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APPENDIX B3

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**QUALITY ASSURANCE OBJECTIVES AND DATA VALIDATION TECHNIQUES
FOR WASTE CHARACTERIZATION SAMPLING AND ANALYTICAL METHODS**

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3 **FOR WASTE CHARACTERIZATION SAMPLING AND ANALYTICAL METHODS**

4 B3-1 Validation Methods

5 The Permittees shall require the generator/storage sites (**sites**) to perform validation of all data
6 (qualitative as well as quantitative) so that data used for Waste Isolation Pilot Plant (**WIPP**)
7 compliance programs will be of known and acceptable quality. Validation includes a quantitative
8 determination of precision, accuracy, completeness, and method detection limits (as appropriate)
9 for analytical data (headspace Volatile Organics Compounds (**VOC**), total VOCs, Semivolatile
10 Organic Compounds (**SVOC**), and metals data). Quantitative data validations shall be performed
11 according to the conventional methods outlined below (equations B3-1 through B3-8). These
12 quantitative determinations will be compared to the Quality Assurance Objectives (**QAOs**)
13 specified in Sections B3-2 through B3-9. A qualitative determination of comparability and
14 representativeness will also be performed.

15 The qualitative data or descriptive information generated by radiography and visual examination
16 is not amenable to statistical data quality analysis. However, radiography and visual examination
17 are complementary techniques yielding similar data for determining the waste matrix code. The
18 waste matrix code is determined to ensure that the container is properly included in the
19 appropriate waste stream.

20 Data validation will be used to assess the quality of waste characterization data collected based
21 upon project precision, accuracy, completeness, comparability, and representativeness
22 objectives. These objectives are described below:

23 Precision

24 Precision is a measure of the mutual agreement among multiple measurements of a single
25 analyte, either by the same method or by different methods. Precision is either expressed as the
26 relative percent difference (**RPD**) for duplicate measurements or as the percent relative standard
27 deviation (**%RSD**) for three or more replicate measurements. For duplicate measurements, the
28 precision expressed as the RPD is calculated as follows:

29
$$RPD = \frac{C_1 - C_2}{\frac{(C_1 + C_2)}{2}} \times 100 \quad (B3-1)$$

30
31 where C_1 and C_2 are the two values obtained by analyzing the duplicate samples. C_1 is the larger
32 of the two observed values.

1 For three or more replicate measurements, the precision expressed as the %RSD is calculated as
2 follows:

$$3 \quad \%RSD = \frac{s}{y_{mean}} \times 100 \quad (B3-2)$$

4 where s is the standard deviation and y_{mean} is the mean of the replicate sample analyses.

5 The standard deviation, s , is calculated as follows:

$$6 \quad s = \sqrt{\frac{\sum_{i=1}^n (y_i - y_{mean})^2}{n - 1}} \quad (B3-3)$$

7 where y_i is the measured value of the i th replicate sample analysis measurement, and n equals the
8 number of replicate analyses.

9 Another aspect of precision is associated with analytical equipment calibration. In these
10 instances, the percent difference (**%D**) between multiple measurements of an equipment
11 calibration standard shall be calculated as follows:

$$12 \quad \%D = \frac{|C_1 - C_2|}{C_1} \times 100 \quad (B3-4)$$

13 where C_1 is the initial measurement and C_2 is the second or other additional measurement.

14 Accuracy

15 Accuracy is the degree of agreement between a measured analyte concentration (or the average
16 of replicate measurements of a single analyte concentration) and the true or known
17 concentration. Accuracy is determined as the percent recovery (**%R**).

18 For situations where a standard reference material is used, the %R is calculated as follows:

$$19 \quad \%R = \frac{C_m}{C_{srm}} \times 100 \quad (B3-5)$$

20 where C_m is the measured concentration value obtained by analyzing the sample and C_{srm} is the
21 “true” or certified concentration of the analyte in the sample.

1 For measurements where matrix spikes are used, the %R is calculated as follows:

$$2 \quad \%R = \frac{S - U}{C_{sc}} \times 100 \quad (B3-6)$$

3 where S is the measured concentration in the spiked aliquot, U is the measured concentration in
4 the unspiked aliquot, and C_{sc} is the actual concentration of the spike added.

5 Method Detection Limit

6 The method detection limit (**MDL**) is the minimum concentration of an analyte that can be
7 measured and reported with 99 percent confidence that the analyte concentration is greater than
8 zero. The MDL for all quantitative measurements (except for those using Fourier Transform
9 Infrared Spectroscopy [**FTIRS**]) is defined as follows:

$$10 \quad MDL = t_{(n-1, 1-\alpha=99)} \times s \quad (B3-7)$$

11 where $t_{(n-1, 1-\alpha=99)}$ is the t-distribution value corresponding to a 99 percent confidence level with n-
12 1 degrees of freedom, n is the number of observations, and s is the standard deviation of replicate
13 measurements.

14 For headspace-gas analysis using FTIRS, MDL is defined as follows:

$$15 \quad MDL = 3s \quad (B3-8)$$

16 where s is the standard deviation. Initially, a minimum of seven samples spiked at a level of three
17 to five times the estimated MDL and analyzed on non-consecutive days must be used to establish
18 the MDLs. MDLs should be updated using the results of the laboratory control sample or on-line
19 control samples.

20 Completeness

21 Completeness is a measure of the amount of valid data obtained from the overall measurement
22 system compared to the amount of data collected and submitted for analysis. Completeness must
23 be expressed as the number of samples analyzed with valid results as a percent of the total
24 number of samples submitted for analysis. Completeness, expressed as the percent complete
25 (**%C**), is calculated as follows:

$$26 \quad \%C = \frac{V}{n} \times 100 \quad (B3-9)$$

1 where V is the number of valid sampling or analytical results obtained and n is the number of
2 samples submitted for analysis.

3 Comparability

4 Comparability is the degree to which one data set can be compared to another. Comparability of
5 data generated at different sites will be ensured through the use of standardized, approved
6 testing, sampling, preservation, and analytical techniques and by meeting the QAOs specified in
7 Sections B3-2 through B3-9.

8 The comparability of waste characterization data shall be ensured through the use of
9 generator/storage site data usability criteria. The Permittees shall ensure that data usability
10 criteria are consistently established and used by the generator/storage sites to assess the usability
11 of analytical and testing data. The criteria shall address, as appropriate, the following:

- 12 • Definition or reference of criteria used to define and assign data qualifier flags based on
13 Quality Assurance Objective results,
- 14 • Criteria for assessing the useability of data impacted by matrix interferences,
- 15 • Criteria for assessing the useability of data based upon positive and negative bias as
16 indicated by quality control data, of data qualifiers, and qualifier flags,
- 17 • Criteria for assessing the useability of data due to
 - 18 • Severe matrix effects,
 - 19 • Misidentification of compounds,
 - 20 • Gross exceedance of holding times,
 - 21 • Failure to meet calibration or tune criteria
- 22 • Criteria for assessing the useability of data that does not meet minimum detection limit
23 requirements.

24 The Permittees shall be responsible for evaluating generator/storage site data useability and shall
25 assess implementation through the generator/storage site audit.

26 Representativeness

27 Representativeness is the degree to which sample data represent a characteristic of a population,
28 parameter variations at a sampling point, or an environmental condition. Representativeness is a
29 qualitative parameter that concerns the proper design of the sampling program.

30 Representativeness of waste containers from waste streams subjected to headspace gas,
31 homogeneous solids, and soil/gravel sampling and analysis will be validated, through
32 documentation, that a true random sample with an adequate population was identified and
33 collected consistent with Permit Attachment B2, Section B2-1. Since representativeness is a

1 quality characteristic that expresses the degree to which a sample or group of samples represents
2 the population being studied, the random selection of waste containers ensures representativeness
3 on a Program level. The Permittees shall require the Site Project Manager to document that the
4 selected waste containers from within a waste stream were randomly selected. Sampling
5 personnel shall verify that proper procedures are followed to ensure that samples are
6 representative of the waste contained in a particular waste container or a waste stream.

7 Identification of Tentatively Identified Compounds

8 In accordance with SW-846 convention, identification of compounds detected by gas
9 chromatography/mass spectrometry methods that are not on the list of target analytes shall be
10 reported. Both composited and individual container headspace gas, volatile analysis
11 (TCLP/Totals), and semi-volatile (TCLP/Totals) shall be subject to tentatively identified
12 compound (TIC) reporting. These TICs for GC/MS Methods are identified in accordance with
13 the following SW-846 criteria:

- 14 • Relative intensities of major ions in the reference spectrum (ions greater than 10% of the
15 most abundant ion) should be present in the sample spectrum.
- 16 • The relative intensities of the major ions should agree within ± 20 percent.
- 17 • Molecular ions present in the reference spectrum should be present in the sample
18 spectrum.
- 19 • Ions present in the sample spectrum but not in the reference spectrum should be reviewed
20 for possible background contamination or presence of coeluting compounds.
- 21 • Ions present in the reference spectrum but not in the sample spectrum should be reviewed
22 for possible subtraction from the sample spectrum because of background contamination
23 or coeluting peaks.
- 24 • The reference spectra used for identifying TICs shall include, at minimum, all of the
25 available spectra for compounds that appear in the 20.4.1.200 NMAC (incorporating 40
26 CFR Part 261) Appendix VIII list. The reference spectra may be limited to VOCs when
27 analyzing headspace gas samples.
- 28 • TICs for headspace gas analyses that are performed through FTIR analyses shall be
29 identified in accordance with the specifications of SW-846 Method 8410.

30 TICs shall be reported as part of the analytical batch data reports for GC/MS Methods in
31 accordance with the following minimum criteria:

- 32 • a TIC in an individual container headspace gas or solids sample shall be reported in the
33 analytical batch data report if the TIC meets the SW-846 identification criteria listed
34 above and is present with a minimum of 10% of the area of the nearest internal standard.

- 1 • a TIC in a composited headspace gas sample that contains 2 to 5 individual container
2 samples shall be reported in the analytical batch data report if the TIC meets the SW-846
3 identification criteria listed above and is present with a minimum of 2% of the area of the
4 nearest internal standard.

- 5 • a TIC in a composited headspace gas sample that contains 6 to 10 individual container
6 samples shall be reported in the analytical batch data report if the TIC meets the SW-846
7 identification criteria listed above and is present with a minimum of 1% of the area of the
8 nearest internal standard.

- 9 • a TIC in a composited headspace gas sample that contains 11 to 20 individual container
10 samples shall be reported in the analytical batch data report if the TIC meets the SW-846
11 identification criteria listed above and is present with a minimum of 0.5% of the area of
12 the nearest internal standard.

13 TICs that meet the SW-846 identification criteria, are reported in 25 percent of all waste
14 containers sampled from a given waste stream, and that appear in the 20.4.1.200 NMAC
15 (incorporating 40 CFR §261) Appendix VIII list, will be compared to acceptable knowledge data
16 to determine if the TIC is a listed waste in the waste stream. TICs identified through headspace
17 gas analyses that meet the Appendix VIII list criteria and the 25 percent reporting criteria for a
18 waste stream will be added to the headspace gas waste stream target list regardless of the
19 hazardous waste listing associated with the waste stream. TICs reported from the Totals VOC or
20 SVOC analyses may be excluded from the target analyte list for a waste stream if the TIC is a
21 constituent in an F-listed waste whose presence is attributable to waste packaging materials or
22 radiolytic degradation from acceptable knowledge documentation. If a listed waste constituent
23 TIC cannot be attributed to waste packaging materials, radiolysis, or other origins, the
24 constituent will be added to the target analyte list and new hazardous waste numbers will be
25 assigned, if appropriate. TICs subject to inclusion on the target analyte list that are toxicity
26 characteristic parameters shall be added to the target analyte list regardless of origin because the
27 hazardous waste designation for these numbers is not based on source. However, for toxicity
28 characteristic and non-toxic F003 constituents, the site may take concentration into account when
29 assessing whether to add a hazardous waste number. If a target analyte list for a waste stream is
30 expanded due to the presence of TICs, all subsequent samples collected from that waste stream
31 will be analyzed for constituents on the expanded list.

32 B3-2 Headspace-Gas Sampling

33 Quality Assurance Objectives

34 The precision and accuracy of the container headspace-gas sampling operations must be assessed
35 by analyzing field QC headspace-gas samples. These samples must include equipment blanks,
36 field reference standards, field blanks, and field duplicates. If the QAOs described below are not
37 met, a nonconformance report must be prepared, submitted, and resolved (Section B3-13).

1 Precision

2 The precision of the headspace-gas sampling and analysis operation must be assessed by
3 sequential collection of field duplicates for manifold sampling operations or simultaneous
4 collection of field duplicates for direct canister sampling operations for VOCs determination.
5 Corrective actions must be taken if the RPD exceeds 25 percent for any analyte found greater
6 than the PRQL in both of the duplicate samples.

7 Accuracy

8 A field reference standard must be collected using headspace-gas sampling equipment to assess
9 the accuracy of the headspace-gas sampling operation at a frequency of one field reference
10 standard for every 20 containers sampled or per sampling batch. Corrective action must be taken
11 if the %R of the field-reference standard is less than 70 or greater than 130.

12 Field blanks must also be collected at a frequency of 1 field blank for every 20 containers or
13 sampling batch sampled to assess possible contamination in the headspace gas sampling method.
14 Equipment blanks must also be collected at a frequency of 1 equipment blank for each
15 equipment cleaning batch to assess possible contamination in the equipment cleaning method.
16 Corrective actions must be taken if the blank exceeds three times the MDLs listed for any of the
17 compounds listed in Table B3-2.

