SW-846 is not intended to be an analytical training manual. Therefore, method procedures are written based on the assumption that they will be performed by analysts who are formally trained in at least the basic principles of chemical analysis and in the use of the subject technology.

In addition, SW-846 methods, with the exception of required method use for the analysis of method-defined parameters, are intended to be guidance methods which contain general information on how to perform an analytical procedure or technique which a laboratory can use as a basic starting point for generating its own detailed standard operating procedure (SOP), either for its own general use or for a specific project application. The performance data included in this method are for guidance purposes only, and are not intended to be and must not be used as absolute QC acceptance criteria for purposes of laboratory accreditation.

1.0 SCOPE AND APPLICATION

1.1 This method describes a procedure using a competitive immunoassay for the quantitative determination in water of triazine herbicides as the following compound:

<table>
<thead>
<tr>
<th>Analyte</th>
<th>CAS No. a</th>
</tr>
</thead>
<tbody>
<tr>
<td>Atrazine</td>
<td>1912-24-9</td>
</tr>
</tbody>
</table>

a Chemical Abstracts Service Registry Number

This method provides a single quantitative result, reported as atrazine, for all triazine herbicide compounds detected. However, the extent to which other triazine herbicides and other compounds are detected may vary between commercial testing products (see Secs. 1.4 and 4.0).

1.2 Testing products are commercially available from several manufacturers. The testing product evaluated by EPA for this method employs a competitive immunoassay. Other products differ in a number of respects, including the format of the test (tubes versus microtiter plates), the reagents used, and the specific steps in the test procedure.

1.3 The lower limit of quantitation submitted by the manufacturer of the testing product described in Sec. 6.2 was 0.03 µg/L for drinking water samples and is provided for guidance purposes only. The actual method sensitivity may be highly dependent on the sample matrix and may not always be achievable.

1.4 Since immunoassay methods use antibody molecules that can bind to more than the target analyte, an immunoassay has a tendency to overestimate the concentration of the target analyte when other analytes are present that may bind with the antibody. The commercially-available testing product evaluated for this method is based on an immunochemical reaction that will also respond to other triazine compounds. These other triazine compounds are often included in pesticide formulations containing atrazine. Thus, the
specificity of this procedure for atrazine is partly a function of the cross-reactivity of those other compounds (see Table 1). Therefore, as with other analytical techniques such as single-column gas chromatography, it is advisable to confirm positive test results near or above a regulatory action limit when the presence of other triazines is suspected.

1.5 Prior to employing this method, analysts are advised to consult the base method for each type of procedure that may be employed in the overall analysis (e.g., Method 4000) and the manufacturer’s instructions for additional information on quality control procedures, development of QC acceptance criteria, calculations, and general guidance. Analysts also should consult the disclaimer statement at the front of the manual and the information in Chapter Two for guidance on the responsibilities of the analyst for demonstrating that the techniques employed are appropriate for the analytes of interest, in the matrix of interest, and at the levels of concern.

In addition, analysts and data users are advised that, except where explicitly specified in a regulation, the use of SW-846 methods is not mandatory in response to Federal testing requirements. The information contained in this method is provided by EPA as guidance to be used by the analyst and the regulated community in making judgments necessary to generate results that meet the data quality objectives for the intended application.

1.6 Use of this method is restricted to use by, or under supervision of, analysts appropriately experienced and trained in the performance and interpretation of immunoassay methods. Each analyst must demonstrate the ability to generate acceptable results with this method.

2.0 SUMMARY OF METHOD

2.1 An accurately measured volume of sample (as little as 200 µL for some testing products) is mixed with a volume of enzyme-atrazine conjugate reagent in a test tube or a microtiter plate that has an anti-atrazine antibody immobilized on the surface, or in a vessel into which particles (magnetic particles for one testing product) with an immobilized antibody on the surface are added. The conjugate "competes" with the atrazine present in the sample for binding to the immobilized anti-atrazine antibody. The mixture is incubated at the temperature and for the length of time described in the manufacturer's instructions. (Testing products may employ other solid-phase support configurations, or even eliminate the solid-phase support. The summary here is intended to be generic and not limit the development of other testing products).

2.2 Unbound conjugate and sample analyte that may be present in the tubes or wells are removed by washing with organic-free reagent water or a wash solution specified by the manufacturer. A signal-generating substrate/chromogen reagent is added and the tube or plate is incubated as described in the manufacturer's instructions. In the case of the testing product described in Sec. 6.2, a magnetic field is applied to the tubes to retain the magnetic particle coated with antibody and any bound enzyme conjugate present during the wash step. (Other testing products may use different configurations.)

