

Development of Procedures To Assess The Performance Of External Leak Detection Devices

Executive Summary - Draft

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DEVELOPMENT OF PROCEDURES TO ASSESS THE PERFORMANCE OF EXTERNAL LEAK DETECTION DEVICES

EXECUTIVE SUMMARY

DRAFT

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SECTION 1

INTRODUCTION

Underground storage systems comprised of tank, piping, and ancillary components that contain hydrocarbons or other hazardous materials represent a potential source of environmental contamination. Proper system design, installation, operation, and maintenance, along with a leak detection program, can minimize the detrimental effects of leaking underground storage tanks and piping. Devices capable of detecting petroleum hydrocarbons lost from underground storage tank (UST) systems can be used inside an UST system, i.e., in-tank, or external to an UST system, i.e., out-of-tank. In-tank (internal) UST leak detectors generally detect losses with liquid-level sensors. Out-of-tank (external) UST detection systems measure the presence of liquid- or vapor-phase hydrocarbons. Early detection of liquid- or vapor-phase petroleum hydrocarbons allows leaking UST systems or components to be removed from service and repaired or replaced, thereby minimizing both environmental impairment and economic loss of product.

Historically, external (out-of-tank) petroleum hydrocarbon leak and release detection devices have not been extensively used and, therefore, are used primarily in conjunction with new underground storage tank installations. However, most external leak and release detection systems can be retrofitted at existing facilities. Since existing UST facilities are potentially at greater risk of failure (due to age), it is extremely important that leak and release detection devices be used to detect releases of petroleum hydrocarbons from UST installations.

There are numerous commercially available external leak and release detecting devices (for petroleum hydrocarbons) designed exclusively for use with UST systems; however, there are no established performance specifications

or certification procedures for assessing the capabilities of these devices. The U.S. Environmental Protection Agency (EPA) is currently promulgating regulations and guidelines that address underground storage tanks. One aspect of this program is to develop simple, benchmark test procedures that can be used to assess the performance of external petroleum hydrocarbon leak and release detection devices. Performance test results for external leak and release detectors in actual field use may vary from performance measured by these benchmark tests.

Radian Corporation, as a subcontractor to Camp Dresser McKee under contract to EPA/EMSL, is providing technical support in developing test procedures for performance evaluation of out-of-tank (external) leak or release detection equipment (components) for petroleum hydrocarbons. Several tasks critical to this program are complete or currently underway. Results of a survey of vendors of external petroleum leak monitoring devices for use with underground storage tanks have already been published (1). Standard test procedures are under development to allow objective comparisons of external leak and release detector performance. Draft performance methods designed to be simple benchmarks for comparison of external leak and release detectors under prescribed conditions have been submitted to EPA (2).

OBJECTIVES

The overall objective of Radian's UST work is to support EPA's regulatory development. Radian's key contributions are to identify parameters that affect leak detector performance and to develop standard test procedures for assessing external petroleum hydrocarbon detector performance within the meaningful ranges for each performance parameter. The objective of the current task (Work Assignment 02) was to develop and execute draft standard test procedures for performance evaluation of external detection systems for leak and release monitoring. Procedures that address specificity, accuracy, and response time for both vapor-phase and liquid-phase external detectors have been submitted to EPA in a draft report (2).

The performance evaluation test procedures submitted to EPA for review were written as a laboratory test plan for Radian personnel involved in developing formal procedures. Each procedure will require validation before it is used by the public. These performance evaluation procedures will also require revision to provide further clarification, example calculations, and an increased level of detail for performance by an operator. A final collection of performance evaluation test procedures for external leak and release detection components will be submitted in a single document at a future date for EPA review and validation.

REPORT ORGANIZATION

Preliminary conclusions and a summary of results from initial execution of the performance evaluation test procedures for a limited number of external leak and release detection devices are given in Section 2. Section 3 provides a description of the technical approach for this work assignment in the context of prior work. Section 4 gives a preliminary discussion of the results from initial testing of external leak and release detection equipment using the test procedures for evaluating performance. References, mathematical formulas used for vapor-phase and liquid-phase detector performance calculations (Appendix A), and a glossary are given at the end of this summary report.

SECTION 2

PRELIMINARY SUMMARY CONCLUSIONS

Test procedures used to date to assess the performance of external petroleum hydrocarbon leak and release detection systems were developed to provide a uniform, inexpensive, benchmark protocol for comparing parameters such as specificity, accuracy, bias, precision, and detection time. Although only 13 detectors have been tested by Radian, the procedures appear to be applicable to a wide variety of external petroleum hydrocarbon detectors. Categorization of external detectors provided adequate separation of operating principles for ease of performance testing. Intermittent and continuous detectors, separated into liquid-phase and vapor-phase categories, were easily tested with the same procedures.

No major problems or obstacles were reported by Radian personnel performing these initial test runs, and it is likely that these procedures could be performed easily by manufacturers or technicians knowledgeable in UST technology.

Preliminary test results of the performance evaluation procedures indicate that external vapor-phase and liquid-phase detectors respond to most petroleum hydrocarbon components; however, specific detection systems (especially vapor-phase sensors) may have better response factors for certain compound classes (aromatics or aliphatics). These results also indicate that the 250 ppmv concentration used for specificity testing of vapor-phase detectors may be too low for evaluation of some types of qualitative-output detectors. All the liquid-phase detectors responded within 24 hours or less to a 0.5-inch thickness of petroleum product layer floating on water.

Liquid-phase accuracy, bias, precision, and response time results appear to be independent of the composition of the hydrocarbon test liquid; therefore, the simulated standard gasoline mixture probably provides a good representation of a commercial product while offering the advantages of a uniform, laboratory-generated test liquid. In general, the liquid-phase detectors that were tested responded in the presence of a hydrocarbon. A larger range of accuracy was observed for the vapor-phase detectors that were tested.

Detection times for vapor-phase detectors ranged from less than 15 seconds to 24 minutes, with most of the responses occurring in less than 1 minute. Vapor-phase fall time values were all under 15 minutes, and most values were under 1 minute. Detection times for liquid-phase detectors were generally less than 2 hours and were less than 15 minutes for most detectors when the hydrocarbon layer thickness was above the vendor's specifications for lower detection limit.

Future testing of a wider array of external leak and release detection devices may require modification to test containers and chambers to accommodate all detector probe configurations. New performance evaluation procedures may be required to address the scientific principles of operation of future detection equipment as technology presents new modes of external leak and release monitoring.

For test procedures to progress past the draft stage, validation tests will be required to evaluate each procedure's acceptability, and possibly, the relationship of leak detector performance in the laboratory to leak detector performance in the field.

SECTION 3

SUMMARY OF TECHNICAL APPROACH

VENDOR SURVEY AND PERFORMANCE CRITERIA (WORK ASSIGNMENTS 44 AND 70)

The external petroleum hydrocarbon leak and release detection devices that were identified in a prior study (1) were categorized by function into four groups (i.e., intermittent and continuous devices for both vapor-phase and liquid-phase petroleum hydrocarbon detection). Separate performance evaluation test procedures are being developed for vapor-phase and liquid-phase petroleum hydrocarbon detection systems, but intermittent and continuous detection systems of each type are being tested using identical procedures. Parameters that describe petroleum hydrocarbon leak and release detection device performance have been identified (3) and are being used to develop performance evaluation test methods. Parameters related to long-term performance (reliability), safety, and cost were not considered in the development of the present test procedures.

FIRST DRAFT PERFORMANCE EVALUATION TEST PROCEDURES (WORK ASSIGNMENT 92)

In March 1987, a first draft of example performance evaluation procedures was submitted to EPA for review (4). These procedures addressed: 1) specificity of vapor-phase leak and release detection equipment and 2) accuracy and response time for liquid-phase detectors. These test procedures were based upon information from the literature, vendor and manufacturer testing, and engineering and scientific knowledge of the equipment to be tested.

Prior to writing the first draft, performance parameters were grouped together, where possible, to save time and resources and to decrease the

variability of reproduced test procedure performances. The performance parameters were combined in some cases, as shown in Table 1, so that initially a total of 10 performance evaluation test procedures were to be written. Procedures have been written in EPA's standard format. The generic performance procedure outline is given in Table 2. The following paragraphs describe each of the outlined sections' objectives.

The <u>Scope and Application</u> section of each procedure names and describes the performance parameter(s) that the procedure was designed to measure, such as accuracy, signal-to-noise ratio, or specificity. The applicability and limitations of the procedure are discussed. Procedures are specific to either vapor-phase or liquid-phase detectors but are usually applicable to both intermittent and continuous external leak and release detectors. Technical limitations of the method are given.