18 Completeness

19 Sampling completeness shall be expressed as the number of valid samples collected as a percent
20 of the total number of samples collected for each waste stream. A valid sample is defined as a
21 sample collected in accordance with approved sampling methods and the container was properly
22 prepared for sampling (e.g., the polyliner was vented to the container headspace). The Permittees
23 shall require participating sampling facilities to achieve a minimum 90 percent completeness.
24 The amount and type of data that may be lost during the headspace-gas sampling operation
25 cannot be predicted in advance. The Permittees shall require the Site Project Manager to evaluate
26 the importance of any lost or contaminated headspace-gas samples and take corrective action as
27 appropriate.

28 Comparability

29 Consistent use and application of uniform procedures and equipment, as specified in Permit
30 Attachment B1 and application of data useability criteria, should ensure that headspace gas
31 sampling operations are comparable when sampling headspace at the different sampling
32 facilities. The Permittees shall require each site to take corrective actions if uniform procedures,
33 equipment, or operations are not followed without approved and justified deviations. In addition,
34 laboratories analyzing samples must successfully participate in the Performance Demonstration
35 Program (**PDP**) (DOE, 2003).

1 Representativeness

2 Specific headspace-gas sampling steps to ensure samples are representative include:

- 3 • Selection of the correct Drum Age Criteria (DAC) Scenario and waste packaging
4 configuration and meeting DAC equilibrium times.
- 5 • A sample canister cleaning and leak check after assembly
- 6 • Sampling equipment cleaning or disposal after use
- 7 • Sampling equipment leak check after sample collection
- 8 • Use of sample canisters with passivated internal surfaces
- 9 • Use of low-internal-volume sampling equipment
- 10 • Collection of samples with a low-sample volume to available headspace volume ratio
11 (less than 10 percent of the headspace when the headspace can be determined)
- 12 • Careful and documented pressure regulation of all activities specified in Attachment B1,
13 Section B1-1
- 14 • Performance audits
- 15 • Collection of equipment blanks, field reference standard, field blanks, and field
16 duplicates at the specified frequencies.
- 17 • Manifold pressure sensors and temperature sensors calibrated before initial use and
18 annually using NIST, or equivalent standards.
- 19 • OVA calibrated daily, prior to first use, or as necessary according to manufacturers
20 specifications.

21 Failure to perform the checks at the prescribed frequencies would result in corrective actions.

22 B3-3 Sampling of Homogeneous Solids and Soils/Gravel

23 Quality Assurance Objectives

24 To ensure that sampling is conducted in a representative manner on a waste-stream basis for
25 waste containers containing homogeneous solids and soil/gravel, samples must be collected
26 randomly in both the horizontal and vertical planes of each container's waste. For waste
27 containers that contain homogeneous solids and soil/gravel in smaller containers (e.g., 1 gal
28 [4.0 L] poly bottles) within the waste container, one randomly chosen smaller container must be
29 sampled from each container.

1 Precision

2 Sampling precision must be determined by collecting and sampling field duplicates (e.g., co-
3 located cores or co-located samples as described in Permit Attachment B1-2b(1)) once per
4 sampling batch or once per week during sampling operations, whichever is more frequent. A
5 sampling batch is a suite of homogeneous solids and soil/gravel samples collected consecutively
6 using the same sampling equipment within a specific time period. A sampling batch can be up to
7 20 samples (excluding field QC samples), all of which must be collected within 14 days of the
8 first sample in the batch. The Permittees shall require the Site Project Manager to calculate and
9 report the RPD between co-located core/samples.

10 The recommended method for establishing acceptance criteria for co-located cores and co-
11 located samples is the F-test method because the F-Test: 1) does not require potentially arbitrary
12 groupings into batches, 2) is based on exact distributions, and 3) is more likely to detect a change
13 in the process. When a sufficient number of samples are collected (25 to 30 pairs of co-located
14 cores or samples), control charts of the RPD will be developed for each constituent and for each
15 waste matrix or waste type (e.g., pyrochemical salts or organic sludges). The limits for the
16 control chart will be three standard deviations above or below the average RPD. Once
17 constructed, RPDs for additional co-located pairs will be compared with the control chart to
18 determine whether or not the co-located cores are acceptable. Periodically, the control charts will
19 be updated using all available data.

20 The statistical test will involve calculating the variance for co-located cores and samples by
21 pooling the variances computed for each pair of duplicate results. The variance for the waste
22 stream will be computed excluding any data from containers with co-located cores, because the
23 test requires the variance estimates to be independent. All data must be transformed to normality
24 prior to computing variances and performing the test. The test hypothesis is evaluated using the F
25 distribution and the method for testing the difference in variances.

26 Accuracy

27 Sampling accuracy through the use of standard reference materials shall not be measured.
28 Because waste containers containing homogeneous solids and soil/gravel with known quantities
29 of analytes are not available, sampling accuracy cannot be determined. However, sampling
30 methods and requirements described are designed to minimize sample degradation and hence
31 maximize sampling accuracy.

32 Sampling accuracy as a function of sampling cross-contamination will be measured. Equipment
33 blanks will be collected at a frequency of once per equipment cleaning batch. Corrective actions
34 must be taken if the blank exceeds three times the MDLs (PRDLs for metals) listed for any of the
35 compounds or analytes listed in Tables B3-4, B3-6, and B3-8. Equipment blanks will be
36 collected from the following equipment types:

- 37 • Fully assembled coring tools
- 38 • Liners cleaned separately from coring tools
- 39 • Miscellaneous sampling equipment that is reused (bowls, spoons, chisels)

1 Completeness

2 Sampling completeness shall be expressed as the number of valid samples collected as a percent
3 of the total number of samples collected for each waste stream. A valid sample is any sample that
4 is collected from a randomly selected container using randomly selected horizontal and vertical
5 planes in accordance with approved sampling methods. The Permittees shall require participating
6 sampling facilities to achieve a minimum 90 percent completeness.

7 Comparability

8 Consistent use and application of uniform procedures, sampling equipment, and measurement
9 units must ensure that sampling operations are comparable. Consistent application of data
10 useability criteria will also ensure comparability. In addition, the Permittees shall require
11 laboratories analyzing samples to successfully participate in the PDP (DOE, 2005).

12 Representativeness

13 Specific steps to ensure the representativeness of samples include the following for both waste
14 containers and smaller containers:

- 15
- Coring tools and sampling equipment must be clean prior to sampling.
 - The entire depth of the waste minus a site defined approved safety factor must be cored,
17 and the core collected must have a length greater than or equal to 50 percent of the depth
18 of the waste. This is called the core recovery and is calculated as follows:

19

$$\text{Core recovery (percent)} = \frac{y}{x} \times 100 \quad (\text{B3-10})$$

20 where

21 x = the depth of the waste in the container
22 y = the length of the core collected from the waste.

- 23
- Coring operations and tool selection should be designed to minimize alteration of the in-
24 place waste characteristics. Minimal waste disturbance must be verified by visually
25 examining the core and describing the observation (e.g., undisturbed, cracked, or
26 pulverized) in the field logbook.

27 If core recovery is less than 50 percent of the depth of the waste, a second coring location
28 shall be randomly selected. The core with the best core recovery shall be used for sample
29 collection.

1 One randomly selected container within a container will be chosen if the container contains
2 individual waste containers.

3 B3-4 Non Destructive Examination Methods

4 Quality Assurance Objectives

5 The QAOs for non destructive examination (**NDE**) are detailed in this section. NDE can be either
6 radiography or visual examination (**VE**). If the QAOs described below are not met, then
7 corrective action shall be taken. It should be noted that NDE does not have a specific MDL
8 because it is primarily a qualitative determination. The objective of NDE for the program is to
9 determine the physical waste form, the absence of prohibited items, and additional waste
10 characterization techniques that may be used based on the Summary Category Groups (i.e.,
11 S3000, S4000, S5000). The Permittees shall require each site to describe all activities required to
12 achieve these objectives in the site quality assurance project plan (**QAPjP**) and standard
13 operating procedures (**SOP**).

14 B3-4a Radiography

15 Data to meet these objectives must be obtained from a video and audio recorded scan provided
16 by trained radiography operators at the sites. Results must also be recorded on a radiography data
17 form. The precision, accuracy, completeness, and comparability objectives for radiography data
18 are presented below.

19 Precision

20 Precision is maintained by reconciling any discrepancies between two radiography operators
21 with regard to identification of the waste matrix code, liquids in excess of TSDF-WAC limits,
22 and compressed gases through independent replicate scans and independent observations.
23 Additionally, the precision of radiography is verified prior to use by tuning precisely enough to
24 demonstrate compliance with QAOs through viewing an image test pattern.

25 Accuracy

26 Accuracy is obtained by using a target to tune the image for maximum sharpness and by
27 requiring operators to successfully identify 100 percent of the required items in a training
28 container during their initial qualification and subsequent requalification.

29 Completeness

30 A video and audio media recording of the radiography examination and a validated radiography
31 data form will be obtained for 100 percent of the waste containers subject to radiography. All
32 video and audio media recordings and radiography data forms will be subject to validation as
33 indicated in Section B3-10.

1 Comparability

2 The comparability of radiography data from different operators shall be enhanced by using
3 standardized radiography procedures and operator qualifications.

4 B3-4b Visual Examination

5 Results must be recorded on a VE data form. The precision, accuracy, completeness, and
6 comparability objectives for VE data are presented below.

7 Precision

8 Precision is maintained by reconciling any discrepancies between the operator and the
9 independent technical reviewer with regard to identification of waste matrix code, liquids in
10 excess of TSDF-WAC limits, and compressed gases.

11 Accuracy

12 Accuracy is maintained by requiring operators to pass a comprehensive examination and
13 demonstrate satisfactory performance in the presence of the VE expert during their initial
14 qualification and subsequent requalification.

15 Completeness

16 A validated VE data form will be obtained for 100 percent of the waste containers subject to VE.

17 Comparability

18 The comparability of VE data from different operators shall be enhanced by using standardized
19 VE procedures and operator qualifications.

20 B3-5 Gas Volatile Organic Compound Analysis

21 Quality Assurance Objectives

22 The development of data quality objective (**DQOs**) specifically for this program has resulted in
23 the QAOs listed in Table B3-2. The specified QAOs represent the required quality of data
24 necessary to draw valid conclusions regarding program objectives. WAP-required limits, such as
25 the program required quantitation limits (**PRQL**) associated with VOC analysis, are specified to
26 ensure that the analytical data collected satisfy the requirements of all data users. A summary of
27 the Quality Control Samples and the associated acceptance criteria is included in Table B3-3.
28 Key data-quality indicators for laboratory measurements are defined below.

1 Precision

2 Precision shall be assessed by analyzing laboratory duplicates and replicate analyses of
3 laboratory-control samples and PDP blind-audit samples. Results from measurements on these
4 samples must be compared to the criteria listed in Table B3-2. These QC measurements will be
5 used to demonstrate acceptable method performance and to trigger corrective action when
6 control limits are exceeded.

7 Accuracy

8 Accuracy as %R shall be assessed for the laboratory operations by analyzing PDP blind-audit
9 samples and laboratory-control samples. Results from these measurements must be compared to
10 the criteria listed in Table B3-2. These QC measurements will be used to demonstrate acceptable
11 method performance and to trigger corrective action when control limits are exceeded.

12 Calibration

13 GC/MS Tunes, Initial Calibrations, and Continuing Calibration will be performed and evaluated
14 using the procedures and criteria specified in Table B3-3. These criteria will be used to
15 demonstrate acceptable calibration and to trigger corrective action when control limits are
16 exceeded.

17 Method Detection Limit

18 MDLs shall be expressed in nanograms for VOCs and must be less than or equal to those listed
19 in Table B3-2. MDLs shall be determined based on the method described in Section B3-1. The
20 detailed procedures for MDL determination shall be included in site SOPs.

21 Program Required Quantitation Limit

22 Laboratories must demonstrate the capability to quantitate analytes at or below the PRQLs given
23 in Table B3-2. Laboratories shall set the concentration of at least one calibration standard below
24 the PRQL. The detailed procedures for PRQL demonstration shall be included in laboratory
25 SOPs.

26 Completeness

27 Laboratory completeness shall be expressed as the number of samples analyzed with valid results
28 as a percent of the total number of samples submitted for analysis. A composited sample is
29 treated as one sample for the purposes of completeness, because only one sample is run through
30 the analytical instrument. Valid results are defined as results that meet the data useability criteria
31 based on application of the Quality Control Criteria specified in Tables B3-2 and B3-3; and meet
32 the detection limit, calibration representativeness, and comparability criteria within this section.
33 The Permittees shall require that participating laboratories meet the completeness criteria
34 specified in Table B3-2.

1 Comparability

2 For VOC analysis, data generated through analysis of samples from different sites shall be
3 comparable. The Permittees shall require each site to achieve comparability by using
4 standardized methods and traceable standards and by requiring all sites to successfully
5 participate in the PDP (DOE, 2003).

6 Representativeness

7 Representativeness for VOC analysis shall be achieved by collecting sufficient numbers of
8 samples using clean sampling equipment that does not introduce sample bias. Samples must be
9 collected as described in Permit Attachment B1.

10 B3-6 Total Volatile Organic Compound Analysis

11 Quality Assurance Objectives

12 The development of DQOs specifically for this program has resulted in the QAOs listed in Table
13 B3-4. The specified QAOs represent the required quality of data necessary to draw valid
14 conclusions regarding program objectives. WAP-required limits, such as the PRQL associated
15 with VOC analysis, are specified to ensure that the analytical data collected satisfy the
16 requirements of all data users. Key data-quality indicators for laboratory measurements are
17 defined below.

18 Precision

19 Precision shall be assessed by analyzing laboratory duplicates or matrix spike duplicates,
20 replicate analyses of laboratory control samples, and PDP blind-audit samples. Results from
21 measurements on these samples must be compared to the criteria listed in Table B3-4. These QC
22 measurements will be used to demonstrate acceptable method performance and to trigger
23 corrective action when control limits are exceeded.