2.3 In an enzyme immunoassay, a stop solution is added to the tubes or wells of the plate to terminate the signal generating activity of the enzyme conjugate reagent. The absorbance is measured at a wavelength specified by the manufacturer. The test is interpreted by measuring the signal produced by a sample and determining the concentration from a dose-response curve constructed from standards tested at the same time. For a competitive immunoassay, the color (signal) developed during the test is inversely proportional to the concentration of atrazine in the sample.
3.0 DEFINITIONS

See Method 4000 and the glossary at the end of this method for definitions to basic immunoassay and procedure-specific terms. Also refer to the manufacturer's instructions and Chapter One for other definitions that may be relevant.

4.0 INTERFERENCES

4.1 Compounds that are chemically similar may cause a positive test result (false positive) for atrazine. This phenomenon is known as cross-reactivity. The testing product used in preparation of this method has been evaluated for cross-reactivity by the manufacturer. Table 1 provides the percent reactivity at which known cross-reactants will give a comparable response to that of atrazine when they are present in the sample.

4.1.1 The presence of cross-reacting compounds will result in an increase in the calculated concentration of the sample being analyzed and therefore will influence the incidence of false positive results. Thus, from the standpoint of monitoring compliance with a regulatory action limit, cross-reactivity is not a significant concern for test results below the action limit.

4.1.2 As with techniques such as single-column gas chromatography, in instances where the presence of other triazine compounds is known or suspected, it may be advisable to confirm positive results near or above the regulatory action limit using another analytical technique. However, false negative results are generally not a concern with immunoassay techniques.

4.2 Non-specific interferences such as sample pH, temperature, osmolarity, solvents, surfactants, and the presence of metal ions can effect immunoassay performance. Samples should be tested at the pH and temperature range specified by the testing product manufacturer. Review the product literature with regard to other potential interferences.

4.3 Storage temperatures may alter the useful life of the testing product reagents and supplies. Follow the manufacturer's directions for storage and use of all reagents and supplies.

5.0 SAFETY

5.1 This method does not address all safety issues associated with its use. The laboratory is responsible for maintaining a safe work environment and a current awareness file of OSHA regulations regarding the safe handling of the chemicals listed in this method. A reference file of material safety data sheets (MSDSs) should be available to all personnel involved in these analyses.

5.2 Some reagents may contain dilute acid solutions. Avoid contact with eyes, skin, and mucous membranes.

6.0 EQUIPMENT AND SUPPLIES

6.1 Each commercially-available testing product will supply or specify the apparatus and materials necessary for successful completion of the test. Most testing products supply the equipment and supplies specific to the immunoassay, including the tubes or plates containing
the immobilized antibody, and the immunochemical reagents. Do not mix the equipment, supplies, and reagents from the testing products for different analytes, or from the testing products from different manufacturers. Testing products contain immunochemical reagents that are evaluated by the manufacturer on a lot-specific basis. Do not mix the reagents from one lot with those from another lot unless expressly allowed by the manufacturer. Other equipment that may be necessary, but is not supplied with the testing product, includes common laboratory items such as precision pipetting devices, vortex mixers, etc.

6.2 The following immunoassay testing product was submitted to and evaluated by EPA, and found by the Agency to meet the performance specifications necessary for its intended application, i.e., drinking water monitoring:

Atrazine RaPID Assay® (Ohmicron Environmental Diagnostics, Inc.).

Additional testing products may be available from other manufacturers or in different formats. As additional testing products are evaluated by EPA and found to provide appropriate performance, information will be made available by the Office of Solid Waste regarding all of those testing products that are capable of meeting the performance requirements for this method. However, this procedure will not be revised solely to include information on such additional testing products. That information will be made available through the OSW Methods Team Website at www.epa.gov/SW-846.

7.0 REAGENTS AND STANDARDS

As with the equipment and supplies, each commercially-available testing product will supply or specify the reagents necessary for successful completion of the test. This includes the calibrators (standards) employed in the immunoassay. As noted in Sec. 6.1, do not mix the equipment, supplies, and reagents from the testing products for different analytes, or from the testing products of different manufacturers. Store all reagents and standards according to the manufacturer's instructions, and, where applicable, discard any which are past the expiration date assigned by the manufacturer or have been stored outside of the manufacturer's specified temperature range.

