



Designation: D7112 – 12

Standard Test Method for Determining Stability and Compatibility of Heavy Fuel Oils and Crude Oils by Heavy Fuel Oil Stability Analyzer (Optical Detection)¹

This standard is issued under the fixed designation D7112; the number immediately following the designation indicates the year of original adoption or, in the case of revision, the year of last revision. A number in parentheses indicates the year of last reapproval. A superscript epsilon (ϵ) indicates an editorial change since the last revision or reapproval.

1. Scope*

1.1 This test method covers an automated procedure involving titration and optical detection of precipitated asphaltenes for determining the stability and compatibility parameters of refinery residual streams, residual fuel oils, and crude oils. Stability in this context is the ability to maintain asphaltenes in a peptized or dissolved state and not undergo flocculation or precipitation. Similarly, compatibility relates to the property of mixing two or more oils without precipitation or flocculation of asphaltenes.

1.2 This test method is applicable to residual products from atmospheric and vacuum distillation, from thermal, catalytic, and hydrocracking processes, to products typical of Specifications D396, Grades No. 5L, 5H, and 6, and D2880, Grades No. 3-GT and 4-GT, and to crude oils, providing these products contain 0.05 mass % or greater concentration of asphaltenes.

1.3 This test method is not relevant to oils that contain less than 0.05 % asphaltenes, and would be pointless to apply to unstable oils that already contain flocculated asphaltenes.

1.4 The values stated in SI units are to be regarded as standard. No other units of measurement are included in this standard.

1.5 *This standard does not purport to address all of the safety concerns, if any, associated with its use. It is the responsibility of the user of this standard to establish appropriate safety and health practices and determine the applicability of regulatory limitations prior to use.*

2. Referenced Documents

- 2.1 *ASTM Standards:*²
D396 Specification for Fuel Oils

¹ This test method is under the jurisdiction of ASTM Committee D02 on Petroleum Products, Liquid Fuels, and Lubricants and is the direct responsibility of Subcommittee D02.14 on Stability and Cleanliness of Liquid Fuels.

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² For referenced ASTM standards, visit the ASTM website, www.astm.org, or contact ASTM Customer Service at service@astm.org. For *Annual Book of ASTM Standards* volume information, refer to the standard's Document Summary page on the ASTM website.

- D2880 Specification for Gas Turbine Fuel Oils
D4057 Practice for Manual Sampling of Petroleum and Petroleum Products
D4175 Terminology Relating to Petroleum, Petroleum Products, and Lubricants
D4177 Practice for Automatic Sampling of Petroleum and Petroleum Products
D6299 Practice for Applying Statistical Quality Assurance and Control Charting Techniques to Evaluate Analytical Measurement System Performance
D6560 Test Method for Determination of Asphaltenes (Heptane Insolubles) in Crude Petroleum and Petroleum Products

3. Terminology

3.1 Definitions:

3.1.1 For definitions of some terms used in this test method, such as crude oil, repeatability, reproducibility, and residual fuel oil, refer to Terminology D4175.

3.1.2 *asphaltenes, n*—(rarely used in the singular), *in petroleum technology*, represent an oil fraction that is soluble in a specified aromatic solvent but separates upon addition of an excess of a specified paraffinic solvent.

3.1.2.1 *Discussion*—In this test method, the aromatic solvent is xylene and the paraffinic solvent is n-heptane.

3.1.3 *compatibility, n*—of crude oils and of heavy fuel oils, the ability of two or more crude oils or fuel oils to be blended together within specified ratios without evidence of separation, such as flocculation or separation of asphaltenes.

3.1.4 *flocculation, n*—of asphaltenes in crude oils or heavy fuel oils, the aggregation of colloiddally dispersed asphaltenes into larger, visible masses that may or may not settle.

3.1.5 *stability reserve, n*—of crude oils, heavy fuel oils, and residual streams containing asphaltenes, the property of an oil to maintain asphaltenes in a peptized (colloiddally dispersed) state and prevent their flocculation when stored or when blended with other oils.

3.1.5.1 *Discussion*—An oil with a high stability reserve can be stored for a long period of time or blended with a range of other oils without flocculation of asphaltenes.