24 Accuracy

25 Accuracy as %R shall be assessed for the laboratory operations by analyzing laboratory control
26 samples, matrix spikes, surrogate compounds, and PDP blind-audit samples. Results from these
27 measurements for matrix spikes samples must be compared to the %R criteria listed in Table B3-
28 4. Results for surrogates and internal standards are evaluated as specified in the SW-846 method
29 (EPA 1996) or Table B3-5. These QC measurements will be used to demonstrate acceptable
30 method performance and to trigger corrective action when control limits are exceeded.

31 Laboratory blanks shall be assessed to determine possible laboratory contamination and are
32 evaluated as specified in Table B3-5. These QC measurements will be used to demonstrate
33 acceptable levels of laboratory contamination and to trigger corrective action when control limits
34 are exceeded.

1 Calibration

2 GC/MS Tunes, Initial Calibrations, and Continuing Calibration will be performed and evaluated
3 using the procedures and criteria specified in Table B3-5 and the SW-846 method (EPA 1996).
4 These criteria will be used to demonstrate acceptable calibration and to trigger corrective action
5 when control limits are exceeded.

6 Method Detection Limit

7 MDLs shall be expressed in milligrams per kilogram (mg/kg) for VOCs and must be less than or
8 equal to those listed in Table B3-4. The detailed procedures for MDL determination shall be
9 included in site SOPs.

10 Program Required Quantitation Limit

11 Laboratories must demonstrate the capability to quantitate analytes in samples at or below the
12 PRQLs given in Table B3-4. Laboratories shall set the concentration of at least one calibration
13 standard below the PRQL. The detailed procedures for PRQL demonstration shall be included in
14 laboratory SOPs.

15 Completeness

16 Laboratory completeness shall be expressed as the number of samples analyzed with valid results
17 as a percent of the total number of samples submitted for analysis. Valid results are defined as
18 results that meet the data useability criteria based upon application of the Quality Control
19 Criteria specified in Tables B3-4 and B3-5 and meet the calibration, detection limit,
20 representativeness, and comparability criteria within this section. Participating laboratories must
21 meet the completeness criteria specified in Table B3-4.

22 Comparability

23 For VOC analysis, data generated through analysis of samples from different sites shall be
24 comparable. The Permittees shall require sites to achieve comparability by using standardized
25 SW-846 sample preparation and methods that meet the QAO requirements in Tables B3-4 and
26 B3-5, traceable standards, and by requiring all sites to successfully participate in the PDP (DOE,
27 2005). Generator/storage sites may use the most recent version of SW-846. Any changes to SW-
28 846 methodology that results in the elimination of sample preparation or analytical methods in
29 use at generator/storage sites must be addressed as a corrective action to address the
30 comparability of data before and after the SW-846 modification.

31 Representativeness

32 Representativeness for VOC analysis shall be achieved by collecting unbiased samples. Samples
33 must be collected as described in Permit Attachment B1.

1 B3-7 Total Semivolatile Organic Compound Analysis

2 Quality Assurance Objectives

3 The development of DQOs specifically for this program has resulted in the QAOs listed in Table
4 B3-6. The specified QAOs represent the required quality of data necessary to draw valid
5 conclusions regarding program objectives. WAP-required limits, such as the PRQLs, are
6 specified to ensure that the analytical data collected satisfy the requirements of all data users. A
7 summary of Quality Control Samples and associated acceptance criteria for this analysis is
8 included in Table B3-7. Key data-quality indicators for laboratory measurements are defined
9 below.

10 Precision

11 Precision shall be assessed by analyzing laboratory duplicates or matrix spike duplicates,
12 replicate analyses of laboratory control samples, and PDP blind-audit samples. Results from
13 measurements on these samples must be compared to the criteria listed in Table B3-6. These QC
14 measurements will be used to demonstrate acceptable method performance and to trigger
15 corrective action when control limits are exceeded.

16 Accuracy

17 Accuracy as %R shall be assessed for the laboratory operations by analyzing laboratory control
18 samples, matrix spikes, surrogate compounds, and PDP blind-audit samples. Results from these
19 measurements for matrix spikes samples must be compared to the %R criteria listed in Table B3-
20 6. Results for surrogates and internal standards are evaluated as specified in the SW-846 method
21 (EPA 1996) or Table B3-7. These QC measurements will be used to demonstrate acceptable
22 method performance and to trigger corrective action when control limits are exceeded.

23 Laboratory blanks shall be assessed to determine possible laboratory contamination and are
24 evaluated as specified in Table B3-7. These QC measurements will be used to demonstrate
25 acceptable levels of laboratory contamination and to trigger corrective action when control limits
26 are exceeded.

27 Calibration

28 GC/MS Tunes, Initial Calibrations, and Continuing Calibration will be performed and evaluated
29 using the procedures and criteria specified in Table B3-7 and the SW-846 method (EPA 1996).
30 These criteria will be used to demonstrate acceptable calibration and to trigger corrective action
31 when control limits are exceeded.

32 Method Detection Limit

33 MDLs shall be expressed in mg/kg for SVOCs and must be less than or equal to those listed in
34 Table B3-6. The detailed procedures for MDL determination shall be included in site SOPs.

1 Program Required Quantitation Limit

2 Laboratories must demonstrate the capability to quantitate analytes in samples at or below the
3 PRQLs given in Table B3-6. Laboratories shall set the concentration of at least one calibration
4 standard below the PRQL. The detailed procedures for PRQL demonstration shall be included in
5 laboratory SOPs.

6 Completeness

7 Laboratory completeness shall be expressed as the number of samples analyzed with valid results
8 as a percent of the total number of samples submitted for analysis. Valid results are defined as
9 results that meet the data useability criteria based on application of the Quality Control Criteria
10 specified in Tables B3-6 and B3-7 and meet the detection limit, calibration, representativeness,
11 and comparability criteria within this section. The Permittees shall require participating
12 laboratories to meet the level of completeness specified in Table B3-6.

13 Comparability

14 For SVOC analysis, data generated through analysis of samples from different sites shall be
15 comparable. The Permittees shall require sites to achieve comparability by using standardized
16 SW-846 sample preparation and methods that meet the QAO requirements in Tables B3-6 and
17 B3-7, traceable standards, and by requiring all sites to successfully participate in the PDP (DOE,
18 2005). Generator/storage sites may use the most current version of SW-846 if the methods are
19 consistent with QAO requirements. Any changes to SW-846 methodology that results in the
20 elimination of sample preparation or analytical methods in use at generator/storage sites must be
21 addressed as a corrective action to address the comparability of data before and after the SW-846
22 modification.

23 Representativeness

24 Representativeness for SVOC analysis shall be achieved by collecting unbiased samples.
25 Samples must be collected as described in Permit Attachment B1.

26 B3-8 Total Metal Analysis

27 Quality Assurance Objectives

28 The development of DQOs for the program has resulted in the QAOs listed in Table B3-8. The
29 specified QAOs represent the required quality of data necessary to draw valid conclusions
30 regarding program objectives. WAP-required limits, such as the PRQLs associated with metal
31 analysis, are specified to ensure that the analytical data collected satisfy the requirements of all
32 data users. A summary of Quality Control Samples and the associated acceptance criteria for this
33 analysis is provided in Table B3-9. Key data-quality indicators for laboratory measurements are
34 defined below.

1 Precision

2 Precision shall be assessed by analyzing laboratory sample duplicates or laboratory matrix spike
3 duplicates, replicate analyses of laboratory-control samples, and PDP blind-audit samples.
4 Results from measurements on these samples must be compared to the criteria listed in Table
5 B3-8. These QC measurements will be used to demonstrate acceptable method performance and
6 to trigger corrective action when control limits are exceeded.

7 Accuracy

8 Accuracy shall be assessed through the analysis of laboratory matrix spikes, PDP blind-audit
9 samples, serial dilutions, interference check samples, and laboratory-control samples. Results
10 from these measurements must be compared to the criterion listed in Table B3-8 and B3-9. These
11 QC measurements will be used to demonstrate acceptable method performance and to trigger
12 corrective action when control limits are exceeded.

13 Laboratory blanks and calibration blanks shall be assessed to determine possible laboratory
14 contamination and are evaluated as specified in Table B3-9. These QC measurements will be
15 used to demonstrate acceptable levels of laboratory contamination and to trigger corrective
16 action when control limits are exceeded.

17 Calibration

18 Mass Tunes (for ICP MS only), Standards Calibration, Initial Calibration verifications, and
19 Continuing Calibrations will be performed and evaluated using the procedures and criteria
20 specified in Table B3-9 and the SW-846 method (EPA 1996). These criteria will be used to
21 demonstrate acceptable calibration and to trigger corrective action when control limits are
22 exceeded.

23 Program Required Detection Limits

24 PRDLs, expressed in units of micrograms per L ($\mu\text{g/L}$), are the maximum values for instrument
25 detection limits (**IDL**) permissible for program support under the WAP. IDLs must be less than
26 or equal to the PRDL for the method used to quantitate a specific analyte. Any method listed in
27 Table B-5 of the Waste Analysis Plan (Permit Attachment B) may be used if the IDL meets this
28 criteria. For high concentration samples, an exception to the above requirements may be made in
29 cases where the sample concentration exceeds five times the IDL of the instrument being used.
30 In this case, the analyte concentration may be reported even though the IDL may exceed the
31 PRDL. IDLs shall be determined semiannually (i.e., every six months). Detailed procedures for
32 IDL determination shall be included in laboratory SOPs.

33 Program Required Quantitation Limit

34 The Permittees shall require participating laboratories to demonstrate the capability of analyte
35 quantitation at or below the PRQLs in units of mg/kg wet weight (given in Table B3-8). The

1 PRDLs are set an order of magnitude less than the PRQLs (assuming 100 percent solid sample
2 diluted by a factor of 100 during preparation). The Permittees shall require participating
3 laboratories to set the concentration of at least one QC or calibration standard at or below the
4 solution concentration equivalent of the PRQL. Detailed calibration procedures shall be included
5 in site SOPs.

6 Completeness

7 Laboratory completeness shall be expressed as the number of samples analyzed with valid results
8 as a percent of the total number of samples submitted for analysis. Valid results are defined as
9 results that meet the data useability criteria based upon application of the Quality Control
10 Criteria specified in Tables B3-8 and B3-9 and meet the detection limit, calibration,
11 representativeness, and comparability criteria within this section. The Permittees shall require
12 participating laboratories to meet the completeness specified in Table B3-8.

13 Comparability

14 For metals analysis, data generated through analysis of samples from different sites shall be
15 comparable. Comparability will be achieved by using standardized SW-846 sample preparation
16 and methods that meet QAO requirements in Tables B3-8 and B3-9, demonstrating successful
17 participation in the PDP (DOE, 2005), and use of traceable standards. Generator/storage sites
18 may use the most recent SW-846 update. Any changes to SW-846 methodology that results in
19 the elimination of sample preparation or analytical methods in use at generator/storage sites must
20 be addressed as a corrective action to address the comparability of data before and after the SW-
21 846 modification.

22 Representativeness

23 Representativeness for metals analysis shall be achieved by the collection of unbiased samples
24 and the preparation of samples in the laboratory using representative and unbiased methods.
25 Samples must be collected as described in Permit Attachment B1.

26 B3-9 Acceptable Knowledge

27 Acceptable knowledge documentation provides primarily qualitative information that cannot be
28 assessed according to specific data quality goals that are used for analytical techniques. QAOs
29 for analytical results are described in terms of precision, accuracy, completeness, comparability,
30 and representativeness. Appropriate analytical and testing results may be used to augment the
31 characterization of wastes based on acceptable knowledge. To ensure that the acceptable
32 knowledge process is consistently applied, the Permittees shall require sites to comply with the
33 following data quality requirements for acceptable knowledge documentation:

- 34 • Precision - Precision is the agreement among a set of replicate measurements without
35 assumption of the knowledge of a true value. The qualitative determinations, such as
36 compiling and assessing acceptable knowledge documentation, do not lend themselves to

1 statistical evaluations of precision. However, the acceptable knowledge information will
2 be addressed by the independent review of acceptable knowledge information during
3 internal and external audits.

- 4 • Accuracy - Accuracy is the degree of agreement between an observed sample result and
5 the true value. The percentage of waste containers which require reassignment to a new
6 waste matrix code and/or designation of different hazardous waste numbers based on
7 sampling and analysis data and discrepancies identified by the Permittees during waste
8 confirmation will be reported as a measure of acceptable knowledge accuracy.

- 9 • Completeness - Completeness is an assessment of the number of waste streams or
10 number of samples collected to the number of samples determined to be useable through
11 the data validation process. The acceptable knowledge record must contain 100 percent
12 of the required information (Permit Attachment B4-3). The useability of the acceptable
13 knowledge information will be assessed for completeness during audits.

- 14 • Comparability - Data are considered comparable when one set of data can be compared to
15 another set of data. Comparability is ensured through sites meeting the training
16 requirements and complying with the minimum standards outlined for procedures that are
17 used to implement the acceptable knowledge process. All sites must assign hazardous
18 waste numbers in accordance with Permit Attachment B4-3b and provide this
19 information regarding its waste to other sites who store or generate a similar waste
20 stream.

- 21 • Representativeness - Representativeness expresses the degree to which sample data
22 accurately and precisely represent characteristics of a population. Representativeness is a
23 qualitative parameter that will be satisfied by ensuring that the process of obtaining,
24 evaluating, and documenting acceptable knowledge information is performed in
25 accordance with the minimum standards established in Permit Attachment B4. Sites also
26 must assess and document the limitations of the acceptable knowledge information used
27 to assign hazardous waste numbers (e.g., purpose and scope of information, date of
28 publication, type and extent to which waste parameters are addressed).

29 The Permittees shall require each generator/storage site to comply with the nonconformance
30 notification and reporting requirements of Section B3-13 if the results of sampling and analysis
31 specified in Permit Attachment B are inconsistent with acceptable knowledge documentation.