In addition, in order to demonstrate the method performance described in Sec. 9.0, the following reagents and standards will be necessary.

7.1 Organic-free reagent water -- All references to water in this method refer to organic-free reagent water, as defined in Chapter One. Organic-free reagent water is used for the preparation of the initial demonstration of capability test, the laboratory control sample, and other quality control tests. These tests are in addition to any control material(s) supplied by the manufacturer.

7.2 Atrazine spiking solution -- a solution of atrazine in a water-miscible solvent is necessary for spiking into organic-free reagent water to prepare the initial demonstration of proficiency test, the laboratory control sample, and other quality control tests. This solution may be provided by the manufacturer. If not provided, the laboratory should prepare a spiking solution or purchase one from a commercial source. Consult the manufacturer's instructions to identify solvents that may interfere with the testing product and do not use them. The concentration of this solution should be approximately 0.3 µg/mL, such that a 100 µL volume spiked into a 10 mL volume of reagent water will yield a concentration of 3 µg/L. Other volumes and concentrations may be employed, provided that the laboratory can demonstrate that the volume of solvent used does not affect the test performance and meets the quality objectives of the application for which the test is being used.
7.3 Solutions for adjusting the pH of samples before extraction, where such pH adjustment is specified by the manufacturer.

7.3.1 Sulfuric acid solution (1:1 v/v), \(H_2SO_4\) -- Slowly add 50 mL of \(H_2SO_4\) (sp. gr. 1.84) to 50 mL of organic-free reagent water.

7.3.2 Sodium hydroxide solution (2N), \(NaOH\) -- Dissolve 8 g of \(NaOH\) in organic-free reagent water and dilute to 100 mL.

8.0 SAMPLE COLLECTION, PRESERVATION, AND STORAGE

Sample collection, preservation, and storage considerations may vary by EPA program and may be specified in the regulation that requires compliance monitoring for a given contaminant. Where sample collection and handling considerations are specified in the regulation, follow those directions. Otherwise, use the following sample collection and handling guidelines.

8.1 Sample collection

The immunoassay testing products employ very small (<1 mL) sample volumes. Therefore, sample collection procedures should focus on the volume necessary to ensure that the sample represents the source.

8.1.1 Samples should be collected in pre-cleaned glass containers.

8.1.2 When sampling from a water tap, open the tap and allow the system to flush until the water temperature has stabilized (usually 2 to 5 mins). Adjust the flow to about 500 mL/min, and collect samples from the flowing stream. When sampling from an open body of water, fill the sample container with water from a representative area.

8.2 Sample preservation

8.2.1 If residual chlorine is present, add 80 mg of sodium thiosulfate per liter of sample to the bottle, prior to collecting the sample.

8.2.2 Retard microbiological degradation by adjusting the pH of the samples to <2 with hydrochloric acid at the time of sample collection. Before analysis, readjust the pH of the samples to the pH specified by the manufacturer with 2N \(NaOH\). The pH of the entire collected sample should be adjusted, not just the smaller volume utilized for the analysis.

8.3 Sample storage -- Samples should be stored at \(\#6\) EC until analysis, but must be warmed to the temperature specified by the manufacturer for analysis.

9.0 QUALITY CONTROL

9.1 Follow the manufacturer’s instructions for the quality control procedures specific to use of the testing product. Also, refer to Chapter One for additional guidance on quality assurance (QA) and quality control (QC) protocols that may be applicable. Any effort involving the collection of analytical data should include development of a structured and systematic planning document, such as a Quality Assurance Project Plan (QAPP) or a Sampling and Analysis Plan (SAP), which translates project objectives and specifications into directions for
those that will implement the project and assess the results. Each laboratory should maintain a formal quality assurance program. The laboratory should also maintain records to document the quality of the data generated. All data sheets and quality control data should be maintained for reference or inspection.

9.2 As noted in Sec. 1.2, the specific formats of the commercially-available testing products vary by manufacturer. As a result, those testing products evaluated and accepted by EPA represent performance-based analytical methods. Therefore, it is imperative that the manufacturer's instructions and specifications be followed. Follow the manufacturer's instructions for the testing product being used for quality control procedures specific to the testing product used. The following discussion of quality control requirements relies heavily on the analyst's knowledge and understanding of the manufacturer's instructions.