*A Summary of Changes section appears at the end of this standard

3.2 Definitions of Terms Specific to This Standard:

3.2.1 *aromatic solvent equivalent (xylene equivalent), SE, n*—the lowest aromatic solvent (xylene) content, expressed as a volume %, in a mixture containing aromatic and paraffinic solvents (xylene and *n*-heptane) which, when mixed with oil, will not result in flocculation of asphaltenes. See *flocculation ratio*.

3.2.1.1 *Discussion*—SE is defined as $FR_{5/1}$ multiplied by 100 %, as shown in Eq 2.

3.2.2 *evaporation correction coefficient, n*—the rate of evaporation of aromatic solvent (xylene) from the sample cup, measured in grams per hour.

3.2.3 *flocculation ratio (FR), n*—the lowest aromatic solvent (xylene) concentration, expressed as a proportion of xylene to xylene plus *n*-heptane which, when mixed with an oil, will not result in flocculation of asphaltenes. See 15.1, Eq 1.

3.2.4 $FR_{5/1}$, *n*—the flocculation ratio at a dilution of 5 mL of xylene and *n*-heptane solvent mixture to 1 g of oil.

3.2.4.1 *Discussion*—The ratio 5 to 1 is used internally by a number of oil companies involved with the stability and compatibility of heavy fuel oils and crude oils. This ratio is chosen so that a *P*-value of six represents an $FR_{5/1}$ of zero.

3.2.5 *insolubility number, I_N, n*—a crude oil blending model parameter which can be used to determine if blends of oils are compatible or incompatible. See *solubility blending number*.

3.2.5.1 *Discussion*—Insolubility numbers for individual oils are determined and calculated from the density of the oil, aromatic solvent equivalent value and volume of paraffinic solvent (*n*-heptane) that can be added to 5 mL of oil without asphaltene precipitation. The equations are given under Calculation of Results (see 15.2).

3.2.6 *maximum flocculation ratio, FR_{max}, n*—of asphaltenes in residual fuel oils and crude oils, the minimum required solvency power of a solvent mixture, expressed as a ratio by volume of aromatic solvent (xylene) to aromatic solvent plus paraffinic solvent (*n*-heptane) to keep the asphaltenes in an oil colloiddally dispersed.

3.2.6.1 *Discussion*— FR_{max} is determined from a plot of flocculation ratios versus the oil concentration in solvent, extrapolated to infinite dilution of the sample at the y-axis (where $(1/X) = 0$. See Eq 3).

3.2.7 *oil matrix, n*—that portion of a sample of heavy fuel oil or crude oil that surrounds and colloiddally disperses the asphaltenes.

3.2.7.1 *Discussion*—For purposes of this test method, an oil sample is considered to be composed of an oil matrix (sometimes called an oil medium) and asphaltenes.

3.2.8 *P-value, n*—of refinery residual steams, residual fuel oils and crude oils, an indication of the stability or available solvency power of an oil with respect to precipitation of asphaltenes.

3.2.8.1 *Discussion*—Since the equation defining *P*-value is $P = (1 + X_{min})$, where X_{min} is the minimum volume of paraffinic solvent, *n*-heptane, (in mL) needed to be added to 1 g of oil to result in flocculation of asphaltenes, the smallest *P*-value is 1, which means the oil is unstable and can precipitate asphaltenes without addition of any paraffinic solvent. A higher *P*-value

indicates that an oil is more stable with respect to flocculation of asphaltenes. *P*-value by this test method relates specifically to xylene and *n*-heptane as the aromatic and paraffinic solvents, respectively.

3.2.9 P_a , *n*—the *P*-value of an asphaltene, which is the peptizability or ability of an asphaltene to remain colloiddally dispersed.

3.2.10 P_o , *n*—the *P*-value of an oil matrix. See *oil matrix*.

3.2.11 *peptize, v*—of an oil or cutter stock, to dissolve an asphaltene or to maintain an asphaltene in colloiddal dispersion.

3.2.12 *solubility blending number, S_{BN}, n*—a crude oil blending model parameter which can be used to determine if blends of oils are incompatible or compatible. See *insolubility number*.