32 The Permittees shall require each site to address quality control by tracking its performance with
33 regard to the use of acceptable knowledge by: 1) assessing the frequency of inconsistencies
34 among information, and 2) documenting acceptable knowledge inconsistencies identified
35 through radiography, visual examination, headspace-gas analyses, and solidified waste analyses.
36 In addition, the acceptable knowledge process and waste stream documentation must be
37 evaluated through internal assessments by generator/storage site quality assurance organizations
38 and assessments by auditors external to the organization (i.e., the Permittees).

1 B3-10 Data Review, Validation, and Verification Requirements

2 Procedures shall be developed for the review, validation, and verification of data at the data
3 generation level; the validation and verification of data at the project level; and the verification
4 of data at the Permittee level. Data review determines if raw data have been properly collected
5 and ensures raw data are properly reduced. Data validation verifies that the data reported satisfy
6 the requirements of this WAP and is accompanied by signature release. Data verification
7 authenticates that data as presented represent the sampling and analysis activities as performed
8 and have been subject to the appropriate levels of data review. The requirements presented in this
9 section ensure that WAP records furnish documentary evidence of quality.

10 The Permittees shall require the sites to generate the following Batch Data Reports for data
11 validation, verification, and quality assurance activities:

- 12 • A Testing Batch Data Report or equivalent includes all data pertaining to radiography or
13 visual examination for up to 20 waste containers without regard to waste matrix. Table
14 B3-11 lists all of the information required in Testing Batch Data Reports (identified with
15 an “X”) and other information that is necessary for data validation, but is optional in
16 Testing Batch Data Reports (identified with an “O”).
- 17 • A Sampling Batch Data Report or equivalent includes all sample collection data
18 pertaining to a group of no more than 20 headspace gas or homogeneous waste samples
19 that were collected for chemical analysis. Table B3-12 lists all of the information
20 required in Sampling Batch Data Reports (identified with an “X”) and other information
21 that is necessary for data validation, but is optional in Sampling Batch Data Reports
22 (identified with an “O”).
- 23 • An Analytical Batch Data Report or equivalent includes analytical data from the analysis
24 of TRU-mixed waste for up to 20 headspace gas or homogeneous waste samples.
25 Analytical Batch Data Reports or equivalent that contain results for composited
26 headspace gas samples must contain sufficient information to identify the containers that
27 were composited for each composite sample and the sample volume that was taken from
28 each waste container. Because Analytical Batch Data Reports are generated based on the
29 number of samples analyzed, an Analytical Batch Data Report may contain results that
30 are applicable to more than 20 containers depending on how many composite samples are
31 part of the report, but may not exceed a total of 20 samples analyzed. Table B3-13 lists
32 all of the information required in Analytical Batch Data Reports (identified with an “X”) and
33 other information that is necessary for data validation, but is optional in Analytical
34 Batch Data Reports (identified with an “O”).

35 Raw analytical data need not be included in Analytical Batch Data Reports, but must be
36 maintained in the site project files and be readily available for review upon request. Raw
37 data may include all analytical bench sheet and instrumentation readouts for all
38 calibration standard results, sample data, QC samples, sample preparation conditions and
39 logs, sample run logs, and all re-extraction, re-analysis, or dilution information pertaining

1 to the individual samples. Raw data may also include calculation records and any
2 qualitative or semi-quantitative data collected for a sample and that has been recorded on
3 a bench sheet or in a log book.

- 4 • An On-line Batch Data Report or equivalent contains the combined information from the
5 Sampling Batch Data Report and Analytical Batch Data Report that is relevant to the on-
6 line method used.

7 B3-10a Data Generation Level

8 The following are minimum requirements for raw data collection and management which the
9 Permittees shall require for each site:

- 10 • All raw data shall be signed and dated in reproducible ink by the person generating it.
11 Alternately, unalterable electronic signatures may be used.
- 12 • All data must be recorded clearly, legibly, and accurately in field and laboratory records
13 (bench sheets, logbooks), and include applicable sample identification numbers (for
14 sampling and analytical labs).
- 15 • All changes to original data must be lined out, initialed, and dated by the individual
16 making the change. A justification for changing the original data may also be included.
17 Original data must not be obliterated or otherwise disfigured so as not to be readable.
18 Data changes shall only be made by the individual who originally collected the data or an
19 individual authorized to change the data.
- 20 • All data must be transferred and reduced from field and laboratory records completely
21 and accurately.
- 22 • All field and laboratory records must be maintained as specified in Table B-6 of
23 Attachment B.
- 24 • Data must be organized into a standard format for reporting purposes (Batch Data Report),
25 as outlined in specific sampling and analytical procedures.
- 26 • All electronic and video data must be stored appropriately to ensure that waste container,
27 sample, and associated QC data are readily retrievable. In the case of classified
28 information, additional security provisions may apply that could restrict retrievability.
29 The additional security provisions will be documented in generator/storage site
30 procedures as outlined in the QAPjP in accordance with prevailing classified information
31 security standards.

1 Data review, validation, and verification at this level involves scrutiny and signature release from
2 qualified independent technical reviewer(s)¹ as specified below. Individuals conducting this data
3 review, validation, and verification must use checklists that address all of the items included in
4 this section. Checklists must contain or reference tables showing the results of sampling,
5 analytical or on-line batch QC samples, if applicable. Checklists must reflect review of all QC
6 samples and quality assurance objective categories in accordance with criteria established in
7 Tables B3-2 through B3-9 (as applicable to the methods validated). Completed checklists must
8 be forwarded with Batch Data Reports to the project level. Analytical raw data must be available
9 and reviewed by the data generation level reviewer.

10 B3-10a(1) Independent Technical Review

11 The independent technical review ensures by review of raw data that data generation and
12 reduction are technically correct; calculations are verified correct; deviations are documented;
13 and QA/QC results are complete, documented correctly, and compared against WAP criteria.
14 This review validates and verifies all of the work documented by the originator.

15 One hundred percent of the Batch Data Reports must receive an independent technical review.
16 This review shall be performed by an individual other than the data generator who is qualified to
17 have performed the initial work. The independent technical review must be performed as soon as
18 practicably possible in order to determine and correct negative quality trends in the sampling or
19 analytical process. However at a minimum, the independent technical review must be performed
20 before any waste associated with the data reviewed is managed, stored, or disposed at WIPP,
21 unless the data are being obtained from waste sampling and analysis as containers are being
22 retrieved or generated after initial WSPF approval as described in Attachment B2, Section B2-1.
23 The reviewer(s) must release the data as evidenced by signature, and as a consequence ensure the
24 following:

- 25 • Data generation and reduction were conducted in a technically correct manner in
26 accordance with the methods used (procedure with revision). Data were reported in the
27 proper units and correct number of significant figures.
- 28 • Calculations have been verified by a valid calculation program, a spot check of verified
29 calculation programs, and/or 100 percent check of all hand calculations. Values that are
30 not verifiable to within rounding or significant difference discrepancies must be rectified
31 prior to completion of independent technical review.
- 32 • The data have been reviewed for transcription errors.
- 33 • The testing, sampling, or analytical data QA documentation for Batch Data Reports is
34 complete and includes, as applicable, raw data, DAC and equilibrium calculations and

¹ Independent technical review is performed by a competent individual who is not directly responsible for performing the work.

1 times, calculation records, chain-of-custody (COC) forms, calibration records (or
2 references to an available calibration package), QC sample results, and copies or originals
3 of gas canister sample tags. Corrective action will be taken to ensure that all Batch Data
4 Reports are complete and include all necessary raw data prior to completion of the
5 independent technical review.

6 • QC sample results are within established control limits, and if not, the data have been
7 appropriately qualified in accordance with data useability criteria. Data outside of
8 established control limits will be qualified as appropriate, assigned an appropriate
9 qualifier flag, discussed in the case narrative, and included as appropriate in calculations
10 for completeness. QC criteria that were not met are documented.

11 • Reporting flags (Table B3-14) were assigned correctly.

12 • Sample holding time and preservation requirements were met, or exceptions documented.

13 • Radiography tapes have been reviewed (independent observation) on a waste container
14 basis at a minimum of once per testing batch or once per day of operation, whichever is
15 less frequent (Attachment B1, Section B1-3). The radiography tape will be reviewed
16 against the data reported on the radiography form to ensure that the data are correct and
17 complete.

18 • Field sampling records are complete. Incomplete or incorrect field sampling records will
19 be subject to resubmittal prior to completion of the independent technical review.

20 • QAOs have been met according to the methods outlined in Sections B3-2 through B3-9.

21 B3-10b Project Level

22 Data validation and verification at this level involves scrutiny and signature release from the Site
23 Project Manager (or designee). The Permittees shall require each site to meet the following
24 minimum requirements for each waste container. Any nonconformance identified during this
25 process shall be documented on a nonconformance report (Section B3-13).

26 The Site Project Manager shall ensure that a repeat of the data generation level review,
27 validation, and verification is performed on the data for a minimum of one randomly chosen
28 waste container quarterly (every three months). This exercise will document that the data
29 generation level review, validation, and verification is being performed according to
30 implementing procedures.

31 B3-10b(1) Site Project Manager Review

32 The Site Project Manager Review is the final validation that all of the data contained in Batch
33 Data Reports from the data generation level are complete and have been properly reviewed as
34 evidenced by signature release and completed checklists.

- 1 One hundred percent of the Batch Data Reports must have Site Project Manager signature
2 release. At a minimum, the Site Project Manager signature release must be performed before any
3 waste associated with the data reviewed is managed, stored, or disposed at WIPP, unless the data
4 are being obtained from waste sampling and analysis as containers are being retrieved or
5 generated as described in Permit Attachment B2, Section B2-1. This signature release must
6 ensure the following:
- 7 • The validity of the DAC assignment made at the data generation level based upon an
8 assessment of the data collection and evaluation necessary to make the assignment.
 - 9 • Testing batch QC checks (e.g., replicate scans, measurement system checks) were
10 properly performed. Radiography data are complete and acceptable based on evidence of
11 videotape review of one waste container per day or once per testing batch, whichever is
12 less frequent, as specified in B1-3.
 - 13 • Sampling batch QC checks (e.g., equipment blanks, field duplicates, field reference
14 standards) were properly performed, and meet the established QAOs and are within
15 established data useability criteria.
 - 16 • Analytical batch QC checks (e.g., laboratory duplicates, laboratory blanks, matrix spikes,
17 matrix spike duplicates, laboratory control samples) were properly performed and meet
18 the established QAOs and are within established data useability criteria.
 - 19 • On-line batch QC checks (e.g., field blanks, on-line blanks, on-line duplicates, on-line
20 control samples) were properly performed and meet the established QAOs and are within
21 established data useability criteria.
 - 22 • Proper procedures were followed to ensure representative samples of headspace gas and
23 homogeneous solids and soil/gravel were taken.
 - 24 • Data generation level independent technical review, validation, and verification have
25 been performed as evidenced by the completed review checklists and appropriate
26 signature releases.
 - 27 • Batch data review checklists are complete.
 - 28 • Batch Data Reports are complete and data are properly reported (e.g., data are reported in
29 the correct units, with the correct number of significant figures, and with qualifying
30 flags).
 - 31 • Verify that data are within established data assessment criteria and meet all applicable
32 QAOs (Sections B3-2 through B3-9).

1 B3-10b(2) Prepare Site Project Manager Summary and Data Validation Summary

2 To document the project-level validation and verification described above, the Permittees shall
3 require each Site Project Manager (or designee) to prepare a Site Project Manager Summary and
4 a Data Validation Summary. These reports may be combined to eliminate redundancy. The Site
5 Project Manager Summary includes a validation checklist for each Batch Data Report. Checklists
6 for the Site Project Manager Summary must be sufficiently detailed to validate all aspects of a
7 Batch Data Report that affect data quality. The Data Validation Summary provides verification
8 that, on a per waste container or sample basis as evidenced by Batch Data Report reviews, all
9 data have been validated in accordance with the site QAPjP. The Data Validation Summary must
10 identify each Batch Data Report reviewed (including all waste container numbers), describe how
11 the validation was performed and whether or not problems were detected (e.g., nonconformance
12 reports), and include a statement indicating that all data are acceptable. Summaries must include
13 release signatures.

14 Once the data have received project-level validation and verification or when the Site Project
15 Manager decides the sample no longer needs to be retained, the Site Project Manager must
16 ensure that the laboratory is notified. Samples must be retained by the laboratory until this
17 notification is received. Gas sample canisters may then be released from storage for cleaning,
18 recertification, and subsequent reuse. Sample tags must be removed and retained in the project
19 files before recycling the canisters. If the Site Project Manager requests that samples or canisters
20 be retained for future use (e.g., an experimental holding time study), the same sample
21 identification and COC forms shall be used and cross-referenced to a document which specifies
22 the purpose for sample or canister retention.

23 B3-10b(3) Prepare Waste Stream Characterization Package

24 In the event the Permittees request detailed information on a waste stream, the Site Project
25 Manager will provide a Waste Stream Characterization Package. The Site Project Manager must
26 ensure that the Waste Stream Characterization Package (Section B3-12b(3)) will support waste
27 characterization determinations.

28 B3-10c Permittee Level

29 The final level of data verification occurs at the Permittee level and must, at a minimum, consist
30 of reviewing a sample of the Batch Data Reports during audits of generator/storage sites and
31 Permittee approved laboratories to verify completeness. During such audits, the Permittees are
32 responsible for the verification that Batch Data Reports include the following:

- 33 • Project-level signature releases
- 34 • Listing of all waste containers being presented in the report
- 35 • Listing of all testing, sampling, and analytical batch numbers associated with each waste
36 container being reported in the package

- 1 • Analytical Batch Data Report case narratives
- 2 • Site Project Manager Summary
- 3 • Data Validation Summary
- 4 • Complete summarized qualitative and quantitative data for all waste containers with data
- 5 flags and qualifiers.

