/ Manufacturer's Expiration Date
Required Quality Control	QC Procedure	Frequency
Required Quality Control	See Chapter 7 of this method	See Chapter 7 of this method
Sample Collection	Preservation	Maximum Hold Time
Jumpio Joneonon	0-10°C	5 days

Method Reference

Ohio EPA Method 701.0 and 701.0-A, Version 2.4, May 2025

Survey Requirements

- For Method 701.0, each analyst must perform an acceptable annual MDL and submit the assay calibration report, curve and test report to dwlabcert@epa.ohio.gov.
- For Method 701.0-A, the laboratory must perform an acceptable annual MDL and submit the assay calibration report, curve and test report to dwlabcert@epa.ohio.gov.
- All reagents, standards and solutions used for this method will be audited for correct labeling and dating.
- All records will be audited.

1.0 General Method Summary

This method is used for the determination of total (extracellular and intracellular) Microcystins – ADDA in surface water, ground water and finished drinking water using enzyme-linked immunosorbent assay (ELISA).

The Ohio EPA Total (Extracellular and Intracellular) Microcystins – ADDA by ELISA Analytical Methodology is an immunoassay for the detection of microcystins in water samples. This test is an indirect competitive ELISA allowing the congener-independent detection of microcystins and nodularins. It is based on the recognition of microcystins, nodularins and their congeners by specific antibodies. Microcystins, nodularins and their congeners when present in a sample and a microcystin-protein analogue immobilized on the plate compete for the binding sites of antibodies in solution. After a washing step and addition of the substrate solution, a color signal is generated. The intensity of the blue color is inversely proportional to the concentration of the microcystins present in the sample. The color reaction is stopped after a specified time and the color is evaluated using a microplate reader at 450 nm. The concentrations of the samples are determined by interpolation using the standard curve constructed with each run.

Interferences

Due to the high variability of compounds found in water samples, test interferences caused by matrix effects cannot be completely ruled out. Ohio EPA continues to work with U.S. EPA and other experts to identify and provide more guidance on potential interferences.

2.0 Equipment and Supplies

- a. Glass or polyethylene terephthalate glycol (PETG) sampling container: 100mL
 - Cleaning of approved sample collection containers is acceptable if the laboratory can
 demonstrate effectiveness of the cleaning procedure by collecting and analyzing
 reagent water in 5% per batch of the cleaned containers. The reagent water results
 must be less than the reporting limit. The laboratory must document this procedure and
 must maintain these records. The Sample Bottle Cleaning Record on page 10 may
 be used to document the required information.
- b. Class 'A' Volumetric Flask: 500 mL
- Bench sheets or logbooks: See pages 10 16.
- d. Micropipette: Capable of 10 100 µL
- e. Multi-Channel Pipette: 50 300 µL
- f. Stepping Pipette: 100 500 µL (optional)
- g. Pipette Tips: Appropriate size for the pipette
- h. Multi-Channel Pipette Reagent Reservoir: Minimum 50 mL capacity
- i. Microplate Reader: Capable of analyzing at 450 nm
- j. Glass Vials: 4.0 mL and 40.0 mL
- k. Syringe Filters: 25 mm glass fiber, 0.45 μm or 1.2 μm pore size
- I. Gastight Luer-lock Syringes: 5.0 mL
- m. ELISA Sealing Teflon Tape or equivalent

- n. Freezer: Must maintain a temperature of ≤ 0.0 °C
- o. Refrigerator: Must maintain a temperature of 4.0 ± 2.0 °C
- p. Chlorine DPD Meter or Low-Range Chlorine Test Strips
- q. pH Meter or pH Test Strips 0-14 pH units
- r. Water Bath: Must maintain a temperature of 35.0 ± 0.5 °C
- s. Cyanotoxin Automated Assay System (CAAS) (optional)