The <u>Summary</u> section provides a brief overview of operational concepts and principles that are necessary for performing the procedure.

The <u>Interferences</u> section is a detailed guide on how to detect and minimize potential interferences. Specific recommendations on how to measure the effects of contamination are made in this section.

The <u>Safety</u> section provides information and guidelines that are needed to perform a procedure safely. This section discusses minimum safety practices that must be implemented to reduce the possibilities of an accident or exposure.

The <u>Apparatus and Materials</u> section describes equipment, materials, and specifications for each. Descriptions of test containers or chambers, gas rotameters, manometers, and timers are included in this section. Materials such as tubing and fittings are covered also.

FABLE 1. PROPOSED PERFORMANCE EVALUATION TEST PROCEDURE MATRIX FOR EXTERNAL LEAK AND RELEASE DETECTION EQUIPMENT

- 1) Test Procedure for Measuring Accuracy and Response Time for Liquid-Phase Out-of-Tank Petroleum Hydrocarbon Leak and Release Detectors
- 2) Test Procedure for Measuring Accuracy and Response Time for Vapor-Phase Out-of-Tank Petroleum Hydrocarbon Leak and Release Detectors
- 3) Test Procedure for Determining the Specificity of Liquid-Phase Out-of-Tank Petroleum Hydrocarbon Leak and Release Detectors
- 4) Test Procedure for Determining the Specificity of Vapor-Phase Out-of-Tank Petroleum Hydrocarbon Leak and Release Detectors
- 5) Test Procedure for Determining the Detection Limit and the Signal-to-Noise Ratio of Liquid-Phase Out-of-Tank Petroleum Hydrocarbon Leak and Release Detectors
- 6) Test Procedure for Determining the Detection Limit and the Signal-to-Noise Ratio of Vapor-Phase Out-of-Tank Petroleum Hydrocarbon Leak and Release Detectors
- 7) Test Procedure for Measuring Drift for Liquid-Phase Out-of-Tank Petroleum Hydrocarbon Leak and Release Detectors
- 8) Test Procedure for Measuring Drift for Vapor-Phase Out-of-Tank Petroleum Hydrocarbon Leak and Release Detectors
- 9) Test Procedure for Determining the Effect of Potential Interferences of Liquid-Phase Out-of-Tank Petroleum Hydrocarbon Leak and Release Detectors
- 10) Test Procedure for Determining the Effect of Potential Interferences of Vapor-Phase Out-of-Tank Petroleum Hydrocarbon Leak and Release Detectors

^aSubmitted to EPA in draft form for review in March 1988.

TABLE 2. OUTLINE FOR PERFORMANCE EVALUATION TEST PROCEDURES FOR EXTERNAL LEAK AND RELEASE DETECTION EQUIPMENT

1. Scope and Application

Method Covers (e.g., detection limit)
Application (e.g., continuous gas-phase detectors)
Limitations
Experienced Personnel to Employ Method

2. Summary of Method

Brief Discussion of Method Principles Optional Method(s)

3. Interferences

Sources of Contamination Practices/Materials to Avoid QC Checks for Contamination

4. Safety

Compliance to OSHA Regulations Special Chemical Concerns Safety Procedures

5. Apparatus and Materials

Major Systems
Testing equipment
Functioning ranges
Specifications
Alternative systems

6. Test Gases or Liquids

Chemicals/Gases

Define if necessary
Purchasing information
Procedures to make solutions
Storage requirements
Expiration
QC reagents discussed in QC section

7. Calibration

Per Instrument Used
Stepwise procedures
Alternative Procedures (include equations/define variables)

(Continued)

TABLE 2. (Continued)

8. Quality Control

Established QC Program
Minimum requirements
Record keeping
Special procedures
Accuracy and precision
Blanks
Spikes
Ongoing system control

9. Procedure

Stepwise Guide Through Method, Including:
Operating conditions
Estimated responses
Calibration
QC
Atypical procedures

10. Calculations

Discussion of All Calculations Definition All Variables Used

11. Method Performance (i.e., how reliable is this method)

Reliability of Method
Specific ranges (if applicable)
Operator
Overall
Method Accuracy

12. Definitions

For terms that have meanings that are particular to the method or are ambiguous

13. References

The <u>Reagent</u> section describes requirements for all chemicals that are used in a procedure. This section is entitled "Test Gases" or "Test Liquids" for vapor-phase or liquid-phase methods, respectively. Requirements include concentrations, tolerances, and quality or purity. Storage recommendations and necessary mixing instructions are also given.

The <u>Calibration</u> section provides a stepwise guide for calibrating equipment. This equipment may include rotameters, timers, thermocouples, etc.

Frequency and acceptance criteria for calibrations are provided. Calibration of actual petroleum hydrocarbon detectors is only briefly discussed in this section. Complete calibration instructions for detectors should be provided by the manufacturer or vendor.

The <u>Quality Control</u> section describes minimum steps and acceptance criteria that are necessary to establish that procedure results are reliable. Quality control steps may include replicate analyses, record keeping, and blank testing. Frequency, acceptance criteria, and corrective actions for quality control methods are listed in this section.

The <u>Procedure</u> section includes detailed step-by-step instructions of how to perform the procedure. This section guides the technician through equipment assembly, quality control procedures, calibration, testing, and expected results.

The <u>Calculations</u> section provides all the equations necessary to perform the procedure and reduce resulting data. Equations for quality control, data manipulation, and data reporting are provided.

The <u>Method Performance</u> section gives the technician an impression of the procedure's reliability. This section addresses procedure accuracy, repeatability, reproducibility, range, applicability, ruggedness, testing duration, and limitations. (This section can be completed only after validation testing

and is incomplete in the methods that have thus far been submitted for review.)

The <u>Definitions</u> section is a list of terms and their meanings that apply specifically to these methods.

References provide a list of references that were used in writing the procedure.

DEVICE ACQUISITION (WORK ASSIGNMENT 02)

After sets of representative vapor-phase and liquid-phase detection devices were identified and categorized, suppliers were contacted. In August 1987, Radian Corporation began acquiring a limited number of external leak and release detection devices from vendors to undergo initial execution of newly drafted performance evaluation test procedures. Efforts were made to acquire several devices from each category of sensor type established in the prior vendor survey (1).

Several selection criteria were established to guide selection of the external leak and release detection devices to undergo initial performance testing. The selection criteria were: detector type, number of devices in service, specificity for UST leak detection market, and commercial availability. Vendors were contacted; agreements with Radian to borrow the sensor systems for testing were executed; and the devices were sent to the Radian laboratories for performance evaluation testing. A more detailed description of the device acquisition plan was presented in a draft technical note to EPA (5) and subsequently updated in monthly progress reports.

Devices that were acquired represented several categories of scientific operating principles within the larger categories of liquid-phase and vapor-phase leak and release detection devices. The following types of external petroleum hydrocarbon liquid-phase leak and release detection devices were

acquired: electrical conductivity sensors, interface probes, productpermeable probes, and product-soluble sensors. External petroleum hydrocarbon
vapor-phase leak and release detection devices were representative of the
following sensor classes: metal oxide semiconductors, diffusion sensors,
catalytic sensors, photoionization sensors, and infrared sensors.

SECOND DRAFT PERFORMANCE EVALUATION TEST PROCEDURES (WORK ASSIGNMENT 02)

After the draft performance evaluation test procedures were executed on a limited basis in the Radian laboratories, these procedures were modified from the procedures given in the Quality Assurance Project Plan for Work Assignment 02 (6). The second draft performance evaluation test procedures were submitted to EPA in March 1988 (2). Modifications included minor changes to the test apparatuses to accommodate different sensor shapes and sizes.

PRELIMINARY LEAK DETECTION PERFORMANCE DATA

Only 13 external petroleum hydrocarbon leak and release detectors were tested using the draft procedures. No more than two detectors representing any one operating principle were tested. Therefore, the discussion of results (Section 4) of this initial phase of testing using draft procedures is limited. No specific conclusions have been made about the adequacy of operating principles from performance of the draft test procedures for performance evaluation. Since these methods have not been validated, these data should be treated as preliminary.

VALIDATION OF PERFORMANCE EVALUATION TEST PROCEDURES

Validation tests will be developed and performed in the next phase of this program. Individual tests for validating each of the draft performance evaluation test procedures will be developed. The major objective of these validation test procedures will be to evaluate the ability of each of the draft procedures to produce valid test data within acceptable error limits.

The results from performing the validation tests (in a future work assignment) will be used to identify any limitations in the performance evaluation test procedures.

The exact details of each validation test will depend on the associated performance test procedure. The overall approach to be used in developing the validation process will be the same for all cases and is patterned after the successful approach developed by the EPA for the new source performance test methods. The overall approach consists of the following steps:

- Technical examination;
- Laboratory evaluation;
- Ruggedness testing; and
- Validation testing.