6 For each Waste Stream Profile Form (**WSPF**) submitted for approval, the Permittees must verify
7 that each submittal (i.e., WSPF and Characterization Information Summary) is complete and
8 notify the originating site in writing of the WSPF approval. The Permittees will maintain the data
9 as appropriate for use in the regulatory compliance programs. For subsequent shipments made
10 after the initial WSPF approval, the verification will also include WWIS internal limit checks
11 (Attachment B, Section B-5a(1)).

12 B3-11 Reconciliation with Data Quality Objectives

13 Reconciling the results of waste testing and analysis with the DQOs provides a way to ensure
14 that data will be of adequate quality to support the regulatory compliance programs.
15 Reconciliation with the DQOs will take place at both the project level and the Permittees' level.
16 At the project level, reconciliation will be performed by the Site Project Manager, while at the
17 Permittees' level, reconciliation will be performed as described below.

18 B3-11a Reconciliation at the Project Level

19 The Permittees shall require each Site Project Manager to ensure that all data generated and used
20 in decision making meet the DQOs provided in Section B-4a(1) of Permit Attachment B. To do
21 so, the Site Project Manager must assess whether data of sufficient type, quality, and quantity
22 have been collected. The Site Project Manager must determine if the variability of the data set is
23 small enough to provide the required confidence in the results. The Site Project Manager must
24 also determine if, based on the desired error rates and confidence levels, a sufficient number of
25 valid data points have been determined (as established by the associated completeness rate for
26 each sampling and analytical process). In addition, the Site Project Manager must document that
27 random sampling of containers was performed for the purposes of waste stream characterization.

28 For each waste stream characterized, the Permittees shall require each Site Project Manager to
29 determine if sufficient data have been collected to determine the following WAP-required waste
30 parameters, as applicable:

- 31 • Waste matrix code
- 32 • Waste material parameter weights
- 33 • If each waste container of waste contains TRU radioactive waste

- 1 • Mean concentrations, UCL₉₀ for the mean concentrations, standard deviations, and the
2 number of samples collected for each VOC in the headspace gas of waste containers in
3 the waste stream
- 4 • Mean concentrations, UCL₉₀ for the mean concentrations, standard deviations, and
5 number of samples collected for VOCs, SVOCs, and metals in the waste stream
- 6 • Whether the waste stream exhibits a toxicity characteristic (TC) under 40 CFR Part 261,
7 Subpart C
- 8 • Whether the waste stream contains listed waste found in 20.4.1.200 NMAC incorporating
9 40 CFR Part 261, Subpart D
- 10 • Whether the waste stream can be classified as hazardous or nonhazardous at the 90-
11 percent confidence level
- 12 • Whether an appropriate packaging configuration and DAC were applied and documented
13 in the headspace gas sampling documentation, and whether the drum age was met prior to
14 sampling.
- 15 • Whether all TICs were appropriately identified and reported in accordance with the
16 requirements of Section B3-1 prior to submittal of a WSPF for a waste stream or waste
17 stream lot.
- 18 • Whether the overall completeness, comparability, and representativeness QAOs were met
19 for each of the analytical and testing procedures as specified in Sections B3-2 through
20 B3-9 prior to submittal of a WSPF for a waste stream or waste stream lot.
- 21 • Whether the PRQLs for all analyses were met prior to submittal of a WSPF for a waste
22 stream or waste stream lot.

23 If the Site Project Manager determines that insufficient data have been collected to make the
24 determinations listed above, additional data collection efforts must be undertaken. The
25 reconciliation of a waste stream shall be performed, as described in Permit Attachment B4, prior
26 to submittal of WSPF and Characterization Information Summary to the Permittees for that
27 waste stream. The Permittees shall not manage, store, or dispose a TRU mixed waste stream at
28 WIPP unless the Site Project Manager determines that the WAP-required waste parameters listed
29 above have been met for that waste stream.

30 The statistical procedure presented in Permit Attachment B2 shall be used by participating Site
31 Project Managers to evaluate and report waste characterization data from the analysis of
32 homogeneous solids and soil/gravel. The procedure, which calculates UCL₉₀ values, shall be
33 used to assess compliance with the DQOs in Attachment B, Section B-4a(1) as well as with
34 RCRA regulations. The procedure must be applied to all laboratory analytical data for total
35 VOCs, total SVOCs, and total metals. For RCRA regulatory compliance (40 CFR § 261.24), data

1 from the analysis of the appropriate metals and organic compounds shall be expressed as toxicity
2 characteristic leaching procedure (**TCLP**) values or results may also be compared to the TC
3 levels expressed as total values. These total values will be considered the regulatory threshold
4 limit (**RTL**) values for the WAP. RTL values are obtained by calculating the weight/weight
5 concentration (in the solid) of a TC analyte that would give the regulatory weight/volume
6 concentration (in the TCLP extract), assuming 100-percent analyte dissolution.

7 B3-11b Reconciliation at the Permittee Level

8 The Permittees must also ensure that data of sufficient type, quality, and quantity are collected to
9 meet WAP DQOs. The Permittees will ensure sufficient data have been collected to determine if
10 the waste characterization information is adequate to demonstrate the Permittees' compliance
11 with Attachment B, Section B-4a(1). This is performed during Permittees' review of the WSPF
12 and Characterization Information Summary.

13 B3-12 Data Reporting Requirements

14 Data reporting requirements define the type of information and the method of transmittal for data
15 transfer from the data generation level to the project level and from the project level to the
16 Permittees.

17 B3-12a Data Generation Level

18 Data shall be transmitted by hard copy or electronically (provided a hard copy is available on
19 demand) from the data generation level to the project level. Transmitted data shall include all
20 Batch Data Reports and data review checklists. The Batch Data Reports and checklists used must
21 contain all of the information required by the testing, sampling, and analytical techniques
22 described in Permit Attachments B1 through B6 , as well as the signature releases to document
23 the review, validation, and verification as described in Section B3-10. All Batch Data Reports
24 and checklists shall be in approved formats, as provided in site-specific documentation.

25 Batch Data Reports shall be forwarded to the Site Project Manager. All Batch Data Reports shall
26 be assigned serial numbers, and each page shall be numbered. The serial number used for Batch
27 Data Reports can be the same as the testing, sampling, or analytical batch number.

28 QA documentation, including raw data, shall be maintained in either testing, sampling, and
29 analytical facility files, or site project files for those facilities located on site in accordance with
30 the document storage requirements of site approved site QAPjPs. Permittee approved
31 laboratories shall forward testing, sampling, and analytical QA documentation along with Batch
32 Data Reports to the site project office for inclusion in site project files.

33 B3-12b Project Level

34 The site project office shall prepare a WSPF for each waste stream certified for shipment to
35 WIPP based on information obtained from acceptable knowledge and Batch Data Reports, if

1 applicable. In addition, the site project office must ensure that the Characterization Information
2 Summary and the Waste Stream Characterization Package (when requested by the Permittees)
3 are prepared as appropriate. The Site Project Manager must also verify these reports are
4 consistent with information found in analytical batch reports. Summarized testing, sampling, and
5 analytical data are included in the Characterization Information Summary. The contents of the
6 WSPF, Characterization Information Summary, and Waste Stream Characterization Package are
7 discussed in the following sections.

8 After approval of a WSPF and the associated Characterization Information Summary by the
9 Permittees, the generator/storage site are required to maintain a cross reference of container
10 identification numbers to each Batch Data Report.

11 A Waste Stream Characterization Package shall be transmitted by hard copy or electronically
12 from the Site Project Manager to the Permittees when requested.

13 B3-12b(1) Waste Stream Profile Form

14 The Waste Stream Profile Form (WSPF, Figure B-1) shall include the following information:

- 15 • Generator/storage site name
- 16 • Generator/storage site EPA ID
- 17 • Date of audit report approval by NMED (if obtained)
- 18 • Original generator of waste stream
- 19 • Whether waste is Contact-Handled or Remote-Handled
- 20 • The Waste Stream WIPP Identification Number
- 21 • Summary Category Group
- 22 • Waste Matrix Code Group
- 23 • Waste Material Parameter Weight Estimates per unit of waste
- 24 • Waste stream name
- 25 • A description of the waste stream
- 26 • Applicable EPA hazardous waste numbers
- 27 • Applicable TRUCON codes
- 28 • A listing of acceptable knowledge documentation used to identify the waste stream

- 1 • The waste characterization procedures used and the reference and date of the procedure
- 2 • Certification signature of Site Project Manager, name, title, and date signed

3 B3-12b(2) Characterization Information Summary

4 The Characterization Information Summary shall include the following elements, if applicable:

- 5 • Data reconciliation with DQOs
- 6 • Headspace gas summary data listing the identification numbers of samples used in the
7 statistical reduction, the maximum, mean, standard deviation, UCL₉₀, RTL, and
8 associated EPA hazardous waste numbers that must be applied to the waste stream.
- 9 • Total metal, VOC, and SVOC analytical results for homogeneous solids and soil/gravel
10 (if applicable).
- 11 • TIC listing and evaluation.
- 12 • Radiography and visual examination summary to document that all prohibited items are
13 absent in the waste (if applicable).
- 14 • A complete listing of all container identification numbers used to generate the WSPF,
15 cross-referenced to each Batch Data Report
- 16 • Complete AK summary, including stream name and number, point of generation, waste
17 stream volume (current and projected), generation dates, TRUCON codes, Summary
18 Category Group, Waste Matrix Code(s) and Waste Matrix Code Group, other TWBIR
19 information, waste stream description, areas of operation, generating processes, RCRA
20 determinations, radionuclide information, all references used to generate the AK
21 summary, and any other information required by Permit Attachment B4, Section B4-2b.
- 22 • Method for determining Waste Material Parameter Weights per unit of waste.
- 23 • List of any AK Sufficiency Determinations requested for the waste stream.
- 24 • Certification through acceptable knowledge or testing and/or analysis that any waste
25 assigned the hazardous waste number of U134 (hydrofluoric acid) no longer exhibits the
26 characteristic of corrosivity. This is verified by ensuring that no liquid is present in U134
27 waste.

1 B3-12b(3) Waste Stream Characterization Package

2 The Waste Stream Characterization Package includes the following information:

- 3 • Waste Stream Profile Form (WSPF, Section B3-12b(1))
- 4 • Accompanying Characterization Information Summary (Section B3-12b(2))
- 5 • Complete AK summary (Section B3-12b(2))
- 6 • Batch Data Reports supporting the characterization of the waste stream and any others
7 requested by the Permittees
- 8 • Raw analytical data requested by the Permittees

9 B3-12b(4) WIPP Waste Information System (WWIS) Data Reporting

10 The WWIS Data Dictionary includes all of the data fields, the field format and the limits
11 associated with the data as established by this WAP. These data will be subjected to edit and
12 limit checks that are performed automatically by the database, as defined in the *WIPP Waste
13 Information System User's Manual for Use by Shippers/Generators* (DOE, 2001). If a container
14 was part of a composite headspace gas sample, the analytical results from the composite sample
15 must be assigned as the container headspace gas data results, including associated TICs, for
16 every waste container associated with the composite sample.

17 B3-13 Nonconformances

18 The Permittees shall require the status of work and the WAP activities at participating
19 generator/storage sites to be monitored and controlled by the Site Project Manager. This
20 monitoring and control shall include nonconformance identification, documentation, and
21 reporting.

22 The nonconformances and corrective action processes specified in this section describe
23 procedures between the Permittees and the generator/storage sites.

24 Nonconformances

25 Nonconformances are uncontrolled and unapproved deviations from an approved plan or
26 procedure. Nonconforming items and activities are those that do not meet the WAP
27 requirements, procurement document criteria, or approved work procedures. Nonconforming
28 items shall be identified by marking, tagging, or segregating, and the affected generator/storage
29 site(s) notified. The Permittees shall require participating sites reconcile and correct
30 nonconforming items as appropriate in accordance with the Permittees' Quality Assurance
31 Program Description (**QAPD**). Disposition of nonconforming items shall be identified and

1 documented. The QAPjPs shall identify the person(s) responsible for evaluating and
2 dispositioning nonconforming items and shall include referenced procedures for handling them.

3 Management at all levels shall foster a “no-fault” attitude to encourage the identification of
4 nonconforming items and processes. Nonconformances may be detected and identified by
5 anyone performing WAP activities, including

- 6 • Project staff - during field operations, supervision of subcontractors, data validation and
7 verification, and self-assessment
- 8 • Laboratory staff - during the preparation for and performance of laboratory testing;
9 calibration of equipment; QC activities; laboratory data review, validation, and
10 verification; and self-assessment
- 11 • QA personnel - during oversight activities or audits

12 A nonconformance report shall be prepared for each nonconformance identified. Each
13 nonconformance report shall be initiated by the individual(s) identifying the nonconformance.
14 The nonconformance report shall then be processed by knowledgeable and appropriate
15 personnel. For this purpose, a nonconformance report including, or referencing as appropriate,
16 results of laboratory analysis, QC tests, audit reports, internal memoranda, or letters shall be
17 prepared. The nonconformance report must provide the following information:

- 18 • Identification of the individual(s) identifying or originating the nonconformance
- 19 • Description of the nonconformance
- 20 • Method(s) or suggestions for correcting the nonconformance (corrective action)
- 21 • Schedule for completing the corrective action
- 22 • An indication of the potential ramifications and overall useability the data, if applicable
- 23 • Any approval signatures specified in the site nonconformance procedures

24 The Permittees shall require the Site Project Manager to oversee the nonconformance report
25 process and be responsible for developing a plan to identify and track all nonconformances and
26 report this information to the Permittees. The Site Project Manager is also responsible for
27 notifying project personnel of the nonconformance and verifying completion of the corrective
28 action for nonconformances.