3.2.12.1 *Discussion*—Solubility blending numbers for individual oils are determined and calculated from the density of the oil, aromatic solvent equivalent value, and volume of paraffinic solvent that can be added to 5 mL of oil without asphaltene precipitation. The equations are given under Calculation of Results (see 15.2).

3.2.13 *step size, n*—the volume in mL of each portion of *n*-heptane added to the stock solution in the course of the test procedure.

3.2.14 *stock solution, n*—a solution of a sample dissolved in a specific amount of xylene.

3.3 Symbols:

FR = flocculation ratio

$FR_{5/1}$ = flocculation ratio at a dilution of 5 mL solution (xylene plus *n*-heptane) to 1 g of oil

FR_{max} = maximum flocculation ratio

I_N = insolubility number

P = the *P*-value of an oil

P_a = the *P*-value of an asphaltene

P_o = the *P*-value or peptizing power of an oil matrix

S_{BN} = solubility blending number

SE = xylene equivalent, volume %

X_{min} = *n*-heptane consumption of undiluted oil, in mL/g of oil

4. Summary of Test Method

4.1 Stability and compatibility parameters are determined by titration and optical detection of precipitated asphaltenes. A stock solution is prepared and three different mixtures of the sample oil plus xylene are titrated with *n*-heptane to cause precipitation of asphaltenes. The titrated mixture is continuously circulated through an optical detector which detects precipitated asphaltenes by back-scattering of visible light. The amounts of oil, xylene, and *n*-heptane are used to calculate stability parameters: solvent equivalent, *P*-value, and $FR_{5/1}$. If the density of a crude oil sample is known, then the compatibility parameters (S_{BN} and I_N) of the crude oil may also be calculated.

5. Significance and Use

5.1 Automatic determination of stability parameters using a light back-scattering technique improves accuracy and removes human errors. In manual testing, operators have to

visually compare oil stains on pieces of filter paper to determine if asphaltenes have been precipitated.

5.2 Refinery thermal and hydrocracking processes can be run closer to their severity limits if stability parameters can be calculated more accurately. This gives increased yield and profitability.

5.3 Results from the test method could be used to set a standard specification for stability parameters for fuel oils.

5.4 The compatibility parameters of crude oils can be used in crude oil blending in refineries to determine, in advance, which crude oil blends will be compatible and thus can be used to minimize plugging problems, unit shut downs, and maintenance costs. Determination of crude oil compatibility parameters also enables refineries to select crude oil mixtures more economically.

5.5 This test method can measure stability and compatibility parameters, and determine stability reserve on different blends for particular applications to optimize the blending, storage, and use of heavy fuel oils

NOTE 1—Users of this test method would normally use stability and compatibility parameters to determine stability reserve of residual products, fuel blends and crude oils. However, the interpretation of stability, stability reserve and compatibility is heavily ‘use dependent,’ and is beyond the scope of this test method.

6. Interferences

6.1 Free water present in the oil can cause difficulties with the optical detector and should be removed by centrifuging prior to testing.

6.2 Solid particles, such as coke or wax particles, mud, sand, or catalyst fines, in the oil will not affect the optical detector or interfere with the results.

7. Apparatus

7.1 *PORLA Heavy and Crude Oil Stability and Compatibility Analyzer*^{3,4}—See Figs. 1 and 2.

7.1.1 A portion of the apparatus is shown diagrammatically in Fig. 2 and is comprised of the following parts:

7.1.1.1 *Sample Cup*, light weight, inert cups designed to fit the sample carousel, with a smooth, flat bottom, volume approximately 100 mL. Typically, aluminum cups have been used.

7.1.1.2 *Sample Carousel*, typically a four-position sample cup holder delivering the sample cups sequentially to the measurement position.

³ The PORLA Heavy and Crude Oil Stability and Compatibility Analyzer is covered by Euro patent EP 0737309 and U.S. patent US5715046. Interested parties are invited to submit information regarding the identification of an alternative(s) to this patented item to the ASTM International Headquarters. Your comments will receive careful consideration at a meeting of the responsible technical committee,¹ which you may attend.