Subsequent to the validation testing, final versions of performance evaluation test procedures will be submitted for EPA review. Final validation may require establishing the relationship of laboratory performance to field performance for network leak detection design.

COMPLETION OF PROPOSED PERFORMANCE EVALUATION TEST PROCEDURES

The purpose of developing performance evaluation test procedures is to provide a means of establishing a uniform, consistent database of performance characteristics for external leak and release detection and monitoring devices. Over the long-term course of this project, it is likely that other procedures for new technologies and additional performance parameters to those given in Table 1 will be identified. As performance evaluation procedures are refined and validated, the procedures document will be updated.

SECTION 4

RESULTS

This section is a summary of results and observations made during the preliminary execution of performance evaluation test procedures to determine specificity, accuracy, bias, precision, and response time of external petroleum hydrocarbon leak and release detection systems. The test matrix given in the project plan (6) and procedures given in the draft methods manual (2) were followed.

None of the results presented in this section have been validated or approved for disclosure by EPA or any of its contractors. The performance evaluation results obtained for specific devices were secondary to the task of developing test procedures. Although all due diligence was taken in the execution of these procedures, the results summarized below must be considered preliminary for several reasons. First, the draft procedures performed have not yet been scientifically validated. Changes in the procedures may alter the responses observed for some of the devices. Secondly, the number of devices tested was extremely small compared to the universe of devices on the market. Therefore, no correlation trends assigned to specific operating principles can be made at this time. Although a wide variety of operating principles were represented, no more than two detectors from a single category were tested. Again, the major emphasis of this work was to develop test procedures, not to test the devices.

The following discussion is organized by the performance categories: specificity, accuracy, and response time. Within each performance category, the test matrix, test procedure, and results are organized under subcategories for vapor-phase and liquid-phase detectors. Special efforts have been made in

the data presentation to maintain the anonymity of devices and associated principles of operations, as agreed upon by the participating vendors, Radian Corporation, and EPA Environmental Monitoring Services Laboratory in Las Vegas, Nevada.

RESULTS OF SPECIFICITY TESTING

The significance of specificity (degree of response) of an external leak detector is twofold. If a detector's response is not specific, it may respond to compounds in the environment that are not associated with UST petroleum hydrocarbon releases, causing a false alarm (false positive) situation. On the other hand, if the detector is highly specific (selective), then a low probability of detection may result (false negative). Therefore, one of the most important performance evaluation parameters for external petroleum hydrocarbon leak detection systems is specificity. Specificity answers the question, "What will the detector detect?" Specificity is especially important when the stored product is a single-component hydrocarbon.

Proper network design, site-specific factors, and calibration for external detectors will affect field specificity. These issues were not addressed in these laboratory benchmark test procedures.

Vapor-Phase Detectors - Specificity Response

The specificity of a vapor-phase detector was defined for the procedure as a system's ability to indicate either the presence or concentration of a pure chemical or a mixture of petroleum hydrocarbons at 250 ppmv (certified gas concentration in ultrahigh-purity air). The test gases were monocomponent n-hexane, benzene, toluene, n-butane, isobutane, 2-methylpentane, and 3-methylpentane gas standards. Each of the seven vapor-phase leak detection systems was randomly tested with each gas near 250 ppmv with 20% replicate measurements. The replicate quality control test gas was chosen at random for

each leak detector. Detector responses were initially measured in ultrahighpurity air, monitored for response to a test gas, then purged with ultrahighpurity air according to the test matrix given in the project test plan (6)
using the second draft test procedure (2). Results from these tests were used
to calculate response factors for quantitative-output detectors for each test
gas, and activated versus nonactivated responses for qualitative-output detectors for each test gas. These results are summarized in Table 3.

Three qualitative-output vapor-phase detectors (A, B, and C) failed to respond to any of the test gases at a concentration of 250 ppmv. These detectors were subsequently tested with n-hexane at 500 ppmv, and only Detector C responded at the higher concentration. These detectors had similar modes of operation and do not represent the wide variety of qualitative-output vapor-phase detectors on the market.

The four quantitative-output vapor-phase detectors (D, E, F, and G) showed specificity for the test gases, with differences in response factors. Some detectors were specific to the chemical class of compound (aliphatic versus aromatic), while others showed specificity response factors that appeared to be a combination of chemical class and other factors such as molecular weight. For example, Detector D was most specific (had high response factors) for 2-methylpentane, isobutane, n-hexane, and isopentane, respectively. It was less specific (had lower response factors) for n-butane and toluene. While Detector E's specificity was consistent, response factors exceeded 100% for four of the test gases. The gas concentrations tested were at the lower end of Detector E's operational range. Detector F was most specific for benzene and toluene (aromatic compounds) but demonstrated very low specificity response factors for the other test gases consisting of aliphatic compounds. Detector G had accurate specificity for n-hexane and 2-methy1pentane but lower response factors for the other aliphatic compounds (nbutane, isobutane, and isopentane) and very low response factors for benzene and toluene (common aromatic components in fuels).

TABLE 3. SUMMARY OF VAPOR-PHASE LEAK DETECTORS SPECIFICITY TEST RESULTS^a (Percent Response to Test Gases)

	Background			Test	t Gases			
Detector Identification	UHP Air	n- Butane	Isobutane	Isopentane	n- Hexane	2-Methy1- pentane	Benzene	Toluene
Qualitative-Out	put Detector	: s						
A	NA	NA	NA	NA	b	. NA .	NA	NA
В	na '	NA	NA	NA	b	NA	NA	NA
C	NA	NA	NA .	NA	c	NA	NA	NA
Quantitative-Ou	tput Detecto	ors	•					
D	${}_{ m NR}{}^{ m d}$	11	63	57	61	65	42	17
E	nrd	100	200	200	200	200	100	150
F	nrd	1	1	6	10	12	100	102
G	$NR^{\mathbf{d}}$	68	63	77	96	92	2	14

^aSpecificity, the ability of a detector to indicate the presence or concentration of a test gas at 250 ppmv, is reported in terms of response factors calculated as the percent of the detection system's response compared to the certified concentration of the test gas for quantitative-output detectors and as "activated" (A) or "non-activated" (NA) for qualitative-output detectors.

b_{No} response to either 250 ppmv or 500 ppmv of n-hexane.

 $^{^{\}mathrm{c}}$ Activated response to 500 ppmv of n-hexane but no response to 250 ppmv of n-hexane.

d_{NR} = no response for quantitative-output detectors.

Liquid-Phase Detectors - Specificity Response

For liquid-phase petroleum hydrocarbon leak and release detection systems, specificity was defined in the performance evaluation test procedure as the response to a 1.27-cm (1/2-in.) layer on water of different chemical substances representing components or types of petroleum hydrocarbons commonly stored in UST systems. The test liquids used in the performance evaluation procedure were: n-hexane, xylene(s), toluene, commercial gasoline, a laboratory-mixed "standard" gasoline, commercial diesel fuel, and commercial jet fuel. Each of six external liquid-phase petroleum hydrocarbon leak and release detection devices was randomly tested using each test liquid with 20% replicate measurements. The replicate quality control test liquids were chosen at random for each liquid-phase detector. Detectors were measured for background responses in tap water, then for responses in each test liquid with a complete cleaning of the test apparatus between tests. The matrix from the project test plan (6) and the second draft test procedure (2) were followed. Results from these tests were used to determine if there was a response for specific compounds within 24 hours.

Test results indicated five of the six liquid-phase detectors responded to a 1.27-cm (1/2-in.) thick layer for all seven test liquids on water within 24 hours. The sixth liquid-phase detector responded to all test liquids within 24 hours, except commercial diesel fuel, to which it responded in 26 hours.

The ranges of observed response times during specificity testing are summarized by test liquid in Table 4. The longest time until response for all test liquids was observed for the same detector (Detector D). Likewise, the shortest time required until response to all test liquids was observed for a single detector (Detector B).

Detector D required over 16 hours to respond to n-hexane, xylene(s), diesel fuel, and jet fuel; it responded to toluene, commercial gasoline, and synthetic gasoline mixture in under 2 hours. Detector B showed no specificity

TABLE 4. SUMMARY OF RESPONSE TIMES TO TEST LIQUIDS DURING LIQUID-PHASE SPECIFICITY TESTING

Test Liquid	Ranges of Observed Time Until Response
n-Hexane	<1 sec - 17.5 hr
Xylene(s)	<1 sec - 21.5 hr
Toluene	<1 sec - 1.5 hr
Simulated Gasoline	<1 sec - 1 hr
Commercial Gasoline	<1 sec - 1 hr
Diesel Fuel	<1 sec - 26 hr
Jet Fuel	<1 sec - 16.5 hr

^aThe range is based on the response times for six detectors.

toward any of the test liquids, responding to all of them in less than 1 second. Detector A required over 9 hours to respond to diesel fuel but responded to all other test liquids in less than 20 minutes. Detector E responded to xylene(s), toluene, synthetic gasoline mixture, and commercial gasoline in 2 minutes or less; it required 6 to 7 hours to respond to nhexane, diesel fuel, and jet fuel. Detectors C and F responded to all test liquids in under 10 minutes.