29 Nonconformance to DQOs

30 For any non-administrative nonconformance related to applicable requirements specified in this
31 WAP which are first identified at the Site Project Manager signature release level (i.e., a failure

1 to meet a data quality objective DQO), the Permittees shall receive written notification within
2 five (5) calendar days of identification and shall also receive a nonconformance report within
3 thirty (30) calendar days of identification of the incident. The Permittees shall require the
4 generator/storage site to implement a corrective action which remedies the nonconformance prior
5 to management, storage, or disposal of the waste at WIPP. The Permittees shall send NMED a
6 monthly summary of nonconformances identified during the previous month, indicating the
7 number of nonconformances received and the generator/storage sites responsible.

8 Permittees' Corrective Action Process

9 The Permittees shall initiate a corrective action process when internal nonconformances and
10 nonconformances at the generator/storage sites are identified. Activities and processes that do
11 not meet requirements are documented as deficiencies.

12 When a deficiency is identified by the Permittees, the following process action steps are
13 required:

- 14 • The condition is documented on a Corrective Action Report (**CAR**) by the individual
15 identifying the problem.
- 16 • The Permittees have designated the CAR Initiator and Assessment Team Leader to
17 review the CAR, determine validity of the finding (determine that a requirement has been
18 violated), classify the significance of the condition, assign a response due date, and issue
19 the CAR to the responsible party.
- 20 • The responsible organization reviews the CAR, evaluates the extent and cause of the
21 deficiency and provides a response to the Permittees, indicating remedial actions and
22 actions to preclude recurrence that will be taken.
- 23 • The Permittees review the response from the responsible organization and, if acceptable,
24 communicate the acceptance to the responsible organization.
- 25 • The responsible organization completes remedial actions and actions to preclude
26 recurrence of the condition.
- 27 • After all corrective actions have been completed, the Permittees schedule and perform a
28 verification to ensure that corrective actions have been completed and are effective.
29 When all actions have been completed and verified as being effective, the CAR is closed
30 by the CAR Initiator and Assessment Team Leader on behalf of the Permittees.
- 31 • As part of the planning process for subsequent audits and surveillances, past deficiencies
32 are reviewed and the previous deficient activity or process is subject to reassessment.

1 B3-14 Special Training Requirements and Certifications

2 Before performing activities that affect WAP quality, all personnel are required to receive
3 indoctrination into the applicable scope, purpose, and objectives of the WAP and the specific
4 QAOs of the assigned task. Personnel assigned to perform activities for the WAP shall have the
5 education, experience, and training applicable to the functions associated with the work.
6 Evidence of personnel proficiency and demonstration of competence in the task(s) assigned must
7 be demonstrated and documented. All personnel designated to work on specific aspects of the
8 WAP shall maintain qualification (i.e., training and certification) throughout the duration of the
9 work as specified in this WAP and applicable QAPjPs/procedures. Job performance shall be
10 evaluated and documented at periodic intervals, as specified in the implementing procedures.

11 Personnel involved in WAP activities shall receive continuing training to ensure that job
12 proficiency is maintained. Training includes both education in principles and enhancement of
13 skills. Each participating site shall include in its QAPjP a description of the procedures for
14 implementing personnel qualification and training. All training records that specify the scope of
15 the training, the date of completion, and documentation of job proficiency shall be maintained as
16 QA Records in the site project file.

17 Analytical laboratory line management must ensure that analytical personnel are qualified to
18 perform the analytical method(s) for which they are responsible. The minimum qualifications for
19 certain specified positions for the WAP are summarized in Table B3-10. QAPjPs, or their
20 implementing SOPs, shall specify the site-specific titles and minimum training and qualification
21 requirements for personnel performing WAP activities. QAPjPs/procedures shall also contain the
22 requirements for maintaining records of the qualification, training, and demonstrations of
23 proficiency by these personnel.

24 An evaluation of personnel qualifications shall include comparing and evaluating the
25 requirements specified in the job/position description and the skills, training, and experience
26 included in the current resume of the person. This evaluation also must be performed for
27 personnel who change positions because of a transfer or promotion as well as personnel assigned
28 to short-term or temporary work assignments that may affect the quality of the WAP.
29 QAPjPs/procedures shall identify the responsible person(s) for ensuring that all personnel
30 maintain proficiency in the work performed and identify any additional training that may be
31 required.

32 B3-15 Changes to WAP-Related Plans or Procedures

33 Controlled changes to WAP-related plans or procedures shall be managed through the document
34 control process described in the QAPD. The Site Project Manager shall review all non-
35 administrative changes and evaluate whether those changes could impact DQOs specified in the
36 Permit. After site certification, any changes to WAP-related plans or procedures that could
37 positively or negatively impact DQOs (i.e., those changes that require prior approval of the
38 Permittees as defined in Attachment B5, Section B5-2) shall be reported to the Permittees within
39 five (5) days of identification by the project level review. The Permittees shall send NMED a

1 monthly summary briefly describing the changes to plans and procedures identified pursuant to
2 this section during the previous month.

3 B3-16 List of References

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11 U.S. Department of Energy.

12 DOE. 2005. Performance Demonstration Program Plan for RCRA Constituent Analysis of
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14 Area Office, U.S. Department of Energy.

15 EPA. 1996. *Test Methods for Evaluating Solid Waste, Physical/Chemical Methods*. SW-846,
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17 Environmental Protection Agency.

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19 Multi-Component Alpha Emitting Samples." *Radiochem. Radioanal. Letters*, 16, No. 1:
20 pp. 5-16.

21 Pasternack B. S. and N. H. Harley. 1971. "Detection Limits for Radionuclides in the Analysis of
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TABLES

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**TABLE B3-1
 WASTE MATERIAL PARAMETERS AND DESCRIPTIONS**

Waste Material Parameter	Description
Iron-based Metals/Alloys	Iron and steel alloys in the waste; does not include the waste container materials
Aluminum-based Metals/Alloys	Aluminum or aluminum-based alloys in the waste materials
Other Metals	All other metals found in the waste materials
Other Inorganic Materials	Nonmetallic inorganic waste including concrete, glass, firebrick, ceramics, sand, and inorganic sorbents
Cellulosics	Materials generally derived from high-polymer plant carbohydrates; (e.g., paper, cardboard, wood, and cloth)
Rubber	Natural or man-made elastic latex materials; (e.g., surgeons' gloves, and leaded rubber gloves)
Plastics (waste materials)	Generally man-made materials, often derived from petroleum feedstock; (e.g., polyethylene and polyvinylchloride)
Organic Matrix	Cemented organic resins, solidified organic liquids and sludges
Inorganic Matrix	Any homogeneous materials consisting of sludge or aqueous-based liquids that are solidified with cement, calcium silicate, or other solidification agents; (e.g., wastewater treatment sludge, cemented aqueous liquids, and inorganic particulates)
Soils/gravel	Generally consists of naturally occurring soils that have been contaminated with inorganic waste materials
Steel (packaging materials)	55-gal (208-L) drums
Plastics (packaging materials)	90-mil polyethylene drum liner and plastic bags

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TABLE B3-2
**GAS VOLATILE ORGANIC COMPOUNDS TARGET ANALYTE LIST
AND QUALITY ASSURANCE OBJECTIVES**

Compound	CAS Number	Precision ^a (%RSD or RPD)	Accuracy ^a (%R)	MDL ^{b,d} (ng)	FTIRS MDL ^b (ppmv)	PRQL (ppmv)	Completeness (%)
Benzene	71-43-2	≤25	70-130	10	5	10	90
Bromoform	75-25-2	≤25	70-130	10	5	10	90
Carbon tetrachloride	56-23-5	≤25	70-130	10	5	10	90
Chlorobenzene	108-90-7	≤25	70-130	10	5	10	90
Chloroform	67-66-3	≤25	70-130	10	5	10	90
1,1-Dichloroethane	75-34-3	≤25	70-130	10	5	10	90
1,2-Dichloroethane	107-06-2	≤25	70-130	10	5	10	90
1,1-Dichloroethylene	75-35-4	≤25	70-130	10	5	10	90
cis-1,2-Dichloroethylene	156-59-2	≤25	70-130	10	5	10	90
trans-1,2-Dichloroethylene	156-60-5	≤25	70-130	10	5	10	90
Ethyl benzene ^d	100-41-4	≤25	70-130	10	10	10	90
Ethyl ether	60-29-7	≤25	70-130	10	5	10	90
Methylene chloride	75-09-2	≤25	70-130	10	5	10	90
1,1,2,2-Tetrachloroethane	79-34-5	≤25	70-130	10	5	10	90
Tetrachloroethylene	127-18-4	≤25	70-130	10	5	10	90
Toluene	108-88-3	≤25	70-130	10	5	10	90
1,1,1-Trichloroethane	71-55-6	≤25	70-130	10	5	10	90
Trichloroethylene	79-01-6	≤25	70-130	10	5	10	90
1,1,2-Trichloro-1,2,2-trifluoroethane	76-13-1	≤25	70-130	10	5	10	90
m-Xylene ^c	108-38-3	≤25	70-130	10	5	10	90
o-Xylene	95-47-6	≤25	70-130	10	5	10	90
p-Xylene ^c	106-42-3	≤25	70-130	10	5	10	90
Acetone	67-64-1	≤25	70-130	150	50	100	90
Butanol	71-36-3	≤25	70-130	150	50	100	90
Methanol	67-56-1	≤25	70-130	150	50	100	90
Methyl ethyl ketone	78-93-3	≤25	70-130	150	50	100	90
Methyl isobutyl ketone	108-10-1	≤25	70-130	150	50	100	90

^a Criteria apply to PRQL concentrations.

^b Values based on delivering 10 mL to the analytical system.

^c These xylene isomers cannot be resolved by GC/MS.

^d The ethyl benzene PRQL for FTIRS is 20 ppm

CAS = Chemical Abstract Service

%RSD = Percent relative standard deviation

RPD = Relative percent difference

%R = Percent recovery

MDL = Method detection limit (maximum permissible value), for GC/MS and GC/FID; total number of nanograms delivered to the analytical system per sample (nanograms); for FTIRS based on 1 m sample cell

PRQL = Program required quantitation limit (parts per million/volume basis)

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**TABLE B3-3
 SUMMARY OF LABORATORY QUALITY CONTROL SAMPLES AND
 FREQUENCIES FOR
 GAS VOLATILE ORGANIC COMPOUND ANALYSIS**

QC Sample	Minimum Frequency	Acceptance Criteria	Corrective Action ^a
Method performance samples	Seven (7) samples initially and four (4) semiannually	Meet method QAOs	Repeat until acceptable
Laboratory duplicates or on-line duplicates	One (1) per analytical batch or on-line batch	RPD $\leq 25^b$	Nonconformance if RPD >25
Laboratory blanks or on-line blanks	Daily prior to sample analysis for GC/MS and GC/FID. Otherwise, daily prior to sample analysis and one (1) per analytical batch or on-line	Analyte amounts $\leq 3 \times$ MDLs for GC/MS and GC/FID; \leq PRQL for FTIRS	Flag Data if analyte amounts $> 3 \times$ MDLs for GC/MS and GC/FID; $>$ PRQL for FTIRS
Laboratory control samples or on-line control samples	One (1) per analytical batch or on-line batch	70-130 %R	Nonconformance if %R <70 or >130
GC/MS comparison sample (for FTIRS only)	One (1) per analytical or on-line batch	RPD $\leq 25^b$	Nonconformance if RPD > 25
Blind audit samples	Samples and frequency controlled by the Gas PDP Plan	Specified in the Gas PDP Plan	Specified in the Gas PDP Plan
GC/MS	BFB Tune Every 12 hours	Abundance criteria for key ions are met	Repeat Until Acceptable
GC/MS	Minimum 5-point initial calibration (minimum of 5 standards) Initially and as needed	%RSD of response factor for each target analyte <35	Repeat Until Acceptable
GC/MS	Continuing calibration Every 12 hours	%D for all target analytes ≤ 30 of initial calibration	Repeat Until Acceptable

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QC Sample	Minimum Frequency	Acceptance Criteria	Corrective Action ^a
GC/FID	Minimum 3-point initial calibration (minimum 3 standards) Initially and as needed	Correlation coefficient \geq 0.99 or %RSD <20 for each target analyte and the retention time of each target analyte within an acceptance criteria defined in the method	Repeat Until Acceptable
GC/FID	Continuing calibration Every 12 hours	%RSD \leq 15%	Repeat Until Acceptable

- 1 ^a Corrective action per Section B3-13 when final reported QC samples do not meet the acceptance criteria.
 2 ^b Applies only to concentrations greater than the PRQLs listed in Table B3-2.
- 3 MDL = Method Detection Limit
 4 QAO = Quality Assurance Objective
 5 PDP = Performance Demonstration Program
 6 PRQL = Program Required Quantitation Limit
 7 %R = Percent Recovery
 8 RPD = Relative Percent Difference
 9 BFB = 4-Bromofluorobenzene
 10 %D = Percent difference
 11 %RSD = Percent relative standard deviation