9.3 Routine quality control

Routine quality control procedures associated with this method include the analyses of standards, matrix spike samples, laboratory control samples, method blanks, and duplicate or replicate analyses (as specified by the manufacturer). All of the analyses described below must be conducted simultaneously, e.g., as part of the same batch of samples. A batch of samples consists of up to 20 field samples prepared and analyzed at the same time, or the maximum number of samples that can be analyzed along with the standards, controls, and other analyses specified by the manufacturer using a single testing product, whichever is fewer. The batch must include any duplicate or replicate analyses as well as any additional quality control tests specified by the manufacturer.

9.3.1 Calibration standards must be analyzed concurrently with each batch of samples processed.

9.3.2 Matrix spike (MS) samples should be analyzed with each batch of samples processed. The matrix spike samples should contain atrazine at the regulatory limit of interest (e.g., the MCL for the Drinking Water Program). The sample chosen for spiking should be representative of the field samples being analyzed.

9.3.3 The analyst must evaluate the accuracy of the assay by analyzing a laboratory control sample (LCS) consisting of organic-free reagent water sample spiked at the regulatory limit of concern for atrazine. Under the Drinking Water Program, the LCS must be spiked at 3 µg/L (the MCL for atrazine) with the spiking solution in Sec. 7.2. The mean recovery (bias) of the assay must be between 80 - 120%. If the manufacturer does not supply the spiking solution described in Sec. 7.2, or if another regulatory limit is relevant, then the laboratory is responsible for purchasing or preparing an appropriate spiking solution and performing this test. An LCS must be prepared and analyzed with each batch of samples analyzed, or per manufacturer's instructions.

NOTE: Spiking at 3 µg/L may require that the sample be diluted to be within the calibration range for some testing products, however, it provides data regarding the bias (if any) at the regulatory threshold, as well as indications of the analyst's proficiency at making dilutions.

9.3.4 A method blank, consisting of a volume of organic-free reagent water (see 7.1) equal to that of a field sample, must be analyzed with each batch of samples processed. The method blank should not contain any detectable atrazine.

9.3.5 Samples should be analyzed in duplicate or triplicate, as instructed by the manufacturer. The number of replicate analyses is specified by the manufacturer, and is a
function of the overall precision of the particular testing product. If the manufacturer
determines that, in order to achieve the precision claimed by the manufacturer, a given
number of replicate analyses must be performed, then the laboratory must employ the
specified number of replicate analyses.

9.4 Sample dilutions

If the sample concentration is outside of the calibrated range demonstrated by the initial
calibration and as specified by the manufacturer, then the sample must be diluted to within the
calibration range and retested. As employed in these testing products, the calibration range
specified by the manufacturer is based on a B/B0 in the 0.2 - 0.8 (20 - 80%) range. Given the
nature of the competitive immunoassay, the sample cannot be diluted after color development.
Thus, a diluted aliquot of the original sample must be prepared and analyzed.

NOTE: The B/B0 range of 0.2 - 0.8 is narrower than the simple concentration range of the
calibration standards. Therefore, the decision to dilute a sample for reanalysis must
be based on an evaluation of the B/B0 value of the sample, and not on a simple
comparison of the concentration in the sample and the highest standard in the
calibration.

9.5 Initial demonstration of proficiency

Each laboratory must demonstrate initial proficiency with the testing product that it utilizes,
by generating data of acceptable accuracy and precision for a reference sample containing
atrazine in a clean matrix. The laboratory must also repeat this demonstration whenever new
staff are trained or significant changes in instrumentation are made.

9.5.1 Prepare the reference sample from a spiking solution containing the
analyte of interest (see Sec. 7.2). Given the very small sample volume designated for the
immunoassay, a single 10-mL aliquot will provide sufficient volume for multiple tests and
will minimize the difficulties involved in spiking small volumes of organic-free reagent
water. Prepare a new aliquot each time the initial demonstration is to be performed.

9.5.2 Prepare an aliquot of organic-free reagent water, spiking it with the
solution in Sec. 7.2 to yield a concentration of 3 µg/L. Mix the aliquot well and allow the
spiked sample to stand for at least one hour.

9.5.3 Analyze at least four replicate subsamples of the spiked organic-free
reagent water aliquot using the same procedures used to analyze actual samples (Sec.
11.0). Analyze the number of replicates of each subsample specified by the manufacturer,
e.g., if the manufacturer specifies triplicate analyses of samples, then analyze 12
replicates (4 x 3) of the spiked sample.