RESULTS OF ACCURACY AND RESPONSE TIME TESTING

Accuracy, in general, is the degree of agreement between measured values and the true values. In these methods, the description of accuracy is quite different for qualitative-output versus quantitative-output detectors. Qualitative accuracy in these methods is a measure of positive responses, where 100% accurate means the detector responded positively 100% of the time. Quantitative accuracy is defined as "relative accuracy," following the convention used for continuous emission monitoring systems where 100% relative accuracy means the maximum expected bias (at the 95% confidence level) is ±100% of the true value. Thus, qualitative and quantitative accuracy in these methods cannot be directly compared. Relative accuracy has no meaning for qualitative—output detectors; however, relative accuracy may serve as the prime comparator for quantitative—output detectors and may be determined in the laboratory against known test conditions or in the field versus a reference method.

Bias is the term used for the systematic error inherent in a method, and it can be positive or negative. Precision is the degree of agreement between repeated measurements of the same parameter, and it reflects random errors that are unaffected by bias. Neither bias nor precision are applicable to qualitative devices, which only produce a binary ("on/off") signal.

Detection time is the sum of lag time (the elapsed time from a detector's first contact with the test gas or liquid to the first detectable signal) and rise time (the elapsed time from a detector's first detectable signal in response to hydrocarbon liquid or gas to an output that is within 95% of full

scale or "activated"). Fall time is the elapsed time from a hydrocarbon-free condition after exposure to hydrocarbon until a detector's output returns to within 95% of its original baseline level or there is no detectable signal output.

Several parameters measured in this suite of tests describe accuracy; others describe time-related statistics that are important to the owner of underground storage tanks. Accuracy answers the question, "How close is the hydrocarbon value measured by a leak detector to the actual hydrocarbon level present?" Bias answers the question, "Is the detection device higher or lower than the real value?" Precision answers the question, "Is the test result repeatable and is the uncertainty in the test result large enough to adversely influence any decision process?" Detection time answers the question, "How fast will the detector signal an alarm after it contacts hydrocarbons?" Fall time answers the question, "How long will it take after a positive response for a sensor placed in a clean environment to cease its positive alarm signal?" Answers to these questions are important because they indicate whether an alarm will occur at an acceptable performance level. Again, the following test results pertain to laboratory performance and not field performance.

Vapor-Phase Detectors - Accuracy Testing

Accuracy for qualitative-output vapor-phase detectors was defined in the test procedure as the number of positive responses expressed as a percentage of the total number of tests (five in this case). For quantitative-output detectors, the relative accuracy was calculated as the absolute mean difference between the measured value and the true value, plus the 2.5% error confidence coefficient for each series of five replicate tests, divided by the true value. It is expressed as a percentage of the reference (certified gas) standard concentration. Accuracy for qualitative-output detectors is neither conceptually nor numerically comparable to relative accuracy for quantitative-output detectors.

Bias, for this method, was expressed as the signed (positive or negative) percent difference between the average measured value for a series of five replicate tests and the true value. Precision was expressed in terms of the percent coefficient of variation (CV), which is equal to the standard deviation for each set of five values, divided by the mean of the set, and multiplied by 100.

Each of six external vapor-phase petroleum hydrocarbon leak and release detection systems was tested with certified gas standards of benzene and isopentane. Four concentration ranges (approximately 50, 250, 500, and 1000 ppmv) were tested for both compounds. Benzene and isopentane were used as test gases because they are common components in petroleum products and many types of vapor-phase detectors are sensitive to them. Concentration levels were chosen based on preliminary information of the working concentration ranges for total hydrocarbons in which vapor release detection might be most appropriate.

The test procedure (2) required an initial background test with ultrahigh-purity air to monitor the response of a test gas; record detection time, response value, and fall time; then repeat these steps until all of the tests were replicated five times for each test gas concentration, according to the matrix of 240 total tests in the project test plan (6). Each of the replicate tests was performed under equivalent test conditions. Duplicate analyses consisted of immediate repetition of test conditions for 10% of the tests chosen at random.

Table 5 summarizes vapor-phase accuracy, bias, precision, and response time test results. In general, some of the vapor-phase detectors were very accurate, while other detectors were less accurate. Some of the vapor-phase detection systems were more accurate for a particular test gas, while others showed similar accuracy for both test gases. Some detectors were more accurate at higher test gas concentrations, while others were just as accurate at the lower test gas concentrations as they were at the higher concentrations.

TABLE 5. VAPOR-PHASE ACCURACY, BIAS, PRECISION, AND RESPONSE TIME RESULT SUMMARY

			Accur	всу				
etector ⁸	Test Gas	Concentration (ppmv)	Qualitative Accuracy (%)	Relative Accuracy (%)	Biss (%)	Precision (%)	Detection Time (minutes)	Fall Time (minutes)
ualitative	Detectors				•	٠		
HALLEGEIVE	Detectors		,		•		ND ^f	1775
В	Benzene	48	$\mathtt{NT}^{\mathbf{d}}$		NL^e	NL		ND
В	Benzene	240	NT		NL	NL	ND	ND
В	Benzene	500	NT		NL	. NL	ND	ND
В	Benzene	990	. 0		NL	NL	ND	ИD
_	_	49	NT		NL	NL	ND	ND
В	Isopentane	253	NT .		NL	NL	ND	ND
В	Isopentane	253 499	0		NL	NL	ND	ND
В	Isopentane	•	0		NL	NL	ND	ND
В	Isopentane	991						
С	Benzene	48	NT		NL	NL	ND	ND
Č	Benzene	240	0		NL	NL	ND	ND
C	Benzene	500	100		NL	NL	42.40	3.40
C	Benzene	990	100		NL	NL	9.60	4.40
			•		NL	NL	ND	ND
C	Isopentane	49	0		NL	NL	24.00	3.20
C	Isopentane	253	100		NL NL	NL	42.00	7.80
С	Isopentane	499	100 100		NL	NL	42.40	3.20
С	Isopentane	991	100		NL	WD	12110	••••
Quantitatiy	ve Detectors							
	7	48		44	-38	7	8.00	8.00
D	Benzene	240		58	-55	6	8.00	8.00
D	Benzene	500	an ross	61	-56	8	8.00	8.00
D	Benzene	990		64	-55	16	16.60	8.00
D	Benzene	990		04				

(Continued)..

TABLE 5. (Continued)

			Accur	всу				
Oetector ^a	Test Gas	Concentration (ppmv)	Qualitative Relative Accuracy Accuracy (%) (%)		Bias (%)	Precision (%)	Detection Time (minutes)	Fall Time (minutes)
uantitativ	e Detectors (Con	nt.)						
G	Benzene	48		99	-99	39	1.15	0.30
G	Benzene	240		98	-98	8	0.25	0.25
G	Benzene	500	-	98	-98	. 5	0.60	0.30
G	Benzene	990	معد حط وشي	98	-98	5	0.35	0.35
	Isopentane	49	gas are time	16	-13	3	0.30	0.45
G				17	-16	. 1	0.35	1.05
G G	Isopentane	253		1/	-10			
	•	253 499	*****	20	-18	· 2	0.30	2.30

^aDetector A was not tested because it did not respond during specificity tests.

b Qualitative Accuracy - For qualitative-output detectors, the number of positive responses to the test gas, expressed as a percentage of the number of tests (five) performed. A value of 100% means the detector responded appropriately for all five tests in the series; 0% means the detector failed to respond during any of the tests.

CRelative Accuracy - For quantitative-output detectors, the absolute mean difference between the measured value and the true value for the test gas, plus the 2.5% error confidence coefficient, expressed as a percentage of the true value. A value of 100% for relative accuracy means that, within the 95% confidence interval, the measurement bias can be expected not to exceed +100% of the true value.

d_{NT} = not tested at this concentration.

eNL = not applicable because the detector's output was qualitative.

 f_{ND} = not applicable because the test gas was not detected.

BDetection time and fall time values are only for test gas responses that were above baseline responses with ultrahigh-purity air.