3.0 Reagents

- a. Liquid Disinfectant: Commercially available, Roccal® or equivalent disinfectant.
- b. Analysis Kit (capable of analyzing all microcystin congeners with the ADDA structure). Store kit according to manufacturer's instructions.
- c. DPD-free Chlorine Reagent
- d. Sodium thiosulfate: De-chlorination agent (used for potable water). Commercially available.
- e. Reagent Water: Laboratory-available deionized water. Quality must meet minimum resistivity of $10.0\ M\Omega$.
- f. Ethanol/Dry Ice Mixture (optional)
- g. Sodium Hydroxide Solution (0.1N): In a 1000 mL volumetric flask, dissolve 4.0g sodium hydroxide (NaOH) in 800mL reagent water. Stir to dissolve. Bring to volume with reagent water.
- h. Hydrochloric Acid Solution (0.1N): In a 1000 mL volumetric flask, slowly and carefully add 8.3 mL concentrated hydrochloric acid (HCl) to 800 mL reagent water. Bring to volume with reagent water.

NOTE:

- Reagents, standards and kits must be labeled with the received, opened and expiration dates.
- Prepared reagents must be labeled with content, date made, expiration date, and analyst initials. Prepared reagents must be discarded one year after preparation or the manufacturer's expiration date for items used, whichever comes first.

4.0 Sample Collection/Preservation/Hold Time

a. **Sample Collection:** A minimum of 100 mL should be collected in a glass or polyethylene terephthalate glycol (PETG) container.

NOTE: Samples treated with chlorine or any other oxidizer (e.g., KMnO₄) must be quenched immediately after collection. 10.0 mg sodium thiosulfate added per 100 mL of sample is typically sufficient.

- b. **Preservation:** All samples must be protected from sunlight, cooled to $0.0 10.0^{\circ}$ C immediately after collection and maintained at $0.0 10.0^{\circ}$ C until analysis.
- c. **Holding time:** Drinking Water samples must be analyzed no later than five days from the time of collection.

NOTE: If the sample is <u>not to be used for drinking water compliance</u>, the hold time can be increased by initiating the first freeze cycle of the sample within five days of collection. The sample may be kept in this first freeze cycle indefinitely to be analyzed later.

d. When freezing, allow adequate volume for expansion. Glass sample containers must be placed on their side to prevent breakage.

5.0 Procedure

5.1 Sample Preparation

NOTE: Sample pH and chlorine levels must be checked upon receipt. An additional sample should be taken, or a portion of the sample should be poured off for these analyses.

- 1. Disinfect the work area.
- 2. Sample pH must be within the range of 5 11 pH units. Samples with pH levels outside of this range may produce inaccurate (falsely low) results and must be adjusted as necessary using 0.1N hydrochloric acid (HCl) or 0.1N sodium hydroxide (NaOH) solutions, prior to analysis.
- 3. Samples treated with chlorine: Check samples for residual chlorine. Any water samples not sufficiently quenched (< 0.1 mg/L) must not be analyzed. Unquenched water samples must be recollected and appropriately quenched immediately after collection.

5.2 Sample Lysing Procedure by Freeze/Thaw

- 1. Shake the sample and pour approximately 20.0 mL of the sample into two properly labeled, 40.0 mL vials to begin the three freeze/thaw lysing cycles. Keep the second vial in the third freeze cycle in case original sample vial cracks.
- 2. Place vials in the freezer until completely frozen (To speed up the process, vial(s) may be immersed in a saturated sodium chloride solution or dry ice/ethanol solution).

NOTE: Do not fully submerge the vial(s) if using either solution.

NOTE: Place glass sample vial(s) on its side in freezer to prevent vial(s) from cracking as the water freezes and expands.

3. Once sample is completely frozen, remove from freezer (or sodium chloride solution) and thaw. To speed up the thawing process, vial(s) may be left at room temperature, placed in a container of lukewarm water or placed in a 35.0°C water bath until completely thawed.

NOTE: Do not fully submerge the vial(s).

- 4. Repeat steps 1 and 2, two more times.
- 5. Once sample is completely thawed for the 3rd time, draw 5.0 mL of sample into the syringe and attach a filter.
- 6. Rinse the filter by passing a minimum of 5.0 mL sample through the filter and discard the filtrate.
- 7. Again, draw 5.0 mL of the sample into the syringe, re-attach the rinsed filter, and filter approximately 2.0 mL of sample into two, properly labeled, 4.0 mL vials. Samples are ready for immediate analysis.