**TABLE B3-4
 VOLATILE ORGANIC COMPOUNDS TARGET ANALYTE LIST
 AND QUALITY ASSURANCE OBJECTIVES**

Compound	CAS Number	Precision ^a (%RSD or RPD)	Accuracy ^a (%R)	MDL ^b (mg/kg)	PRQL ^b (mg/kg)	Completeness (%)
Benzene	71-43-2	≤45	37-151	1	10	90
Bromoform	75-25-2	≤47	45-169	1	10	90
Carbon disulfide	75-15-0	≤50	60-150	1	10	90
Carbon tetrachloride	56-23-5	≤30	70-140	1	10	90
Chlorobenzene	108-90-7	≤38	37-160	1	10	90
Chloroform	67-66-3	≤44	51-138	1	10	90
1,4-Dichlorobenzene ^c	106-46-7	≤60	18-190	1	10	90
ortho-Dichlorobenzene ^c	95-50-1	≤60	18-190	1	10	90
1,2-Dichloroethane	107-06-2	≤42	49-155	1	10	90
1,1-Dichloroethylene	75-35-4	≤250	D-234 ^d	1	10	90
trans-1,2-Dichloroethylene	156-60-5	≤50	60-150	1	10	90
Ethyl benzene	100-41-4	≤43	37-162	1	10	90
Methylene chloride	75-09-2	≤50	D-221 ^d	1	10	90
1,1,2,2-Tetrachloroethane	79-34-5	≤55	46-157	1	10	90
Tetrachloroethylene	127-18-4	≤29	64-148	1	10	90
Toluene	108-88-3	≤29	47-150	1	10	90
1,1,1-Trichloroethane	71-55-6	≤33	52-162	1	10	90
1,1,2-Trichloroethane	79-00-5	≤38	52-150	1	10	90
Trichloroethylene	79-01-6	≤36	71-157	1	10	90
Trichlorofluoromethane	75-69-4	≤110	17-181	1	10	90
1,1,2-Trichloro-1,2,2-trifluoroethane	76-13-1	≤50	60-150	1	10	90
Vinyl chloride	75-01-4	≤200	D-251 ^d	1	4	90
m-xylene	108-38-3	≤50	60-150	1	10	90
o-xylene	95-47-6	≤50	60-150	1	10	90
p-xylene	106-42-3	≤50	60-150	1	10	90
Acetone	67-64-1	≤50	60-150	10 ^e	100	90
Butanol	71-36-3	≤50	60-150	10 ^e	100	90
Ethyl ether	60-29-7	≤50	60-150	10 ^e	100	90
Formaldehyde ^f	50-00-0	≤50	60-150	10 ^e	100	90
Hydrazine ^g	302-01-2	≤50	60-150	10 ^e	100	90
Isobutanol	78-83-1	≤50	60-150	10 ^e	100	90
Methanol	67-56-1	≤50	60-150	10 ^e	100	90
Methyl ethyl ketone	78-93-3	≤50	60-150	10 ^e	100	90
Pyridine ^c	110-86-1	≤50	60-150	10 ^e	100	90

^a Applies to laboratory control samples and laboratory matrix spikes. If a solid laboratory control sample material which has established statistical control limits is used, then the established control limits for that material should be used for accuracy requirements.

^b TCLP MDL and PRQL values are reported in units of mg/l and limits are reduced by a factor of 20.

^c Can also be analyzed as a semi-volatile organic compound. If analyzed as a semi-volatile compound, the QAOs of Table B3-6 apply.

^d Detected; result must be greater than zero.

^e Estimate, to be determined.

^f Required only for homogeneous solids and soil/gravel waste from Savannah River Site, if analysis is required to resolve assignment of EPA hazardous waste numbers.

^g Required only for homogeneous solids and soil/gravel waste from Oak Ridge National Laboratory and Savannah River Site, if analysis is required to resolve assignment of EPA hazardous waste numbers.

CAS = Chemical Abstract Service

%RSD = Percent relative standard deviation

RPD = Relative percent difference

%R = Percent recovery

MDL = Method detection limit (maximum permissible value) (milligrams per kilogram)

PRQL = Program required quantitation limit; calculated from the toxicity characteristic level for benzene assuming a 0.9 oz (25-gram [g]) sample, 0.1 gal (0.5 liter [L]) of extraction fluid, and 100 percent analyte extraction (milligrams per kilogram)

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**TABLE B3-5
 SUMMARY OF LABORATORY QUALITY CONTROL SAMPLES AND
 FREQUENCIES FOR VOLATILE ORGANIC COMPOUND ANALYSIS**

QC Sample	Minimum Frequency	Acceptance Criteria	Corrective Action^a
Method performance samples	Seven (7) samples initially and four (4) semiannually	Meet Table B3-4 QAOs	Repeat until acceptable
Laboratory duplicates ^b	One (1) per analytical batch	Meet Table B3-4 precision QAOs	Nonconformance if RPDs > values in Table B3-4
Laboratory blanks	One (1) per analytical batch	Analyte concentrations ≤ 3 × MDLs	Nonconformance if analyte concentrations > 3 × MDLs
Matrix spikes ^b	One (1) per analytical batch	Meet Table B3-4 accuracy QAOs	Nonconformance if %Rs are outside the range specified in Table B3-4
Matrix spike duplicates	One (1) per analytical batch	Meet Table B3-4 accuracy and precision QAOs	Nonconformance if RPDs > values and %Rs outside range specified in Table B3-4
Laboratory control samples	One (1) per analytical batch	Meet Table B3-4 accuracy QAO's	Nonconformance if %R < 80 or > 120
GC/MS Calibration	BFB Tune every 12 hours 5-pt. Initial Calibration initially, and as needed	Abundance criteria met as per method Calibrate according to SW-846 Method requirements: %RSD for CCC ≤ 30, %RSD for all other compounds ≤ 15% Average response factor (RRF) used if %RSD ≤ 15, use linear regression if %RSD > 15; R or R ² ≥ 0.990 if using alternative curve System Performance Check Compound (SPCC) minimum RRF as per SW-846 Method; RRF for all other compounds ≥ 0.01	Repeat until acceptable

QC Sample	Minimum Frequency	Acceptance Criteria	Corrective Action ^a
GC/MS Calibration (continued)	Continuing Calibration every 12 hours	%D ≤ 20 for CCC; SPCC minimum RRF as per SW-846 Method; RRF for all other compounds ≥ 0.01 RT for internal standard must be ± 30 seconds from last daily calibration, internal standard area count must be >50% and <200% of last daily calibration	Repeat until acceptable
GC/FID Calibration	3-pt. Initial Calibration initially and as needed Continuing Calibration every 12 hours	Correlation Coefficient ≥ 0.990 or %RSD ≤ 20 for all analytes %D or %Drift for all analytes ≤ 15 of expected values, RT ± 3 standard deviations from initial RT calibration per applicable SW-846 Method	Repeat until acceptable.
Surrogate compounds	Each analytical sample	Average %R from minimum of 30 samples for a given matrix ±3 standard deviations	Nonconformance if %R < (average %R - 3 standard deviation) or > (average %R + 3 standard deviation)
Blind audit samples	Samples and frequency controlled by the Solid PDP Plan	Specified in the Solid PDP Plan	Specified in the Solid PDP Plan

- 1 ^a Corrective Action per Section B3-13 when final reported QC samples do not meet the acceptance criteria.
 2 Nonconformances do not apply to matrix related exceedances.
 3 ^b May be satisfied using matrix spike duplicate; acceptance criteria applies only to concentrations greater than the
 4 PRQLs listed in Table B3-4.
- 5 MDL = Method detection limit
 6 QAO = Quality assurance objective
 7 PDP = Performance Demonstration Program
 8 %R = Percent recovery
 9 RPD = Relative percent difference

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TABLE B3-6
SEMI-VOLATILE ORGANIC COMPOUND TARGET ANALYTE LIST
AND QUALITY ASSURANCE OBJECTIVES

Compound	CAS Number	Precision ^a (%RSD or RPD)	Accuracy ^a (%R)	MDL ^b (mg/kg)	PRQL ^b (mg/kg)	Completeness (%)
Cresols	1319-77-3	≤50	25-115	5	40	90
1,4-Dichlorobenzene ^{bc}	106-46-7	≤86	20-124	5	40	90
ortho-Dichlorobenzene ^c	95-50-1	≤64	32-129	5	40	90
2,4-Dinitrophenol	51-28-5	≤119	D-172 ^d	5	40	90
2,4-Dinitrotoluene	121-14-2	≤46	39-139	0.3	2.6	90
Hexachlorobenzene	118-74-1	≤319	D-152 ^d	0.3	2.6	90
Hexachloroethane	67-72-1	≤44	40-113	5	40	90
Nitrobenzene	98-95-3	≤72	35-180	5	40	90
Pentachlorophenol	87-86-5	≤128	14-176	5	40	90
Pyridine ^c	110-86-1	≤50	25-115	5	40	90

- 4 CAS = Chemical Abstract Service
 5 %RSD = Percent relative standard deviation
 6 RPD = Relative percent difference
 7 %R = Percent recovery
 8 MDL = Method detection limit (maximum permissible value) (milligrams per kilogram)
 9 PRQL = Program required quantitation limit; calculated from the toxicity characteristic level for nitrobenzene
 10 assuming a 100-gram (g) sample, 0.5 gal (2 liter [L]) of extraction fluid, and 100 percent analyte
 11 extraction (milligrams per kilograms)
- 12 ^a Applies to laboratory control samples and laboratory matrix spikes. If a solid laboratory control sample material
 13 which has established statistical control limits is used, then the established control limits for that material should be
 14 used for accuracy requirements.
 15 ^b TCLP MDL and PRQL values are reported in units of mg/l and limits are reduced by a factor of 20.
 16 ^c Can also be analyzed as a volatile organic compound
 17 ^d Detected; result must be greater than zero

QC Sample	Minimum Frequency	Acceptance Criteria	Corrective Action ^a
GC/ECD Calibration	5-pt. Calibration initially and as needed Continuing Calibration every 12 hours	Correlation Coefficient \geq 0.990 or %RSD < 20 for all analytes %D or %Drift for all analytes \leq 15 of expected values, RT \pm 3 standard deviations of initial RT calibration per applicable SW-846 Method	Repeat until acceptable
Matrix spike duplicates	One (1) per analytical batch	Meet Table B3-6 accuracy and precision QAOs	Nonconformance if RPDs > values and %Rs outside range specified in Table B3-6
Laboratory control samples	One (1) per analytical batch	Meet Table B3-6 accuracy QAO's	Nonconformance if %R < 80 or > 120
Surrogate compounds	Each analytical sample	Average %R from minimum of 30 samples from a given matrix \pm 3 standard deviations	Nonconformance if %R < (average %R - 3 standard deviations) or > (average %R + 3 standard deviations)
Blind audit samples	Samples and frequency controlled by the Solid PDP Plan	Specified in the Solid PDP Plan	Specified in the Solid PDP Plan

- 1 ^a Corrective action per Section B3-13 when final reported QC samples do not meet the acceptance criteria.
2 Nonconformances do not apply to matrix related exceedances.
3 ^b May be satisfied by using matrix spike duplicate; acceptance criteria applies only to concentrations greater than the
4 PRQLs listed in Table B3-6.
- 5 MDL = Method Detection Limit
6 QAO = Quality Assurance Objective
7 PDP = Performance Demonstration Program
8 %R = Percent Recovery
9 RPD = Relative Percent Difference

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**TABLE B3-8
 METALS TARGET ANALYTE LIST
 AND QUALITY ASSURANCE OBJECTIVES**

Analyte	CAS Number	Precision (%RSD or RPD) ^a	Accuracy (%R) ^b	PRDL ^d (µg/L)	PRQL ^c (mg/kg)	Completeness (%)
Antimony	7440-36-0	≤30	80-120	100	100	90
Arsenic	7440-38-2	≤30	80-120	100	100	90
Barium	7440-39-3	≤30	80-120	2000	2000	90
Beryllium	7440-41-7	≤30	80-120	100	100	90
Cadmium	7440-43-9	≤30	80-120	20	20	90
Chromium	7440-47-3	≤30	80-120	100	100	90
Lead	7439-92-1	≤30	80-120	100	100	90
Mercury	7439-97-6	≤30	80-120	4.0	4.0	90
Nickel	7440-02-0	≤30	80-120	100	100	90
Selenium	7782-49-2	≤30	80-120	20	20	90
Silver	7440-22-4	≤30	80-120	100	100	90
Thallium	7440-28-0	≤30	80-120	100	100	90
Vanadium	7440-62-2	≤30	80-120	100	100	90
Zinc	7440-66-6	≤30	80-120	100	100	90

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^a ≤ 30 percent control limits apply when sample and duplicate concentrations are ≥ 10 x IDL for ICP-AES and AA techniques, and ≥ 100 x IDL for Inductively Coupled Plasma—Mass Spectrometry (ICP-MS) techniques. If less than these limits, the absolute difference between the two values shall be less than or equal to the PRQL.

^b Applies to laboratory control samples and laboratory matrix spikes. If a solid laboratory control sample material which has established statistical control limits is used, then the established control limits for that material should be used for accuracy requirements.

^c TCLP PRQL values are reported in units of mg/l and limits are reduced by a factor of 20.

^d PRDL set such that it is a factor of 10 below the PRQL for 100 percent solid samples, assuming a 100x dilution during digestion.