9.5.4 Calculate the mean recovery (X), and the standard deviation of the
recoveries using the total number of replicate results, as described in Sec. 11.14.

9.5.5 Given the total number of replicate analyses performed, the mean
recovery (X) should be in the range of 90 - 110% and the relative standard deviation
should be no more than 10% of the mean recovery. If the results fall outside of these
acceptance limits, recheck all calculations. If no errors are found, repeat the
demonstration until the specifications are met.
9.6 Other quality control considerations

9.6.1 Do not use testing products past their expiration date.

9.6.2 Do not mix the equipment, supplies, and reagents from the testing products for different analytes, or from the testing products from different manufacturers.

9.6.3 Use the testing products within the storage temperature and operating temperature limits specified by the manufacturer.

10.0 CALIBRATION AND STANDARDIZATION

See Secs. 11.2 through 11.5 for calibration information related to this method.

11.0 PROCEDURE

Follow the manufacturer's instructions for the test being used. These instructions are summarized below, however, given the difference in test formats and reagents, the discussion is generic in nature. Where the manufacturer's instructions contradict these instructions or where these instructions do not apply to a specific testing product, follow the manufacturer's instructions.

11.1 Prepare the samples and standards

11.1.1 Bring samples, controls, and reagents to ambient temperature. Verify that the ambient temperature is consistent with the manufacturer's recommendations and limitations for this method. Do not attempt to perform tests outside of the temperature range specified by the manufacturer.

11.1.2 Check the pH of the samples. If necessary, adjust the pH to the range specified by the manufacturer, using 2N NaOH.

11.2 Initial calibration

The analyst must perform an initial calibration. This calibration is performed concurrently with the analysis of samples.

11.2.1 The initial calibration must consist of standards (calibrators) at a minimum of three concentrations that describe the quantitation range of the assay and should preferably span the regulatory limit of interest (e.g., for drinking water, the maximum contaminant level [MCL] is 3.0 µg/L). The standards must fall within the $B/B_0$ range of 0.2 to 0.8. The calibrators are generally provided by the product manufacturer. Calibration curves where all the calibrators are below the regulatory limit are allowed, but dilution and reanalysis of samples will be necessary when the sample concentration is near the regulatory limit. Calibration curves where all the calibrators are above the applicable regulatory limit may not be employed for compliance monitoring.

11.2.2 The testing product must also contain a "zero standard" or diluent solution that contains none of the target analytes. This solution is used to generate the $B_0$ value, but must not be used as one of the three standards required in Sec. 11.1.1.
11.2.3 When the entire dose response of a competitive immunoassay testing product (the absorbance of the solution or other signal specified by the manufacturer) is plotted on the y-axis against the concentration of the calibration standard on the x-axis, the resulting calibration curve will be hyperbolic when plotted on rectilinear paper, sigmoidal when plotted on semi-log paper, and linear when a Logit-log transformation of the data is employed and plotted on rectilinear paper. In addition, since the immunoassay is competitive, the blank (zero standard) will yield the highest response, with the color development inversely proportional to the standard concentration.

A plot of either the Logit B/B₀ or the Logit of the signal (absorbance units) versus the natural log of concentration is a widely used representation of the calibration data that generally yields a linear response curve. It is the basis of most computerized data analysis algorithms for competitive binding assays. The Logit B/B₀ is calculated according to the following formula:

\[
\text{Logit } \left( \frac{B}{B_0} \right) \cdot \log_e \left( \frac{B}{B_0} \right) \cdot \log_e \left( \frac{B}{B_0} \right)
\]

where:

- \(\log_e\) = Natural log or logarithm base e
- \(B\) = Response of the standard or sample
- \(B_0\) = Response of the zero standard

When Logit B/B₀ is plotted against the natural log (\(\log_e\)) of concentration, the results approximate a straight line with a negative slope (see Figure 1c). The transformed calibration data can then be characterized by the slope, intercept, correlation coefficient, and standard error of the line. The following sections describe the use of the Logit-log transformation of the data to prepare a calibration curve. Manufacturers may provide software that performs these calculations and, if provided, such software should be employed according to the manufacturer's instructions.

11.2.3.1 The commercially-available testing products may specify the analysis of standards in duplicate, or even in triplicate in some testing products. Thus, a three-point initial calibration may generate six to nine calibration points. Calculate the mean response (absorbance) at each concentration, and use this in all subsequent calculations.