6.0 Analysis

The accuracy of ELISA analysis is highly dependent upon analyst technique, adequate storage conditions of the test kit, pipetting sequence, accuracy of reagent volumes and maintenance of constant/optimum laboratory temperature during the analysis. The ELISA analysis is a time sensitive procedure. Care must be taken to ensure the reagent addition steps are completed in an efficient manner and incubation times are followed according to manufacturer's instructions.

- 1. Verify kit standards and reagents are used prior to the expiration date.
- 2. The assay procedure must be performed away from direct sunlight.
- 3. Bring samples and standards to room temperature prior to analysis.
- 4. Follow manufacturer's instructions provided with the individual Microcystins ADDA kit for calibration, quality control (QC) and sample analysis procedures.

NOTE: If sample analysis results in a higher concentration than the highest standard in the calibration curve, the sample must be diluted and reanalyzed. If diluted, the sample must be diluted using the LRB to match the matrix and can be prepared in a 4 mL vial using the ratios in the table below.

Samples may not be diluted in the well plate.

Dilution	Sample μL	Laboratory	Adjusted	Maximum
		Reagent Blank	Reporting Limit	Microcystin
		μL	μg/L	μg/L
2	500	500	0.6	10
5	200	800	1.5	25
10	100	900	3.0	50
20	50	950	6.0	100

If a sample is diluted, the final values must be calculated by multiplying the result by the proper dilution factor. Report calculated values.

5. Save and print a copy of the calibration curve and sample results as part of the laboratory's record maintenance protocol. Record the analyst initials, date of analysis, and the kit lot number/expiration date on the results page.

7.0 Quality Control Requirements

7.1 Initial Demonstration of Capability

For both manual and automated analyses, to maintain the reporting limit set in the method, demonstration of the capability to achieve a Method Detection Limit (MDL) less than the reporting limit must be performed by each new analyst seeking certification or whenever there is a change in analytical performance (i.e., a change in instrument hardware or operating conditions).

MDLs must be established for microcystins using a standard with a concentration between one and ten times the reporting limit. To calculate the MDL value, <u>during the same run</u>, take seven replicate aliquots of the standard and process them through the entire analytical method. **The 7 MDL aliquots must be treated as 7 samples; QC aliquots cannot be included in the MDL calculation.** Once the results for the seven replicates have been obtained, calculate the MDL as follows:

 $MDL = (t)*(SD_R)$

Where: t = Student's t value for a 99% confidence interval and a standard

deviation estimate with n-1 degrees of freedom

(t = 3.143 for the seven replicates)

SD_R = Standard deviation of the replicate aliquot analyses

The study will be valid if the resulting value of the MDL is no more than ten times lower than the replicate standard concentration level and does not exceed the replicate standard concentration level, and all QC requirements are met (see section 7.3). The **Total Microcystins-ADDA Method Detection Level (MDL) Report** located on the Laboratory Certification webpage (https://epa.ohio.gov/ddagw/labcert) may be used. Save and print a copy of each MDL study (including associated test data and calibration curves), adding the test kit lot number/expiration date and the analyst's initials as part of the laboratory's record maintenance protocol. This data must also be sent to the Laboratory Certification Section at dwlabcert@epa.ohio.gov for review.

7.2 Annual Demonstration of Capability

MDL studies must pass all QC. Qualifiers are not permitted.

- Manual Analysis MDL Requirements: Annual MDL studies, including associated test data, calibration curves, test kit lot number/expiration date and the analyst's initials, for each certified analyst must be sent to the Laboratory Certification Section at dwlabcert@epa.ohio.gov for review. Save and print a copy of each MDL study as part of the laboratory's record maintenance protocol.
- 2. Automated Analysis MDL Requirements An annual MDL study must be completed and documented for each instrument used for certified drinking water analysis. These annual MDL studies, including associated test data, calibration curves, test kit lot number/expiration date and the analyst's initials, must be sent to the Laboratory Certification Section at dwlabcert@epa.ohio.gov for review. Save and print a copy of each MDL study as part of the laboratory's record maintenance protocol.

7.3 Analyst QC Requirements

Certified Analyst Requirements

All analysts certified for manual analysis are required to generate an MDL study for initial certification and then at least annually.

All analysts certified for automated analysis are required to generate and submit an MDL study for initial certification.