⁴ The sole source of supply of the PORLA Heavy and Crude Oil Stability and Compatibility Analyzer known to the committee at this time is Finnish Measurement Systems Limited, Koskikuja 5, FIN-71570 Syvanniemi, Finland, www.finnmeassys.com. If you are aware of alternative suppliers, please provide this information to ASTM International Headquarters. Your comments will receive careful consideration at a meeting of the responsible technical committee,¹ which you may attend.

7.1.1.3 *Mixer Lift System*, vertically moving lift system, forming a seal with the sample cup in the measurement position and incorporating a mechanical stirrer which starts to rotate when the seal is made. It also incorporates delivery lines for *n*-heptane and xylene addition, the circulation line for passing the sample through the detector and the exhaust line, which empties the sample cup after analysis.

7.1.1.4 *Aromatic Solvent Pump*, accurate and adjustable ceramic piston pump, capable of delivering xylene at a rate of 0.01 to 0.5 mL/s.

7.1.1.5 *Paraffinic Solvent Pump*, accurate and adjustable ceramic piston pump, capable of delivering *n*-heptane at a rate of 0.01 to 0.5 mL/s.

7.1.1.6 *Circulation Pump*, accurate and adjustable ceramic piston pump used to circulate the sample under test through the detector system.

7.1.1.7 *Exhaust Pump*, accurate and adjustable ceramic piston pump used to empty the sample cup at the end of the measurement.

7.1.1.8 *Detector System*, (see Fig. 3) optical detector through which the sample solution is continuously circulated. The detector is composed of a visible light source and a photodiode for recording the light reflecting from asphaltene particles in the test sample.

7.1.1.9 *Hot Plate*, a temperature controlled heating system may be located below the sample cups, which will warm up the sample so that the titration may be performed at an elevated temperature. The temperature of the hot plate should be adjustable between 20 and 100°C.

7.1.2 *Computer*, controls the measurement and calibration programs and is an interface between the operator and the analyzer.

7.1.3 *PORLA Step Measurement Screen*, computer display, allowing data about the sample and operator to be input as well as showing the results of each titration (see Fig. 4).

7.1.4 *Parameter Screen*, computer display, allows all of the measurement cycle parameters to be adjusted from the default values and also allows the pump calibration procedure to be run (see Fig. 5).

8. Reagents and Materials

8.1 *Purity of Reagents*—Reagent grade chemicals shall be used in all tests. Unless otherwise indicated, all reagents shall conform to the specifications of the Committee on Analytical Reagents of the American Chemical Society where such specifications are available.⁵ Other grades may be used, provided it is first determined that the reagents are of sufficiently high purity to permit their use without lessening the accuracy of the determination.

8.2 *Xylene (C₈H₁₀)*—The xylene used is generally a mixture of ortho, meta, and para isomers and may contain some ethyl benzene. (**Warning**—Flammable, health hazard.)

⁵ *Reagent Chemicals, American Chemical Society Specifications*, American Chemical Society, Washington, DC. For suggestions on the testing of reagents not listed by the American Chemical Society, see *Analar Standards for Laboratory Chemicals*, BDH Ltd., Poole, Dorset, U.K., and the *United States Pharmacopeia and National Formulary*, U.S. Pharmacopeial Convention, Inc. (USPC), Rockville, MD.



FIG. 1 PORLA Heavy and Crude Oil Stability and Compatibility Analyzer

8.3 *n*-heptane (C_7H_{16})—(Warning—Flammable, health hazard.)

9. Hazards

9.1 Place the analyzer in a fume hood or similar well ventilated area to minimize exposure of operators to harmful vapors.

9.2 Operators should use proper protective laboratory clothing and gloves to avoid skin exposure to oil samples and solvents. In addition, operators should be careful when handling hot oil containers when preparing the stock solutions from very viscous oils as oil spills on exposed skin will cause burns.

10. Sampling and Test Specimens

10.1 Obtain samples in accordance with procedures described in Practices D4057 or D4177. Ensure that samples are representative of the whole batch of oil.

10.2 A minimum sample size of 40 g is required for a single test. It is preferable to collect a larger sample such as 200 to 500 g to allow for multiple testing, if necessary.