TABLE 5. (Continued)

			Accur	асу				
Detector ^a	Test Gas	Test Gas Concentration (ppmv)	Qualitative Accuracy (%)	Relative Accuracy (%)	Bias (%)	Precision (%)	Detection Time (minutes)	Fall Time (minutes)
Ouantitative	e Detectors (Co	nt.)						
				46	-33	16	6.40	7.20
D	Isopentane	49			-42	17	8.00	8.00
D	Isopentane	253		54	-42 24	3	4.00	7.20
D	Isopentane	499		29	T :	. 7	6.40	8.00
D '	Isopentane	991	*****	42	61		0.40	0.00
				607	100	224	0.25 ^g	0.25 ⁸
E	Benzene	48		687	108	224 26	0.45	0.80
E	Benzene	240	*****	370	254		0.50	1.00
E	Benzene	500		150	150	0		
.E	Benzene	990		173	132	14	0.90	2.15
							o org	0.25 ⁸
E	Benzene	· 48		687	108	224	0.25 ^g	0.25
E	Benzene	240		370	254	26	0.45	0.80
Ē	Benzene	500		150	150	0	0.50	1.00
Ē	Benzene	990	***	173	132	14	0.90	2.15
	20							
E	Isopentane	49		1206	512	91	0.55	0.30
E	Isopentane	253		295	295	0	0.30	0.70
E	Isopentane	499		543	321	43	0.65	1.70
E	Isopentane	991		267	233	8	2.40	1.85
P.	Isopencane)) <u>,</u>						
F	Benzene	48		13	10	3	0.25	0.40
		240		9	-6	3	0.30	0.55
F	Benzene	500		20	-17	4	0.25	0.30
F	Benzene	990		35	-34	1	0.25	0.95
F	Benzene	990		J.J		_		
	-	49		95	-93	28	0.25	0.30
F	Isopentane			97	-96	20	0.30	0.25
F	Isopentane	253		98	-96	25	0.30	0.25
F	Isopentane	499		97	- 97	0	0.25	0.20
F	Isopentane	991		71	-31	•	0.23	, ,

(Continued)

Vapor-phase Detectors A, B, and C were qualitative-output detectors. Detector A was not tested because it did not respond to any of the specificity test gases at 250 ppmv or to hexane at 500 ppmv. Its mode of operation was almost identical to Detector B. Detector B did not respond during accuracy testing to either test gas at any concentration tested. Detector C responded in all of the accuracy tests for 500 ppmv and 990 ppmv of benzene and for 253 ppmv, 499 ppmv, and 991 ppmv of isopentane. (The fact that Detector C responded at 253 ppmv in these tests and did not respond at the same concentration during the specificity tests indicates that this concentration may be very close to the detector's detection limits.)

Vapor-phase Detectors D, E, F, and G were quantitative-output detectors. Detector D's relative accuracy ranged from 29% to 64% for benzene and isopentane. Relative accuracy (RA) values were similar for benzene and isopentane at all concentration levels. Detector E demonstrated poorer relative accuracy (RA = 150% to 1206%). Detector F was more accurate for benzene (RA = 9% to 35%) than for isopentane (RA = 95% to 98%). Detector G's relative accuracy values were more accurate for isopentane (RA = 16% to 25%) than for benzene (RA = 98% to 99%).

Average bias values for quantitative-output vapor-phase detectors ranged from -99% to a high of 512%. In general, isopentane bias values tended to decrease as the test gas concentration increased. The best bias values were -6% and 10% for Detector F in benzene at 240 ppmv and 48 ppmv, respectively. Bias is not applicable to qualitative-output detectors, which produce only a binary ("on/off") output signal, and it was not calculated for vapor-phase Detectors A, B, and C. For Detector D, benzene bias was from -56% to -38%. For isopentane, this detector showed negative bias at 49 ppmv and 253 ppmv and positive bias at 499 ppmv and 991 ppmv. The bias magnitudes for both test gases were similar. For Detector E, bias values ranged from 108% to 512%. Isopentane bias values (RA = 233% to 512%) were higher than benzene bias values (RA = 108% to 254%). Benzene bias values for Detector F were from -34%

to 10%, and the bias decreased with increasing test gas concentration. Detector F isopentane bias results ranged closely from -97% to -93%. As with relative accuracy results, Detector F bias results followed an almost opposite trend from bias results for Detector G. Detector G bias was -99% to -98% for benzene and -13% to -24% for isopentane.

In general, all four quantitative-output detectors were very precise. All but one detector showed similar precision values for both test gases. The poorest precision values (91% and 224%) were observed for the lowest test gas concentrations with a single detector. All other precision values were under 50%. Precision is not applicable to qualitative-output detectors, which produce only a binary ("on/off") output signal, and it was not calculated for vapor-phase Detectors B and C.

Detector D's precision ranged from 3% to 17% CV. Precision values for Detector E ranged from 0% to 224% coefficient of variation (CV), showing less precision at the lowest test concentrations for both gases than at higher concentrations. As previously noted, test gases were at the low end of Detector E's operating range. Precision values for Detector F ranged from 0% to 28% CV. Even though this detector's best precision value was for 991 ppmv isopentane, benzene precision values were better than isopentane precision values overall. Detector G precision values were from 1% to 39% CV. Other than a 39% benzene precision value, values were under 10% for Detector G. This detector was slightly more precise for isopentane than for benzene.

Liquid-Phase Detectors - Accuracy Testing

Accuracy for qualitative-output liquid-phase detectors was defined in the test procedure as the number of positive responses for a particular test liquid at a particular thickness, expressed as a percentage of the total number of tests in the series (five in this case). For quantitative-output liquid-phase detectors, the relative accuracy was calculated as the absolute mean difference between the measured value and the true value, plus the 2.5%

error confidence coefficient for each series of five tests, divided by the true value. It is expressed as a percentage of the known thickness of hydrocarbon layer on water. As with vapor-phase detectors, accuracy, as defined in these methods, is neither conceptually nor numerically comparable for qualitative-output versus quantitative-output detectors.

Bias is expressed as the signed (positive or negative) percent difference between the average measured value for each series of five replicate tests and the true value for the test liquid thickness. Precision is expressed in terms of the percent coefficient of variation, which is equal to the standard deviation for each set of five values divided by the mean of the set and multiplied by 100.

Each of six external liquid-phase petroleum hydrocarbon leak and release detection systems was tested with three thicknesses (0.04, 0.32, and 0.64 cm) each of simulated (laboratory-generated) gasoline mixture and commercial gasoline. The simulated gasoline mixture will provide a uniform protocol for future testing; commercial gasolines frequently have differing product compositions that may affect test results.

The test procedure (2) required an initial background test in tap water to monitor the response to each test liquid and thickness as a function of time, then repeating these steps until all of the tests were replicated five times for each liquid product and thickness in accordance with the matrix of 180 total tests given in the project test plan (6). Each of the replicate tests was performed under identical conditions.

Table 6 summarizes the test results for accuracy, bias, precision, and response time testing for liquid-phase petroleum hydrocarbon detectors.

Detector B was the only detector capable of providing quantitative output. Relative accuracy for this detector ranged from 1120% for the simulated

TABLE 6. LIQUID-PHASE ACCURACY, BIAS, PRECISON, AND RESPONSE TIME RESULTS SUMMARY

		Hydro- Detector							
Criteria	Gasoline Type	carbon Thickness (cm)	A	В	С	ם	Е	F	
	Ci-ulanai	0.04	1007ª	1120 %	100% ^a	100% ^a	100% a	0% ^a	
Accuracy	Simulated	0.32	100%a	147% ^b	100% a	100% ²	100% a	100%	
		0.64	100% ^a	147% 61%	100% ^a 100% ^a 100% ^a	100%ª	100%ª	100% ^a 100% ^a	
		2.01		444% b	100% ^a	100% ^a	100% a	0% ^a 100% ^a 100% ^a	
	Commercial	0.04	100%	97%b	100%a	100% a	100% ^a	100% ^a	
		0.32	1002 ^a 1002 ^a 1002 ^a	87% b 42% b	100% ^a 100% ^a 100% ^a	100% a 100% a 100% a	100% ^a 100% ^a	100% ^a	
		0. 64	100%	4 2%	100%	100%	200,0		
	Simulated	0.04	c	606%	c	c	c	c	
Bias	Simulated	0.32	c	114%	c	С	, c	c	
		0.64	c	45%	c	c	c	c	
		0.04	c	. 102%	c	c	c	c	
	Commercial	0.04	c	57%	c	c	c	С	
		0.32	c	42%	č	c	c	, c	
		0.64		720	•				
Precision	Simulated	0.04	đ	59%	đ	đ	đ	đ	
Precision	SIMULATED	0.32	đ	12%	đ	đ	ď	đ	
		0.64	, ā	9%	đ	đ	đ	đ	
	Commercial	0.04	đ	137%	đ	đ	đ	đ	
	COMMETCIAL	0.32	đ	18%	đ	đ	đ	đ	
		0.64	đ	0%	đ	đ	đ	đ	
						104.60	13.30	f	
Detection	Simulated	0.04	6.18	e	2.73	72.83	0.08	0.08	
Time		0.32	0.83	<0.02	0.32	72.83 57.63	0.32	0.17	
(minutes)		0.64	0.58	<0.02	0.32	5/.03	0.32		
	Commercial	0.04	1.37	e	2.42	121.55	8.13	£	
	Commercial	0.32	1.30	<0.02	0.23	64.22	0.13	0.18	
		0.52	0.70	<0.02	0.27	73.10	0.12	0.10	