11.2.3.2 Following the Logit B/B₀ and log transformations described in Sec. 11.2.3, construct a first order regression line (e.g., \(y = mx + b\)) using Logit B/B₀ as the dependent variable (y-axis) and the log-e concentration as the independent variable (x-axis). Since the slope of the line is negative, the regression cannot be forced through the origin, as the zero standard will yield the highest response and a value of 1.0 for B/B₀. The standards used to construct the regression line all must have B/B₀ values (prior to the Logit transformation) that fall within the 0.2-0.8 range.
The correlation coefficient of the regression (r) must be at least 0.98 in order to employ the calibration curve (manufacturers may provide more stringent linearity requirements for their testing products). If r is less than 0.98, check the expiration dates of all reagents, review the procedures to ensure that all standards were incubated for the same time specified by the manufacturer, and perform a new calibration.

11.3 By convention, the working range of an immunoassay calibration curve is defined as the range of $B/B_0$ from 0.2 to 0.8 (or $%B/B_0$ from 20% to 80%). Samples may be quantitated only within the working range of the curve.

11.4 As noted in Sec. 9.0, a new initial calibration curve must be constructed with each batch of samples assayed.

11.5 Calibration verification

Calibration verification is not performed in the traditional sense because the initial calibration standards are analyzed with each batch of samples each time the analyses are performed.

11.6 Prepare the spectrophotometer, photometer, or signal measurement equipment specified by the manufacturer.

11.7 Assay samples

11.7.1 Dispense the standards, controls, and samples into the container specified by the manufacturer. Be certain to include the replicate analyses specified by the manufacturer and the routine quality control samples described in Sec. 9.3.5 (also in replicate if samples are analyzed in replicate). Determine the maximum number of standards, controls, and samples that can be analyzed simultaneously and limit the number of field samples accordingly.

11.7.2 Dispense the enzyme conjugate reagent into each container as specified by the manufacturer.

11.7.3 Dispense the antibody capture reagent (where appropriate) as specified by the manufacturer.

11.7.4 Immunoassay methods employ kinetic and chromogenic reactions that are temperature sensitive. As a result, take care to perform the assay in the temperature range recommended by the manufacturer. Failure to follow temperature recommendations can lead to anomalous test results.

CAUTION: Do not attempt to process more samples simultaneously than specified by the manufacturer, as the additional processing time will lead to different incubation times for the samples and standards being tested and will produce erroneous results.

11.7.5 Wash each tube or well with washing reagents, as directed by the manufacturer.

11.7.6 Dispense the signal generating and signal terminating reagents (e.g., substrate/chromogen reagent and stop solutions) to each container in accordance with the manufacturer's instructions. Pay careful attention to the incubation times specified by the manufacturer.
manufacturer. Failure to follow incubation time recommendations can lead to erroneous results.

11.7.7 Interpret the test results within the time specified by the manufacturer. Follow the manufacturer's instructions for determining the sample concentration. For instance, read absorbance values (or optical density) at wavelength(s) specified by the manufacturer. Follow the manufacturer's quality control and data acceptance instructions.

11.8 As with the specific formats of the testing products and the reagents and supplies, the specifics of the applicable calculations may vary by manufacturer. Some testing products may provide measuring devices such as optical density readers or spectrophotometers and may include software for performing all the necessary calculations. Other testing products may require the analyst to plot results manually, using graph paper that may or may not be provided with the testing product, and determine sample results by interpolation from a standard curve. Whichever approach is used, the laboratory records (bench notes, etc.) should clearly indicate how the results were obtained and records specific to each determination, whether in hard copy or in electronic form, should be retained by the laboratory to substantiate the results.

11.9 Follow the manufacturer's instructions regarding calculation of all testing product results. Use the calibration curve generated concurrently with the sample analyses.

11.10 Where replicate test results are generated for samples or standards, calculate the mean concentration ($\bar{C}$) as:

$$\text{mean concentration } \bar{C} = \frac{1}{n} \sum_{i=1}^{n} C_i$$

where $C_i$ is the concentration in each replicate and $n$ is the number of replicate analyses.

11.11 For duplicate test results, calculate the relative percent difference (RPD) according to the following equation:

$$\text{RPD} = \left| \frac{C_1 - C_2}{\frac{C_1 + C_2}{2}} \right| \times 100$$

where $C_1$ and $C_2$ are the concentrations of the two replicate determinations.