Operationally Certified Analyst Requirements

Operational certification is not available for this method.

7.4 QC Requirements with Each Analysis

- 1. With each sample batch analysis, the following QC samples must be analyzed:
 - a. Laboratory Reagent Blank (LRB): An aliquot of reagent water that is lysed and filtered to match the sample processing procedure. An LRB must be analyzed with each batch

of samples to verify laboratory, reagents and supplies are free of contaminants. The LRB must contain sodium thiosulfate if drinking water samples are included in the analysis batch. Values exceeding the reporting limit require corrective action and reanalysis of the sample batch.

NOTE: Do not use the LRB provided in the kit.

- b. Low Calibration Range Check (LCRC): An LCRC must be analyzed with each batch of samples to verify accuracy of the calibration curve near the reporting limit. The LCRC may be one of the curve's calibration points and the concentration must be ≥ 0.24 μg/L and ≤ 0.50 μg/L. Acceptance limits must be within ± 40% of the true value. LCRC values exceeding the acceptance limits require corrective action and reanalysis of sample(s) with results below the concentration of an acceptable QCS in the same analytical batch. If reanalysis is not possible, all sample concentration results less than an acceptable QCS analyzed in the same batch must be appropriately qualified with a J or UJ, based on the result and noted in the final report.
- c. Quality Control Standard (QCS): A secondary source QCS must be analyzed with each batch of samples to verify the concentration of the calibration curve. If a QCS is already included in the kit, it may be used if it has a different lot number than the calibration standards and was prepared from a separate primary stock. Acceptance limits must be within ± 25% of true value. QCS values exceeding the acceptance limits require corrective action and reanalysis of sample(s) with results greater than the concentration of an acceptable LCRC in the same analytical batch. If reanalysis is not possible, all sample concentration results greater than an acceptable LCRC analyzed in the same batch must be appropriately qualified (J/UJ) and noted in the final report.

NOTE: If both LCRC and QCS exceed acceptance limits and reanalysis is not possible all results must be appropriately qualified with a J or UJ, based on the result.

- Analyze all calibration standards, QC standards and samples in at least two well replicates.
 The mean of the well replicates must be used in all analytical calculations and reporting of sample results.
- 3. The curve generation must include a calibration concentration point less than or equal to the reporting limit.
- 4. Calibration curve Correlation Coefficient (R) must be ≥ 0.990 or calibration curve Coefficient of Determination (R²) must be ≥0.980 to be acceptable.
- 5. Coefficient of Variation (%CV) for well replicate absorbance values for calibration *standards* and QC *standards* must all be ≤10.0%. It is acceptable to have <u>one</u> calibration or QC standard >10.0% as long as it is ≤15.0%. The zero standard is excluded from this requirement.
- 6. If %CV for more than one calibration or QC standard is >10.0%, or if one is >15.0%, the analytical run is not acceptable. Corrective action and reanalysis of the sample batch is required.

Calculate %CV as follows:

 $%CV = (SD_A/Mean_A) *100$

Where: SD_A = Standard deviation of well replicate absorbances Mean_A = Mean of well replicate absorbances

7. %CV for replicate absorbance values for *MDLs* must be ≤ 15.0%. If the %CV value of any MDL is > 15.0%, the MDL study is not acceptable. Corrective action and reanalysis are required.

- 8. %CV for replicate absorbance values for *samples* must be ≤ 15.0%. If the value of any sample is > 15.0% then reanalyze or qualify the results with the appropriate qualifier (J or UJ) and note in the final report.
- 9. Samples not analyzed within the required holding time must be recollected.

8.0 Qualifiers

- J This qualifier is applied only if the sample result is greater than the reporting limit (RL) and any of the following conditions apply:
 - Sample is collected in improper sample container.
 - Sample is received warm (>10°C).
 - LCRC or QCS are out of acceptance criteria.
- UJ This qualifier is applied only if the sample result is below the reporting limit (RL) and any of the following conditions apply:
 - Sample is collected in improper sample container.
 - Sample is received warm (>10°C).
 - Low LCRC recovery.