10.3 Ensure that the sample is homogeneous before withdrawing an aliquot or test specimen for testing.

10.4 To avoid changes or degradation of oil samples, minimize exposure to air, temperatures above 25°C, and light. Store samples in sealed, opaque containers such as metal cans or dark glass bottles.

11. Preparation of Apparatus

11.1 Review the operations manual for the apparatus,⁶ and follow all recommended steps. The following actions summarize the preparation of the apparatus:

11.1.1 Place the apparatus on a level surface, in a well-ventilated area such as a fume hood.

⁶ Obtain the operator's manual from the supplier of the apparatus.

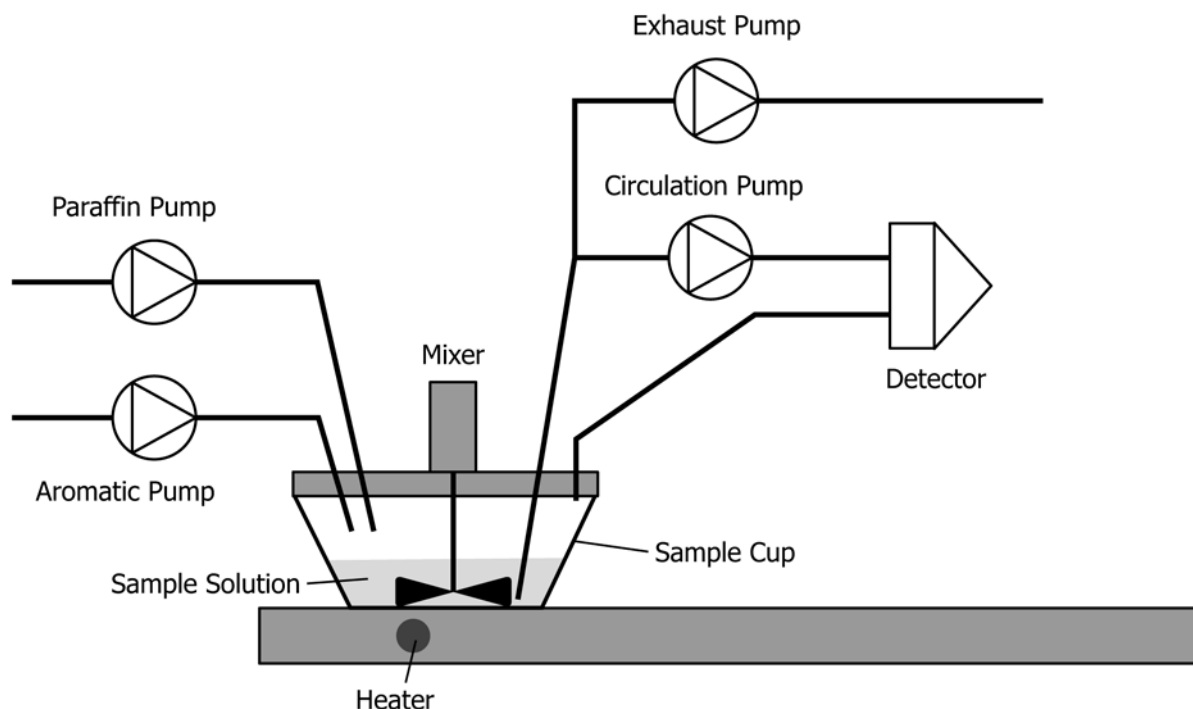


FIG. 2 Schematic Diagram of PORLA Heavy and Crude Oil Stability and Compatibility Analyzer

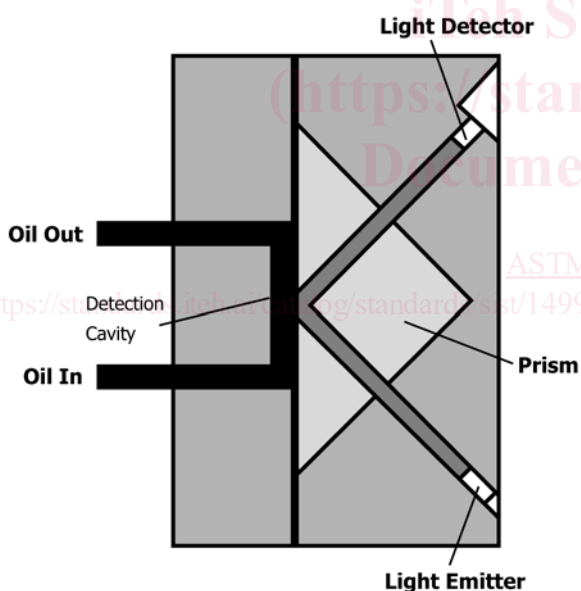


FIG. 3 Schematic Diagram of PORLA Heavy and Crude Oil Stability and Compatibility Analyzer Detector