11.12 When the manufacturer's instructions specify the analyses of three or more replicates, calculate the standard deviation (SD) and the relative standard deviation (RSD) of the replicate results for each sample, according to the following equations:
where \( C_i \) is the concentration in each replicate, \( C \) is the mean concentration, and \( n \) is the number of replicate analyses.

11.13 Accuracy is estimated from the recovery of spiked analytes from the matrix of interest. Laboratory performance in a clean matrix is estimated from the recovery of analytes in the LCS. Calculate the recovery of each spiked analyte in the matrix spike, matrix spike duplicate (if performed) and LCS according to the following formula.

\[
\text{Recovery} = \frac{C_s \times C_u}{C_n} \times 100
\]

where:
- \( C_s \) = Measured concentration of the spiked sample aliquot
- \( C_u \) = Measured concentration of the unspiked sample aliquot (use 0 for the LCS)
- \( C_n \) = Nominal (theoretical) concentration of the spiked sample aliquot

11.14 For the initial demonstration of proficiency (Sec. 9.5) calculate the mean recovery and the standard deviation of the recoveries, using the results from all replicate analyses of the four subsamples. Use the equation in Sec. 11.12 for standard deviation, substituting recovery for concentration.

12.0 DATA ANALYSIS AND CALCULATIONS

See Secs. 11.8 through 11.14 for information on data analysis and calculations.

13.0 METHOD PERFORMANCE

13.1 Performance data and related information are provided in SW-846 methods only as examples and guidance. The data do not represent required performance goals for users of the methods. Instead, performance goals should be developed on a project-specific basis, and the laboratory should establish in-house QC performance criteria for the application of this method. These performance data are not intended to be and must not be used as absolute QC acceptance criteria for purposes of laboratory accreditation.
13.2 In the case of this method (which may be used in either the field or the laboratory), any test kits used must be able to meet the performance specifications for the intended application. Also, follow the manufacturer's instructions for quality control procedures specific to the test kit used. Required performance criteria for a particular testing product may be included in the manufacturer's instructions.

13.3 Table 1 summarizes the cross-reactivity of other triazines relative to atrazine for the testing product listed in Sec. 6.2. Other testing products may have different cross-reactivity characteristics.

13.4 Table 2 summarizes the example results of a collaborative study of the immunoassay testing product described in Sec. 6.2 conducted under the auspices of the AOAC and described in Reference 3. These data are provided for guidance purposes only.

13.5 Figure 1 (1a - 1c) provides three example graphical representations of the calibration of atrazine using a competitive binding immunoassay such as those described here. These data are provided for guidance purposes only.

14.0 POLLUTION PREVENTION

14.1 Pollution prevention encompasses any technique that reduces or eliminates the quantity and/or toxicity of waste at the point of generation. Numerous opportunities for pollution prevention exist in laboratory operation. The EPA has established a preferred hierarchy of environmental management techniques that places pollution prevention as the management option of first choice. Whenever feasible, laboratory personnel should use pollution prevention techniques to address their waste generation. When wastes cannot be feasibly reduced at the source, the Agency recommends recycling as the next best option.

14.2 For information about pollution prevention that may be applicable to laboratories and research institutions consult Less is Better: Laboratory Chemical management for Waste Reduction available from the American Chemical Society, Department of Government Relations and Science Policy, 1155 16th Street, NW, Washington, DC, 20036, http://www.acs.org.

15.0 WASTE MANAGEMENT

The Environmental Protection Agency requires that laboratory waste management practices be conducted consistent with all applicable rules and regulations. The Agency urges laboratories to protect the air, water, and land by minimizing and controlling all releases from hoods and bench operations, complying with the letter and spirit of any sewer discharge permits and regulations, and by complying with all solid and hazardous waste regulations, particularly the hazardous waste identification rules and land disposal restrictions. For further information on waste management, consult The Waste Management Manual for Laboratory Personnel available from the American Chemical Society at the address listed in Sec. 14.2.