(Continued)

TABLE 6. (Continued)

Criteria	Gasoline Type	Hydro- carbon Thickness (cm)	Detector						
			A	В	С	D	E	F	
Fall Time (minutes)	Simulated	0.04 0.32 0.64	8 8 8	h k k	i i i	8 8	& & &	j k k	
	Commercial	0.04 0.32 0.64	g g g	h k k	i i i	8 8 8	& & &	j k k	

Qualitative Accuracy - For qualitative-output detectors, the number of positive responses to the test liquid, expressed as a percentage of the number of tests (five) performed. A value of 100% means the detector responded appropriately for all five tests in the series; 0% means the detector failed to respond during any of the tests.

be Relative Accuracy - For quantitative-output detectors, the absolute mean difference between the measured value and the true value for the test liquid, plus the 2.5% error confidence coefficient, expressed as a percentage of the true value. A value of 100% for relative accuracy means that, within the 95% confidence interval, the measurement bias can be expected not to exceed ±100% of the true value.

^CBias not applicable because device is qualitative.

dPrecision was not applicable because the device was qualitative.

^eImmediate response when any hydrocarbon was detected.

f Not applicable because no response was recorded.

gnot applicable because the probe was destroyed during the test.

h Immediate when activation occurs.

ino deactivation after 24 hours in water. Will deactivate in air.

jNot applicable because there was no response.

kImmediate deactivation (fall time <1 second).

gasoline mixture at 0.04 cm (1/64 in.) to 42% at 0.64 cm (1/4 in.) for commercial gasoline. Poor accuracy results at the 0.04 cm (1/64 in.) thickness were expected because the manufacturer claimed a 0.32 cm (1/8 in.) lower detection limit. Accuracy values at 0.32 cm (1/8 in.) and 0.64 cm (1/4 in.) were much better, ranging from 147% to 42%. Detector B showed similar accuracy for both test liquids.

Bias for external liquid-phase petroleum hydrocarbon detectors was only applicable to the one quantitative-output detector (Detector B) that was tested. Bias values for this detector ranged from 606% at 0.04 cm of simulated gasoline to 42% at 0.64 cm of commercial gasoline. As with the relative accuracy values for this detector, bias values for 0.32 cm and 0.64 cm test liquid thicknesses, which were equal to or higher than the detector's lower detection limit, were significantly better than the values at the 0.04 cm test liquid thickness.

Precision for liquid-phase detectors was only applicable to Detector B because it was the only quantitative-output detector tested. Precision ranged from 137% at 0.04 cm (1/64 in.) to 0% at 0.62 cm (1/4 in.) of commercial gasoline. As with relative accuracy and bias, precision values were much better for test liquid thicknesses that were above the detector's lower detection limit. Precision values at the 0.04 cm thickness were 59% and 137% for the synthetic gasoline mixture and commercial gasoline, respectively. Precision values at test thicknesses of 0.32 cm and 0.64 cm ranged from 18% to 0%.

Only one of five qualitative-output detectors failed to respond to test liquids at each test thickness within 24 hours. Detector F, whose manufacturer reported a detection limit of 0.32 cm (1/8 in.), failed to respond to either test liquid at 0.04 cm (1/64 in.). This same detector, however, did respond in all tests to both test liquids at the two higher test liquid thicknesses. In other words, this detector was 0% accurate at a test liquid thickness of 0.04 cm but 100% accurate at test liquid thickness of 0.32 cm and 0.64 cm.

Vapor-Phase Detectors - Response Time Testing

Detection time and fall time results for vapor-phase detectors were given in Table 5. Vapor-phase detectors typically responded faster than their liquid-phase counterparts. The range of average detection times ranged from less than 15 seconds to 1 hour and 24 minutes. The vast majority of responses occurred in under 1 minute. Vapor-phase fall time values were smaller than detection time values. All fall time values were less than 10 minutes, and most values were under 1 minute.

Liquid-Phase Detectors - Response Time Testing

Detection time for liquid-phase detectors, as shown in Table 6, was generally 2 hours or less. For all but one detector, responses occurred in less than 15 minutes if the test liquid was above the detector's lower detection limit.

Detection times for liquid-phase Detectors B and F were very short (<1 to 11 seconds) at all test liquid thicknesses at which these detectors responded. These detectors did not respond at the 0.04 cm test liquid thickness, which was below the detection limit of both devices.

Detectors A, C, and E responded at all three thicknesses of both test liquids in under 15 minutes. The longest average detection time for a set of five replicate tests from this group of detectors was 13.3 minutes for the simulated gasoline mixture at 0.04 cm. The shortest average detection time was 7 seconds for commercial gasoline at 0.64 cm. Detection times were much higher for test liquid thicknesses at 0.04 cm (1 minute, 22 seconds to 13 minutes, 18 seconds) than for 0.32 cm and 0.64 cm test liquid thicknesses (7 seconds to 1 minute, 18 seconds). There did not appear to be significant differences between simulated versus commercial gasoline or test liquid thicknesses of 0.32 cm versus 0.64 cm. Detector D averaged from 57 minutes to 2

hours for responses. Response times between tests with simulated gasoline were not significantly different from those for commercial gasoline.

Fall time was generally not a significant factor for liquid-phase detectors. Fall time was not applicable to Detectors A, D, and E because they required replacement of a product-soluble component that was destroyed during contact with test liquid. Detector C would not return to an inactivated state when tested for fall time in water. This detector would return to an inactivated state if the probe was tested for fall time in air. Detectors B and F had fall times that were less than 1 second if the detectors responded to the test liquid. If these detectors did not respond to a test liquid, fall time was not applicable.

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APPENDIX A

MATHEMATICAL FORMULAS

Vapor-Phase Detectors

Relative accuracy -- The relative accuracy (RA) of a set of data was calculated as follows:

$$RA = \frac{|\overline{d}| + |cc|}{\nabla_r} \times 100 \tag{1}$$

where: $V_{\underline{r}}$ = reference (theoretical) value; \overline{d} = arithmetic mean of the difference of a data set, Equation 1.1; and

cc = 2.5% error confidence coefficient (one tailed, Equation 1.2).

Mean difference -- The arithmetic mean of the difference (d) of a data set was calculated as follows:

$$\overline{d} = \frac{1}{n} \sum_{i=1}^{n} d_{i}$$
 (1.1)

where: d = measured response - theoretical response.

Confidence coefficient -- The one-tailed 2.5% confidence coefficient (cc) was calculated as follows:

$$cc = t_{0.975} \cdot \frac{S}{\sqrt{n}}$$
 (1.2)

where: S =the standard deviation (n-1) of the data set; and $t_{0.975} = 2.5\%$ (one-tailed) t value = 2.776 for n=5.

Accuracy for qualitative-output detectors—For qualitative-output detectors, accuracy for a particular test atmosphere at a particular concentration was calculated as the number of positive responses expressed as a percentage of the total number of tests (five in this case) at that concentration with that test atmosphere, according to the following formula:

$$A_{q} = 100 \times (r_{p}/5) \tag{2}$$

where: A_q = accuracy in percent; and r_p = number of positive responses.

<u>Bias</u>—Bias is a measure of how much, on the average, the quantitativeoutput detector responds high or low with respect to the true (theoretical) response. Bias was calculated as follows:

$$Bias = 100 \times \left[(\overline{V}_{0} - V_{r})/V_{r} \right]$$
 (3)

where: \bar{V}_0 = the average observed value, $\frac{1}{5} \sum_{i=1}^{5} V_i$; V_i = the individual response to a test atmosphere; and V_i = the reference (theoretical) value.

<u>Detection time</u>—Detection time was calculated according to the following formula:

$$DT = T_2 - T_1 \tag{4}$$

where: DT = elapsed detection time;

T₁ = clock time when the test atmosphere was first added to the test chamber; and

T₂ = clock time when the detector output went from an inactivated state to an activated state for a qualitative-output detector or from a baseline reading to 95% of stable high level output for a quantitative-output detector.