11.1.2 Connect the analyzer and the computer to a suitable ac power source, and connect the analyzer, keyboard, monitor, and mouse to the computer with the cables supplied.

NOTE 2—All of the controls for the instrument except for the main power switch are accessed by means of the keyboard and mouse of the computer.

11.1.3 Fill the aromatic and paraffinic solvent bottles with xylene and *n*-heptane, respectively, and ensure the waste bottle is empty. Connect the three plastic bottles, labeled aromatic solvent, paraffinic solvent and waste, to the appropriate lines from the analyzer.

11.1.4 Switch the analyzer on using the rocker switch on the back panel. A green lamp will illuminate on the upper right of the front panel.

11.1.5 Switch on the computer and allow it to boot up.

11.1.6 Select either PORLA Step or PORLA Test by double clicking the mouse on the appropriate icon to bring up the PORLA Step measurement screen (see Fig. 4) or the system test screen.

12. Calibration and Standardization

12.1 *Pump Calibration*—Perform the pump calibration at the initial set-up and whenever the instrument has been serviced. When in continuous service, perform pump calibrations monthly to verify the pumping rates.

12.1.1 On the PORLA Step measurement screen (Fig. 4) select the Parameter screen (see Fig. 5). Calibrate the xylene and *n*-heptane pumps by starting the automatic procedure under Pump Calibration.

12.1.2 Identify two clean sample cups as ‘xylene’ and ‘*n*-heptane,’ and accurately weigh to two decimal places. These are the tare weights of the cups.

12.1.3 The instrument automatically pumps solvent over a constant period of time (60 s) into a tared sample cup.

12.1.4 Weigh the collected solvent and sample cup and subtract the tare weight of the sample cup to give the weight of solvent pumped in the set time period (60 s).

12.1.5 Enter the weight of solvent collected in the appropriate screen.

12.1.5.1 For the xylene pump rate, the instrument calculates the updated pump rate in g/s and displays it in the appropriate field in the parameter screen.