CAS = Chemical Abstract Service
 %RSD = Percent relative standard deviation
 RPD = Relative percent difference
 %R = Percent recovery
 PRDL = Program required detection limit (i.e., maximum permissible value for IDL) (micrograms per liter)
 PRQL = Program required quantitation limit (milligrams per kilogram)

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**TABLE B3-9
SUMMARY OF LABORATORY QUALITY CONTROL SAMPLES AND
FREQUENCIES FOR METALS ANALYSIS**

QC Sample	Minimum Frequency	Acceptance Criteria	Corrective Action^a
Method performance samples	Seven (7) samples initially and four (4) semiannually	Meet Table B3-8 QAOs	Repeat until acceptable
Laboratory blanks	One (1) per analytical batch	$\leq 3 \times \text{IDL}$ ($\leq 5 \times \text{IDL}$ for ICP-MS) ^b	Redigest and reanalyze any samples with analyte concentrations which are $\leq 10 \times$ blank value and $\geq 0.5 \times$ PRQL
Matrix spikes	One (1) per analytical batch	Meet Table B3-8 accuracy QAOs	Nonconformance if %R outside the range specified in Table B3-8
Matrix spike duplicates	One (1) per analytical batch	Meet Table B3-8 accuracy and precision QAOs	Nonconformance if RPDs > values and %Rs outside range specified in Table B3-8
ICP-MS Tune (ICP-MS Only)	Daily	4 Replicate %RSD ≤ 5 ; mass calibration within 0.9 amu; resolution < 1.0 amu full width at 10% peak height	Nonconformance if %RSD > 5; mass calibration > 0.9 amu; resolution > 1.0 amu
Initial Calibration 1 blank, 1 standard (ICP, ICP-MS) 3 standard, 1 blank (GFAA, FLAA) 5 standard, 1 blank (CVAA, HAA)	Daily	90-110 %R (80-120% for CVAA, GFAA, HAA, FLAA) for initial calibration verification solution. Regression coefficient ≥ 0.995 for FLAA, CVA, GFAA, MAA	Correct problem and recalibrate; repeat initial calibration
Continuing Calibration	Every 10 samples and beginning and end of run	90-110% for continuing calibration verification solution. (80-120% for CVAA, GFAA, HAA, FLAA)	Correct problem and recalibrate; rerun last 10 samples
Internal Standard Area Verification (ICP-MS)	Every Sample	Meet SW-846 Method 6020 criteria	Nonconformance if not reanalyzed at $5 \times$ dilution until criteria are met
Serial Dilution (ICP, ICP-MS)	One (1) per analytical batch	$5 \times$ dilution must be $\leq 10\%$ D of initial value for sample > $50 \times \text{IDL}$	Flag Data if >10% and > $50 \times \text{IDL}$

QC Sample	Minimum Frequency	Acceptance Criteria	Corrective Action ^a
Interference Correction Verification (ICP, ICP-MS)	Beginning and end of run or every 12 hours (8 for ICP) whichever is more frequent	80-120% recovery for analytes Note: Acceptance Criteria and Corrective Action apply only if interferences found in samples at levels greater than ICS A Solution	Correct problem and recalibrate, nonconformance if not corrected
Laboratory Control Samples	One (1) per analytical batch	Table B3-8 accuracy QAOs	Redigest and reanalyze for affected analytes; nonconformance if not reanalyzed
Blind audit samples	Samples and frequency controlled by the Solid PDP Plan	Specified in the Solid PDP Plan	Specified in the Solid PDP Plan

1 ^a Corrective action per Section B3-13 when final reported QC samples do not meet the acceptance criteria.

2 Nonconformances do not apply to matrix related exceedances.

3 ^b Applies only to concentrations greater than the PRQLs listed in Table B3-8.

- 4 IDL = Instrument Detection Limit
- 5 PDP = Performance Demonstration Program
- 6 PRQL = Program Required Quantitation Limit
- 7 %R = Percent Recovery
- 8 RPD = Relative Percent Difference

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**TABLE B3-10
 MINIMUM TRAINING AND QUALIFICATIONS REQUIREMENTS ^a**

Personnel	Requirements ^a
Radiography Operators ^c	Site-specific training based on waste matrix codes and waste material parameters; requalification every 2 years
FTIRS Technical Supervisors ^b FTIRS Operators ^c	Site-specific and on-the-job training based on the site-specific FTIRS system; requalification every 2 years
Gas Chromatography Technical Supervisors ^b Gas Chromatography Operators ^c	B.S. or equivalent experience and 6 months previous applicable experience
Gas Chromatography/Mass Spectrometry Operators ^c Mass Spectrometry Operators ^c	B.S. or equivalent experience and 1 year independent spectral interpretation or demonstrated expertise
Gas Chromatography/Mass Spectrometry Technical Supervisors ^b Mass Spectrometry Technical Supervisors ^b Atomic Absorption Spectroscopy Technical Supervisors ^b Atomic Absorption Spectroscopy Operators ^c Atomic Mass Spectrometry Operators ^c Atomic Emission Spectroscopy Operators ^c	B.S. or equivalent experience and 1 year applicable experience
Atomic Mass Spectrometry Technical Supervisors ^b	B.S. and specialized training in Atomic Mass Spectrometry and 2 years applicable experience
Atomic Emission Spectroscopy Technical Supervisors ^b	B.S. and specialized training in Atomic Emission Spectroscopy and 2 years applicable experience.

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^a Based on requirements contained in *USEPA Contract Laboratory Program Statement of Work for Organics Analysis* (Document Number OLM 01.0) and *Statement of Work for Inorganics Analysis* (Document Number ILM 03.0).

^b Technical Supervisors are those persons responsible for the overall technical operation and development of a specific laboratory technique. QAPjPs shall include the site-specific title for this position.

^c Operators are those persons responsible for the actual operation of analytical equipment. QAPjPs shall include the site-specific title for this position.

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**TABLE B3-11
 TESTING BATCH DATA REPORT CONTENTS**

Required Information	Radiography	Visual Examination	Comment
Batch Data Report Date	X	X	
Batch number	X	X	
Waste container number	X	X	
Waste stream name and/or number	O	O	
Waste Matrix Code	X	X	Summary Category Group included in waste matrix code
Implementing procedure (specific version used)	X	X	If procedure cited contains more than one method, the method used must also be cited. Can use revision number, date, or other means to track specific version used.
Container type	O	O	Drums, Standard Waste Box, Ten Drum Overpack, etc.
Video media reference	X	X	Reference to Video media applicable to each container. For visual examination of newly generated waste, video media not required if two trained operators review the contents of the waste container to ensure correct reporting.
Imaging check	O		
Camera check		O	
Audio check	O	O	
QC documentation	X	X	
Verification that the physical form matches the waste stream description and Waste Matrix Code.	X	X	Summary Category Group included in waste matrix code
Comments	X	X	
Reference to or copy of associated NCRs, if any	X	X	Copies of associated NCRs must be available.

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Required Information	Radiography	Visual Examination	Comment
Verify absence of prohibited items	X	X	
Operator signature and date of test	X	X	Signatures of both operators required for Visual Verification of Acceptable Knowledge
Data review checklists	X	X	All data review checklists will be identified

- 1 LEGEND:
- 2 X - Required in batch data report.
- 3 O - Information must be documented and traceable; inclusion in batch data report is optional.

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**TABLE B3-12
 SAMPLING BATCH DATA REPORT CONTENTS**

Required Information	Headspace Gas	Solid Sampling	Comment
Batch Data Report Date	X	X	
Batch number	X	X	
Waste stream name and/or number	O	O	
Waste Matrix Code		X	Summary Category Group included in Waste Matrix Code
Procedure (specific version used)	X	X	If procedure cited contains more than one method, the method used must also be cited. Can use revision number, date, or other means to track specific version used.
Container number	X	X	
Container type	O	O	Drums, Standard Waste Box, Ten Drum Overpack, etc.
Sample matrix and type	X	X	
Analyses requested and laboratory	X	X	
Point of origin for sampling	X	X	Location where sample was taken (e.g., building number, room)
Sample number	X	X	
Sample size	X	X	
Sample location	X	X	Location within container where sample is taken. (For HSG, specify what layer of confinement was sampled. For solids, physical location within container.)
Sample preservation	X	X	
Person collecting sample	X	X	
Person attaching custody seal	O	O	May or may not be the same as the person collecting the sample
Chain of custody record	X	X	Original or copy is allowed
Sampling equipment numbers	X	X	For disposable equipment, a reference to the lot

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Required Information	Headspace Gas	Solid Sampling	Comment
Drum age	X		Must include all supporting determinative information, including but not limited to packaging date, equilibrium start time, storage temperature, and sampling date/time. If Scenario 3 is used, the packaging configuration, filter diffusivity, liner presence/absence, and rigid liner vent hole diameter used in determining the DAC must be documented. If Scenario 1 and 2 are used together, the filter diffusivity and rigid liner vent hole diameter used in determining the DAC must be documented. If default values are used for retrievably stored waste, these values must clearly be identified as such.
Cross-reference of sampling equipment numbers with associated cleaning batch numbers	O	X	As applicable to the equipment used for the sampling. For disposable equipment, a reference to the lot and procurement records to support cleanliness is sufficient
Drum age	X		
Equilibration time	X		
Verification of rigid liner venting	X		Only applicable to containers with rigid liners
Verification that sample volume taken is small in comparison to the available volume	X		Must include headspace gas volume when it can be estimated
Scale Calibration		O	
Depth of waste		X	For newly generated waste, if a sampling method other than coring is used, this is replaced by documentation that a representative sample has been taken.
Calculation of core recovery		X	For newly generated waste, if a sampling method other than coring is used, this is replaced by documentation that a representative sample has been taken.
Co-located core description		X	For newly generated waste, if a sampling method other than coring is used, this is replaced by documentation that a QC sample has been taken.
Time between coring and subsampling		X	Only applicable to coring.
OVA calibration and reading	O		Only applicable to manifold systems. Must be done in accordance with manufacturer's specifications

Required Information	Headspace Gas	Solid Sampling	Comment
Field Records	X	X	Must contain the following as applicable to the sampling method used: Collection problems, Sequence of sampling collection, Inspection of the solids sampling area, Inspection of the solids sampling equipment, Coring tool test, random location of sub-sample, canister pressure, and ambient temperature and pressure.
Reference to or copy of associated NCRs, if any	X	X	Copies of associated NCRs must be available.
Operator Signature and date and time of sampling	X	X	
Data review checklists	X	X	All data review checklists will be identified

- 1 LEGEND:
- 2 X - Required in batch data report.
- 3 O - Information must be documented and traceable; inclusion in batch data report is optional.

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**TABLE B3-13
ANALYTICAL BATCH DATA REPORT CONTENTS**

Required Information	Headspace Gas	Solid Sampling	Comment
Batch Data Report Date	X	X	
Batch number	X	X	
Sample numbers	X	X	
QC designation for sample	X	X	
Implementing procedure (specific version used)	X	X	If procedure cited contains more than one method, the method used must also be cited. Can use revision number, date, or other means to track specific version used.
QC sample results	X	X	
Sample data forms	X	X	Form should contain reduced data for target analytes and TICs
Chain of custody	X	X	Original or copy
Gas canister tags	X		Original or copy
Sample preservation	X	X	
Holding time		X	
Cross-reference of field numbers to laboratory sample numbers	X	X	
Date and time analyzed	X	X	
Verification of spectra used for results	O	O	Analyst must qualitatively evaluate the validity of the results based on the spectra, can be implemented as a check box for each sample
TIC evaluation	X	X	
Reporting flags, if any	X	X	Table B3-14 lists applicable flags
Case narrative	X	X	
Reference to or copy of associated NCRs, if any	X	X	Copies of associated NCRs must be available.
Operator signature and analysis date	X	X	
Data review checklists	X	X	All data review checklists will be identified

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LEGEND:

- X - Required in batch data report.
- O - Information must be documented and traceable; inclusion in batch data report is optional.

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**TABLE B3-14
 DATA REPORTING FLAGS**

DATA FLAG	INDICATOR
B	Analyte detected in blank (Organics/ Headspace gases)
B	Analyte blank concentration greater than or equal to 20 percent of sample concentration prior to dilution corrections (Metals)
E	Analyte exceeds calibration curve (Organics/ Headspace gases)
J	Analyte less than PRQL but greater than or equal to MDL (Organics/ Headspace gases)
J	Analyte greater than or equal to IDL but less than 5 times the IDL before dilution correction (Metals)
U	Analyte was not detected and value is reported as the MDL (IDL for Metals)
D	Analyte was quantitated from a secondary dilution, or reduced sample aliquot (Organics/ Headspace gases)
Z	One or more QC samples do not meet acceptance criteria
H	Holding time exceeded

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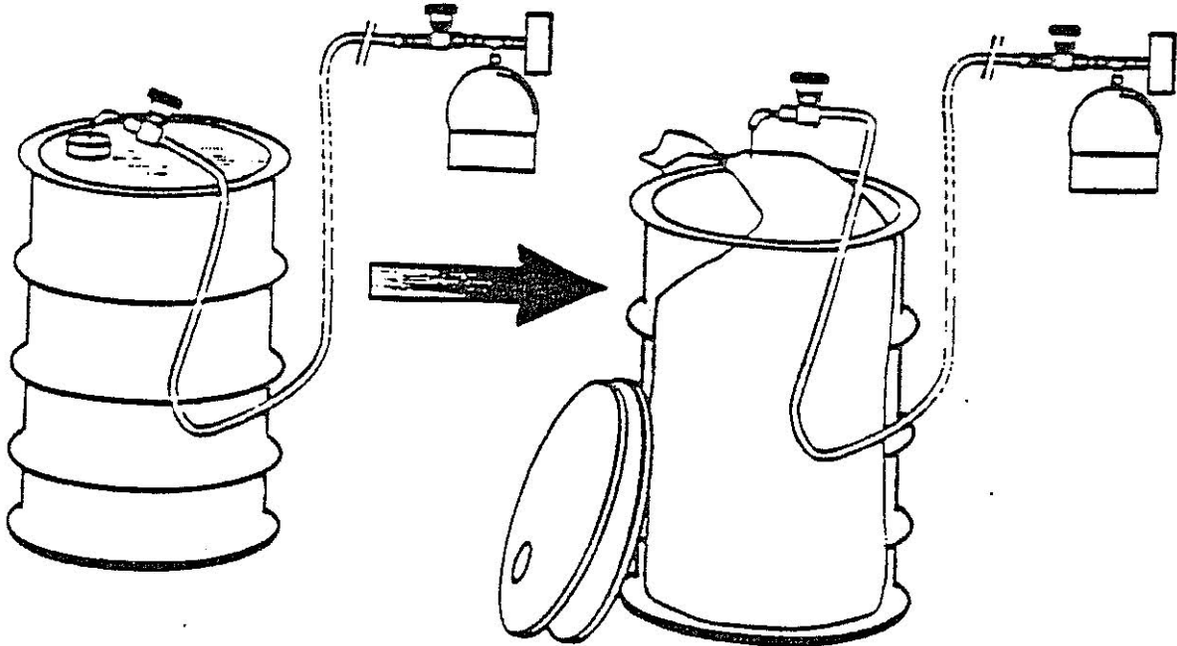
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FIGURES

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Figure B3-1
Overall Headspace-Gas Sampling Scheme Illustrating Manifold Sampling