Fall time -- Fall time was calculated according to the following formula:

$$\mathbf{FT} = \mathbf{T}_2 - \mathbf{T}_1 \tag{5}$$

where: FT = elapsed fall time;

T₁ = clock time when the test atmosphere was switched to ultrahighpurity air; and

T₂ = clock time when the detector output went from an activated state to an inactivated state for a qualitative-output detector or from a high level reading to within 95% of stable baseline level output for a quantitative-output detector.

Relative percent difference—The relative percent difference is a measure of variation between two observations. It is their absolute difference divided by their average, expressed as a percentage, and was calculated as follows:

$$RPD = 200 \times [(\nabla_1 - \nabla_2)/(\nabla_1 + \nabla_2)]$$
(6)

where: RPD = relative percent difference;

 V_1 = larger value; and

 $\nabla_2 = \text{smaller value.}$

<u>Correlation coefficient</u>—The correlation coefficient was calculated as follows:

$$cc = \frac{s_{xy}}{\sqrt{s_{xx} \cdot s_{yy}}}$$
 (7)

where: cc = correlation coefficient; $S_{xy} = n\sum_{i}y_{i} - (\sum_{i})(\sum_{j}y_{i});$ $S_{xx} = n\sum_{i}^{2} - (\sum_{i}x_{i})^{2};$ $S_{yy} = n\sum_{i}^{2} - (\sum_{j}y_{i})^{2};$ $x_{i} = i^{th} \times value; and$ $y_{i} = i^{th} y \times value.$

<u>Coefficient of variation (%)</u>—The percent coefficient of variation is equal to the standard deviation for a set of values, divided by the mean of the set, times 100 and was calculated as follows:

$$CV(\%) = (S/\overline{X}) \times 100 \tag{8}$$

where: CV = coefficient of variation;

S = standard deviation for n values (n-1 degrees of freedom); and

 \bar{X} = the arithmetic mean of n values.

Response factor (RF) -- Response factors were calculated as follows:

$$RF = C_{m}/C_{t}$$
 (9)

where: C_{m} = the vapor-phase detection system response (ppm-v, %, etc.); and C_{+} = the concentration of the test atmosphere gas (ppm-v, %, etc.).

Liquid-Phase Detectors

<u>Water-miscible substances content--</u>The fraction of water-miscible substances in commercial gasoline test product was calculated with the following equation for a 20-mL sample:

$$w = 1 - \frac{(m_u - m_1)}{20.0} \tag{10}$$

where: w = water-miscible substances content;

m. = upper meniscus volume in mL; and

m, = lower meniscus volume in mL.

Area from immersing in water—The cross-sectional area of a probe was measured by determining the volume occupied by the probe when immersed in water. The following equation was used to calculate the cross-sectional area:

$$\mathbf{a_d} = (\nabla_{\mathbf{i}} - \nabla_{\mathbf{p}})/1.27 \tag{11}$$

where:

a_d = cross-sectional area in cm²;

 V_{i} = volume between marks without probe in mL;

 $V_{\rm D}$ = volume between marks with probe in mL; and

1.27 = height of column of water displaced in cm.

Product volume—Before any testing began, a product layer of accurately known thickness was formed on water in the test container. The volume of test product to add to the test container to create a desired thickness of product was calculated with the following equation:

$$v = \frac{t \times (a_c - a_d)}{1 - w} \tag{12}$$

where: v = volume of product to add to container in mL;

t = desired product thickness in cm;

 $a_c = \text{test container cross-sectional area (193.77 cm}^2);$

 a_d = estimated detector cross-sectional area in cm²; and

w = test product water content (w = 0 for synthetic gasoline).

<u>Product thickness</u>—The thickness of a test product layer on water was not measured but was determined from the volume of product added to the test container, test container dimensions, and detector dimensions. The thickness of product on water was determined with the following formula:

$$t = v/(a_c - a_d) \tag{13}$$

where: t = test product thickness in cm;

v = volume of test product in cm³;

 $a_c = \text{container cross-sectional area (193.77 cm}^2);$ and

 a_d = detector cross-sectional area in cm².

Output at 95% of stable high level output—Detection time for quantitative—output detectors was measured from the time the detector contacted test product until it reached 95% of its final stable output. The following formula was used to calculate the 95% of stable high level output, based on the stable baseline level output and the stable high level output.

$$HB = BL + (HL - BL) \times 0.95 \tag{14}$$

where: HB = output at 95% of stable high level output;

BL = stable baseline output; and

HL = stable high level output.

Output within 95% of stable baseline output--Fall time for quantitativeoutput detectors was measured from the time the detector was no longer in contact with test product until it reached 95% of its final stable baseline output. The following formula was used to calculate the output within 95% stable baseline level output based on the stable baseline level output and the stable high level output.

$$OB = BL + (HL - BL) \times 0.05$$
 (15)

where: OB = output within 95% of stable baseline output;

BL = stable baseline output; and

HL = stable high level output.

Relative percent difference—The relative percent difference is a measure of variation between two observations, neither of which is considered a reference value. It is their absolute difference divided by their average, expressed as a percentage, and was calculated as follows:

$$RPD = 200 \times [(\nabla_1 - \nabla_2)/(\nabla_1 + \nabla_2)]$$
 (16)

where: RPD = relative percent difference;

 V_1 = larger value; and

 $\nabla_2 = \text{smaller value.}$

<u>Coefficient of variation</u>—The coefficient of variation indicates the relative degree of variation associated with two or more values. It is often reported as "Relative Standard Deviation." The coefficient of variation was calculated as follows:

$$CV (Z) = (S/\overline{X}) \times 100 \tag{17}$$

where: CV = coefficient of variation;

S = standard deviation of n values (n-1 degrees of freedom); and

 \bar{X} = mean of n values.

Accuracy—Calculation for accuracy of liquid-phase detectors depended on the type of output that the detectors produced. Some detectors produced quantitative output from which the thickness of product on water was determined. Other liquid-phase detectors only produced an alarm or other qualitative signal that signified that petroleum product was present.

Quantitative-output liquid-phase detectors—Accuracy for quantitative—output detectors is a function of systematic error (bias) and random error (precision). Accuracy was calculated as the sum of the absolute value of the arithmetic mean of differences, |d|, and the absolute value of the confidence coefficient, cc. Relative accuracy for liquid-phase detectors, which is the accuracy at a given hydrocarbon thickness divided by the hydrocarbon thickness and multiplied by 100, was calculated for each test product at every hydrocarbon thickness.

Relative accuracy—The relative accuracy (RA) of a set of data was calculated as follows:

$$RA = \frac{|\overline{d}| + |cc|}{\nabla_r} \times 100 \tag{18}$$

where: $V_{\perp} = \text{reference}$ (theoretical) value;

 $\frac{1}{d}$ = arithmetic mean of the difference of a data set, Equation 18.1; and

cc = 2.5% error confidence coefficient, Equation 18.2.

<u>Mean difference</u>—The arithmetic mean of the difference (d) of a data set was calculated as follows:

$$\overline{d} = \frac{1}{n} \sum_{i=1}^{n} d_{i}$$
(18.1)

where: d; = measured response - theoretical.

Confidence coefficient—The one-tailed 2.5% confidence coefficient (cc) was calculated as follows:

$$cc = t_{0.975} \cdot \frac{s}{\sqrt{n}}$$
 (18.2)

where:

S = the standard deviation (n-1) of the data set; and $t_{0.975} = 2.5\%$ t value = 2.776 for n=5.

The relative accuracy was calculated for both commercial gasoline and synthetic gasoline at every test product thickness. Relative accuracy values were plotted against hydrocarbon thickness.

Qualitative-output detectors—For qualitative-output detectors, accuracy for a particular test product at a particular thickness was calculated as the number of positive responses expressed as a percentage of the total number of tests at that thickness with that test product, according to the following formula:

$$A_{q} = 100 \times (r_{p}/5)$$
 (19)

where: $A_q = accuracy in percent; and <math>r_p = number of positive responses.$

<u>Bias</u>—Bias is a measure of how much, on the average, the quantitativeoutput detector responded high or low with respect to the true (theoretical) response. Bias was calculated as follows:

$$Bias = 100 \times [(\overline{V}_{o} - V_{r})/V_{r}]$$
 (20)

where:

$$\overline{V}_{o}$$
 = the average observed value, $\frac{1}{5}$ $\sum_{i=1}^{5}$ V_{i} ;

 V_i = the individual response to a test atmosphere; and

Vr = the reference (theoretical) value.