9.0 Required Documentation

- 9.1 The **Sample Bottle Cleaning Record** on page 10 of this SOP may be used to keep these records. The minimum requirements for documenting each procedure are as follows:
 - a. Analyst
 - b. Date Bottles Cleaned
 - c. Number Cleaned
 - d. Date Tested
 - e. Number Tested
 - f. Results: Number < Reporting Limit or Number > Reporting Limit
- 9.2 The **Reagent/Standard Preparation Record** on page 11 of this SOP may be used to keep records
- 9.3 The **Total Microcystins- ADDA Sample Preparation Record** on page 12 of this SOP may be used to keep these records. The minimum requirements for documenting each procedure are as follows:
 - a. Sample ID
 - b. Sample Location
 - c. Analyst(s)
 - d. Collection: Date and Time
 - e. pH Result (5-11)
 - f. Chlorine Result (<0.10mg/L)
 - g. 1st Freeze
 - h. 2nd Freeze
 - i. Final Freeze
- 9.4 The **Total Microcystins- ADDA Sample Results Record** on page 13 of this SOP may be used to keep these records. The minimum requirements for documenting each procedure are as follows:
 - a. Sample ID
 - b. Analyst

- c. Analysis: Date and Time
- d. Final Assay Calibration: Kit Lot #, R2, LRB, QCS, LCRC
- e. Sample Results from Report (mg/L)
- f. Reported Result (mg/L)
- g. Qualifiers Used
- 9.5 The **Total Microcystins- ADDA Method Detection Limit (MDL) Report** is available on our website at https://epa.ohio.gov/ddagw/labcert. It is not included in this manual since it has built-in formulas to calculate the MDL.

10.0 Optional Documentation

Documentation for daily temperature of refrigerator, freezer and water bath are not required for microcystin analysis. Should you wish to keep track of these, they are attached below.

- 10.1 The Daily Refrigerator Temperature Record, page 14, may be used to keep these records.
- 10.2 The **Daily Freezer Temperature Record**, page 15, may be used to keep these records.
- 10.3 The Water Bath Temperature Record, page 16, may be used to keep these records.

Sample Bottle Cleaning Record

To be recorded for each lot or batch cleaned

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Analyst	Date Bottles Cleaned	Number Cleaned	Date Tested	Results Number Tested Com	<u>.</u>		Comments ¹
	Cleaned	Cleaned			Number < Reporting Limit	Number > Reporting Limit	

¹Note action taken if results are unacceptable.

Reagent/Standard Receipt/Preparation Record

Supplier/Analyst(s) Initials	Type of Reagent/Standard	Reagent/Standard Lot Number	Date Received/Prepared	Reagent/Standard Expiration Date

Total Microcystins-ADDA Sample Preparation Record

Laboratory

Sample ID Sample Location		acation Analyst(a)		Collection		pH Result	Chlorine Result	1st Freeze ¹	2nd Freeze ¹	Final
Sample ID	Sample Location	n Analyst(s)	Date	Time	(5-11)	(<0.10 mg/L)				

¹Indicate completion with a check mark (no date or time required)

Total Microcystins-ADDA Sample Results Record

Laboratory:

		Analys	is		Final Ass	say Calibrati	on		Sampl e Result	Reporte d Result	Qualifier s Used	
Sample ID	Analyst	Date	Time	Kit Lot#	R²	LRB	QCS	LCR C	from Report (mg/L)	(mg/L)		Comments ¹

¹Note action taken if quality control was unacceptable.

To be recorded daily, 4.0 ± 2.0°C	To be	recorded	daily,	4.0	± 2	.0°C
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Laboratory	

Date	Temp (°C)	Comments	Analyst	Date	Temp (°C)	Comments

^{*}Note action taken if temperature is out of range.

Daily Freezer Temperature Record for HABs
To be recorded daily, <0.0°C

3 *

Laboratory	

Analyst	Date	Temp (°C)	Comments	Analyst	Date	Temp (°C)	Comments

^{*}Note action taken if temperature is out of range.

	Water Bath Temperature Record for HABs
	To be recorded daily, 35.0 ± 0.5°C
Laboratory	

Analyst	Date	Temp (°C)	Comments	Analyst	Date	Temp (°C)	Comments

^{*}Note action taken if temperature is out of range.