16.0 REFERENCES


17.0 TABLES, DIAGRAMS, FLOWCHARTS, AND VALIDATION DATA

The following pages contain the tables and figures referenced by this method. A glossary follows the tables and figures.
### TABLE 1
CROSS-REACTIVITY OF RAPID ASSAY TO RELATED COMPOUNDS

<table>
<thead>
<tr>
<th>Compound</th>
<th>CAS #</th>
<th>Percent Reactivity Relative to Atrazine</th>
</tr>
</thead>
<tbody>
<tr>
<td>Atrazine</td>
<td>1912-24-9</td>
<td>100</td>
</tr>
<tr>
<td>Ametryn</td>
<td>834-12-8</td>
<td>185</td>
</tr>
<tr>
<td>Prometryn</td>
<td>7287-19-6</td>
<td>113</td>
</tr>
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<td>Propazine</td>
<td>139-40-2</td>
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<td>Terbuthylazine</td>
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<td>&lt;0.1</td>
</tr>
<tr>
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<td>Spike Conc. (µg/L)(^1)</td>
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</tr>
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<td>---------------------------</td>
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<tr>
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\(^1\) Data for the two field-contaminated water samples represent the amount of atrazine added to the sample and the mean concentration and RSD data represent the amount found in excess of the background field contamination. Provided for guidance purposes only.

\(^2\) Recovery not calculated for unspiked samples.
FIGURE 1
EXAMPLE CALIBRATION DATA FROM A COMPETITIVE IMMUNOASSAY

Figure 1a - Generalized plot of immunoassay signal (test response) versus concentration of calibration standard (µg/L).

Figure 1b - Generalized plot of $B/(B_0-B)$ versus log concentration of calibration standard.
Figure 1c - Generalized plot of Logit \([B/(B_0-B)]\) versus log concentration of calibration standard.
Antibody -- A binding protein which is produced in response to an antigen, and which has the ability to bond with the antigen that stimulated its production.

\[
\% \frac{B}{B_0} = \frac{\text{Response of the standard or sample}}{\text{Response of the zero standard}} \times 100
\]

Competitive immunoassay -- An immunoassay method involving an in-vitro competitive binding reaction.

Cross-reactivity -- The relative concentration of an untargeted substance that would produce a response equivalent to a specified concentration of the targeted compound. In a quantitative immunoassay, it provides an indication of the concentration of cross-reactant that would produce a positive response. Cross-reactivity for individual compounds is often calculated as the ratio of target substance concentration to the cross-reacting substance concentration at 50% inhibition of the immunoassay's maximum signal times 100%.

Dose-response curve -- Representation of the signal generated by an immunoassay (y axis) plotted against the concentration of the target compound (x axis) in a series of standards of known concentration. When plotting a competitive immunoassay in a rectilinear format, the dose-response will have a hyperbolic character. When the log of concentration is used, the plot assumes a sigmoidal shape, and when the log of signal is plotted against the Logit transformation of concentration, a straight line plot is produced.

ELISA -- Enzyme-linked immunosorbent assay or ELISA is an enzyme immunoassay method that uses an immobilized reagent (e.g., antibody adsorbed to a plastic tube) to facilitate the separation of targeted analytes (antibody-bound components) from non-target substances (free reaction components) using a washing step, and an enzyme conjugate to generate the signal used for the interpretation of results.

Enzyme conjugate -- A molecule produced by the coupling of an enzyme molecule to an immunoassay component that is responsible for acting upon a substrate to produce a detectable signal.

Enzyme immunoassay -- An immunoassay method that uses an enzyme conjugate reagent to generate the signal used for interpretation of results. The enzyme mediated response may take the form of a chromogenic, fluorogenic, chemiluminescent or potentiometric reaction. (see Immunoassay and ELISA)

False negatives -- A negative interpretation of the method containing the target analytes at or above the quantitation level. Ideally, an immunoassay test product should produce no false negatives. The false negative rate can be estimated by analyzing split samples using both the test product and a reference method.

False positives -- A positive interpretation for a sample is defined as a positive response for a sample that contains analytes below the action level.
**Immunoperoxidase** -- An analytical technique that uses an antibody molecule as a binding agent in the detection and quantitation of substances in a sample (*see enzyme immunoassay and ELISA*).

**Immunogen** -- A substance having a minimum size and complexity, and that is sufficiently foreign to a genetically competent host to stimulate an immune response.

**Logit** -- A logarithmic transformation of data normalized to the highest observed response. For the competitive immunoassay described in this procedure, the Logit transformation is calculated as:

\[
\text{Logit } \left( \frac{B}{B_0} \right) = \log_e \left( \frac{B}{B_0} \right) - \log_e \left( \frac{B}{B_0} \right)
\]

**Natural log** -- The logarithm, base e, of a number. The natural logarithm may also be represented as "ln" or "log e."