Bias at a particular thickness for a particular test product was the percent difference between the average detector output for a series of tests and the actual thickness of test product. In this calculation, the average detector output reading is used as the observed value (V_0) in Equation 13, and the actual test product thickness is the reference value (V_r). Bias was reported in percent.

Detection time—Detection time was the elapsed time between introduction of the detector probe into the test container and when the detector reached a positive response. The nature of a positive response was dependent on whether the detector had quantitative or qualitative output; however, the calculation for detection time was the same for both types of detectors. Detection time was calculated according to the following formula:

$$DT = T_2 - T_1 \tag{21}$$

where: DT = elapsed detection time;

 T_1 = clock time when liquid was first added to test container; and

T₂ = clock time when detector output went from an inactivated state to an activated state for a qualitative-output detector or from a baseline reading to 95% of stable high level output for a quantitative-output detector.

<u>Fall time</u>—Fall time was the amount of elapsed time between removal of test product from the test container and when the detector reached a negative response. The nature of a negative response was dependent on whether the

detector had quantitative or qualitative output; however, the calculation for detection time was the same for both types of detectors. Fall time was calculated according to the following formula:

$$FT = T_2 - T_1 \tag{22}$$

where: FT = elapsed fall time;

 T_1 = clock time when detector was removed from test container; and T_2 = clock time when detector output went from an activated state to an inactivated state for a qualitative-output detector or from a high level reading to within 95% of stable baseline level output for a quantitative-output detector.

<u>Specificity</u>—Determination of specificity was dependent on whether a detector's output was qualitative or quantitative.

Quantitative-output detectors--Specificity for quantitative-output detectors was defined as the ratio of detector output, or measured thickness, to the actual thickness of hydrocarbon test product expressed as a percentage. The following equation was used to calculate specificity for quantitative-output detectors:

$$S = 100 \times m_{d}/t \tag{23}$$

where:

S = specificity in percent;

m_d = detector's output reading in cm; and

t = hydrocarbon thickness in cm (t = 1.27 cm).

Specificity for qualitative-output devices was reported as "activated" if the detector responded within 24 hours. Otherwise, specificity was reported as "inactivated."

GLOSSARY

Accuracy - The degree of agreement between measured values and the true values. Accuracy in a single measurement reflects both systematic and random error. Accuracy in multiple measurements estimates of the same parameter, i.e., average accuracy, reflects systematic error, or bias.

<u>Activated</u> - Refers to the state of a qualitative detector's response when indicating the presence of hydrocarbons.

<u>Bias</u> - The systematic error inherent in a method. Bias may be positive or negative.

Continuous Petroleum Hydrocarbon Detection Systems (Detectors) - A class of detectors that monitor in a constant, real-time mode, without interruption.

Detection Time - Sum of lag time and rise time.

<u>Fall Time</u> - The elapsed time after the test atmosphere is displaced by ultrahigh-purity air until its output returns to within 95% of its original baseline level or there is no detectable signal output.

Intermittent Petroleum Hydrocarbon Detection Systems (Detectors) - A class of detectors that monitor on a periodic basis. Interruptions may be due to sample transfer, sample analysis, system removal, installation, detection response time, etc. A continuous detector may be operated in an intermittent mode.

<u>Lag Time</u> - The elapsed time from a detector's first contact with the test atmosphere or test liquid to the first detectable signal.

Memory Effects - An effect typically encountered when tests using test atmospheres or test liquids of different concentrations or compounds are performed in succession. The effect is caused by residual hydrocarbon vapors from the previous test significantly interfering with the succeeding test.

Non-activated - Refers to the state of a qualitative output detector's response when indicating that no hydrocarbons are detected.

Petroleum Hydrocarbon Detection System (Detector) - Detection system (probes, lines, control box, readouts, etc.) for hydrocarbon compounds specifically designed or used for vapor-phase petroleum product detection.

<u>Precision</u> - The degree of agreement of repeated measurements of the same parameter. Precision estimates reflect random error and are not affected by bias.

<u>Probe</u> - Component of a detection system that must come into contact with petroleum vapor before the vapor can be detected.

Qualitative Response - A type of detector response that indicates only the presence or absence of hydrocarbons without determining the hydrocarbon concentration.

Quantitative Response - A type of petroleum hydrocarbon leak detection system response that quantifies the concentration of the hydrocarbon present.

Relative Accuracy - The absolute mean difference between a group of measured values and the true value, plus the 2.5% error confidence coefficient, divided by the true value. Relative accuracy is a measure of the maximum (upper 95% limit) expected bias for a series of measurements.

Responses - The detector's indications of the presence or concentration of petroleum hydrocarbons. These can be qualitative or quantitative.

Response Factor - The ratio of the detector response (ppmv, %, etc.) to the test atmosphere or liquid concentration (ppmv, %, etc.).

Response Time - A general term that refers to the more specific terms of lag time and rise time, which together constitute detection time, and fall time.

Rise Time - The elapsed time from a detector's first detectable signal in response to hydrocarbon vapor to an output that is 95% of full scale.

<u>Specificity (gas-phase)</u> - The ability of a detector to detect a particular chemical or class of chemicals at 250 ppm-v.

<u>Specificity (liquid-phase)</u> - The ability of a detector to respond to different substances. For the purposes of this procedure, these substances include commercial gasoline, synthetic gasoline, diesel fuel, jet fuel, n-hexane, xylene(s), and toluene.

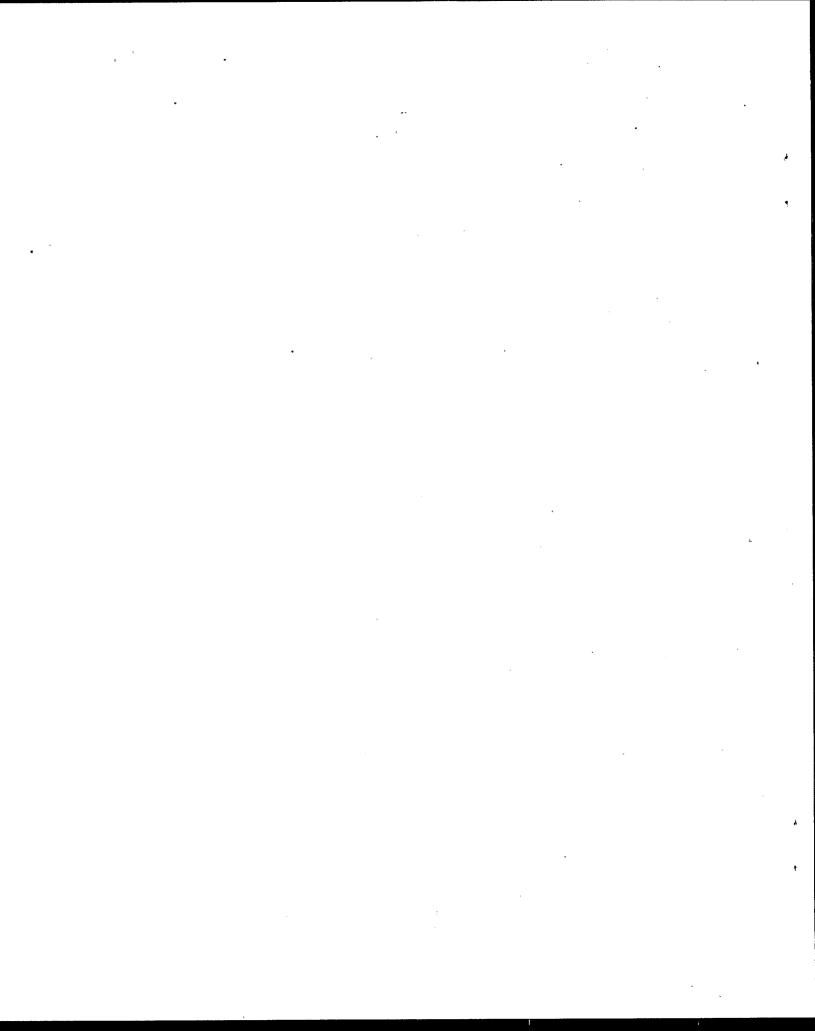
<u>Test Atmospheres</u> - The standard gases (Section 6.0) used to evaluate the performance of detectors.

<u>Test Chamber</u> - The gas-tight or liquid-tight chamber where a detector is exposed to ultrahigh-purity air or water and hydrocarbon test gases or liquids.

<u>Test System</u> - The arrangement of instrumentation and equipment required to perform this test procedure.

For a larger glossary of terms consistent with the field of petroleum and hydrocarbon detection, consult the <u>Survey of Vendors of External Petroleum</u>

Leak Monitoring Devices for <u>Use with Underground Storage Tanks</u> Glossary (2).



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