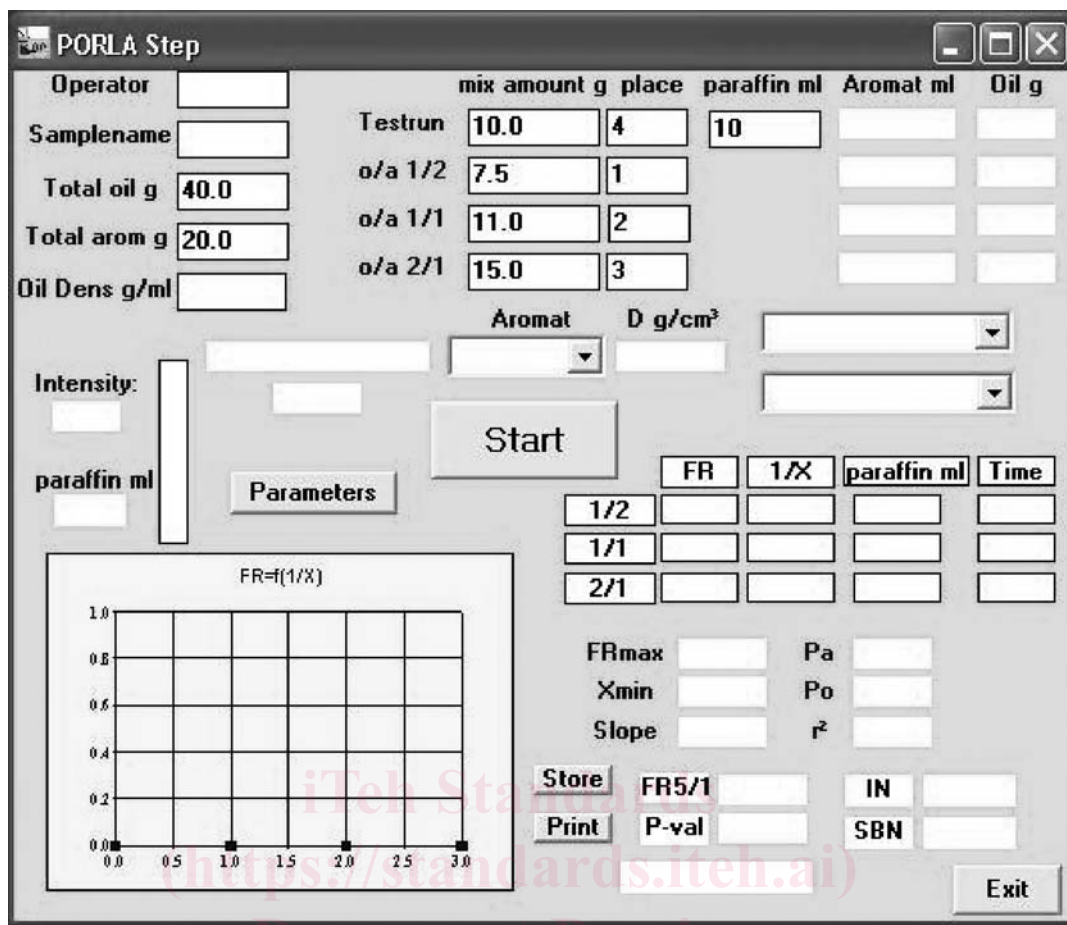


FIG. 4 PORLA Step Measurement Screen

12.1.5.2 For the *n*-heptane pump rate, the instrument calculates the updated pump rate and displays it in mL/s in the appropriate field in the parameter screen.

12.2 *Evaporation Correction Coefficient*—Determine the evaporation correction coefficient for the xylene by weighing (accurate to two decimal places) a sample cup containing a stock solution after 0.5, 1.0, and 1.5 h of standing in the sample carousel and calculate the average solvent loss by evaporation in g/h. Input this information into the PORLA parameter screen, which is accessed by means of the Parameter button in the PORLA Step measurement screen.

12.2.1 Verify the evaporation correction coefficient quarterly when the instrument is in continuous service.

12.2.2 If a specific sample contains volatile components, such as a whole crude oil, use the semi-automatic *Light Ends* mode of operation. In this mode of operation, evaporation is minimized by loading the stock solution cups one at a time into the carousel. See 14.9.

NOTE 3—No significant evaporation of solvent or sample occurs during the titration procedure because the sample cup is sealed in the test position in the instrument. However, evaporation of xylene does occur when several sample cups sit in the sample carousel during the fully automatic mode of operation.

13. Quality Control Monitoring

13.1 Confirm the performance of the instrument and test procedure by analyzing quality control (QC) samples.

13.1.1 The QC sample(s) should be a stable and homogeneous residual fuel oil or crude oil similar in composition and viscosity to the type of samples routinely tested, and containing from 1 to 10 mass % asphaltene. Asphaltene content may be measured by Test Method D6560.

13.2 Prior to monitoring the measurement process, determine the average value and control limits for the QC sample (see Practice D6299 and Section 3 of Manual 7⁷).

13.3 Record QC results and analyze by control charts or other statistically equivalent techniques to ascertain the statistical control status of the total test process.^{7,8} Investigate any out-of-control data for root cause(s). The results of this investigation may, but not necessarily, result in instrument recalibration.

13.4 The frequency of QC testing is dependent on the criticality of the quality being measured, the demonstrated stability of the testing process and customer requirements.⁸ Generally, a QC sample should be analyzed each testing day for routine samples. The QC frequency should be increased if a large number of samples are routinely analyzed. However,

⁷ ASTM MNL 7, *Manual on Presentation of Data Control Chart Analysis*, 6th edition, Section 3: Control Chart for Individuals. Available from ASTM International Headquarters

⁸ In the absence of explicit requirements given in the test method, this clause provides guidance on QC testing frequency.