NISTIR 7383

Selected Procedures for Volumetric Calibrations (2013 Ed)

Harris, Georgia L.

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Office of Weights and Measures
Physical Measurement Laboratory

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U.S. Department of Commerce *Rebecca Blank, Acting Secretary*

National Institute of Standards and Technology

Patrick D. Gallagher, Under Secretary of Commerce for Standards and Technology and Director

Foreword

This NIST IR of Selected Publications was updated from the 2006 version in 2012 and included Good Laboratory Practices, Good Measurement Practices, and Standard Operating Procedures for volumetric calibrations. Procedures were reviewed at 2012 Regional Measurement Assurance Program meetings with input provided by working metrologists who have implemented these procedures in their laboratories. As a result, additional updates were required in 2013.

Updates to this edition include the latest CIPM accepted equations for the calculation of water density in GLP 10. SOP 17 and 20 have been modified so that they apply to control charts for check standards, standard deviation charts, and range charts for any process with replicate measurements (versus only those for volume transfer calibrations as previously written.)

The following Practices and Procedures that were included in the 2006 edition were *removed* in the 2012 publication:

Standard Operating Procedures for:

• Gravimetric Volume Calibrations Using a Single-Pan Balance (13) and Using Equal Arm Balances (15). Contents specific to glassware calibrations have been included in SOP 14. Few laboratories maintain mechanical balances or equal arm balances for gravimetric calibrations, therefore, these procedures will no longer be updated.

The calibration procedures for Small Volume Provers (SOP 26, Volumetric, and SOP-SVP, Gravimetric) have been revised and are replacing SOP 26 for generic "Gravimetric Calibrations of Dynamic Volumetric Standards" which includes Small Volume Provers and alternative standards in this 2013 edition.

Many of these procedures are updates to procedures that were originally published in NBS Handbook 145, Handbook for the Quality Assurance of Metrological Measurements, in 1986, by Henry V. Oppermann and John K. Taylor. The 2006 updates incorporated many of the requirements noted for procedures in ISO Guide 25, ANSI/NCSL Z 540-1-1994, and ISO/IEC 17025 laboratory [quality] management systems. The 2006 changes incorporated 1) uncertainty analyses that comply with current international methods in the Guide to the Expression of Uncertainty in Measurements (GUM) and 2) measurement assurance techniques using check standards. The 2012 updates included updated terminology consistent with the International Vocabulary of Metrology (VIM), primarily associated with metrological traceability. The 2013 update includes data sheets and example uncertainty budget tables in all SOPs.

Note regarding SI units: This publication conforms to the concept of primary use of the International System of Units (SI) recommended in the Omnibus Trade and Competitiveness Act of 1988 by citing SI units before customary units where both units appear together and placing separate sections containing requirements for SI units before

corresponding sections containing requirements for customary units. In some cases, however, laboratory standards and/or trade practice are currently restricted to the use of customary units (e.g., retail sale of gasoline in the United States is still sold by U.S. gallons); therefore, some procedures in this publication specify only customary units. Non-SI units are predominately in common use in State legal metrology laboratories, and/or the petroleum industry for many volumetric measurements, therefore non-SI units have been used to reflect the practical needs of the laboratories performing these measurements as appropriate. SI units are used where practical, and where use or potential conversion errors, will not likely impact the quality of laboratory measurement results.

Endorsement Disclaimer

Certain commercial equipment, instruments, or materials are identified in this publication in order to specify the experimental procedure adequately. Such identification is not intended to imply recommendation or endorsement by the National Institute of Standards and Technology, nor is it intended to imply that the materials or equipment identified are necessarily the best available for the purpose.

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Good Laboratory Practices



GLP 10

Good Laboratory Practice for the Purity of Water

Water is used in two ways in the metrology laboratory. It may be used as a cleaning fluid. It may also be used as a transfer medium in volume transfer calibrations, or as standard reference material for gravimetric volume calibrations. In each case of these cases, the water must be clean; in the case of gravimetric volume calibrations it must be pure, as well.

Dirty water can cause a number of problems, including depositing residues in a volumetric vessel that could cause volumetric errors or could soil its interior. When in doubt of the cleanliness of the water supply, filters should be attached to the supply lines used.

Cleanliness of water may be achieved by removal of physical contaminating substances, especially by filtration. City water is ordinarily clean but may become dirty from the distribution system and especially from prolonged standing in some kinds of pipes and tubing. Hoses used to transfer water from and into large vessels and tanks may need internal cleaning, as well. Flushing to remove visible contamination is all that is usually required prior to use in volume transfer calibrations or prior to further purification.

Clean Water used in Volume Transfer Calibrations

Clean water is all that is necessary when making measurements by volumetric transfer since volumetric comparisons are involved. When the water density value is used in a volume transfer equation, the ratio of the density values of clean water versus pure water is essentially unchanged, despite impurities. The coefficient of expansion for distilled water and pure water are also essentially the same.

Note: the equations used in gravimetric calibrations do not use ratios and systematic errors will result in the calibration relative to impurities in the water. The use of low resolution density meters for evaluating the quality of clean water (instead of using pure water as described) is discouraged for use in gravimetric calibrations.

Pure Water used in Gravimetric Volume Calibrations

Pure water is needed for gravimetric volume calibrations. For gravimetric calibrations, filtration systems alone are *not* adequate. Pure water requires the removal of chemical contaminants and this may be achieved by distillation, reverse osmosis, ion exchange systems, or combinations of these systems. The purity of the water from any given system and maintenance requirement is often dependent on the quality of the source water. Source water should be tested to determine the best type of system to meet laboratory needs. Density calculations may be accurate even when appreciable levels of dissolved salts remain in the water. However, since it is very difficult to know what type and quantity of salts are tolerable before density is affected, it is prudent to use the best system practical within budgetary constraints.

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Water Quality Specifications and Purification Methods

ASTM Type III or IV Reagent Water¹* is recommended as adequate for gravimetric calibration purposes. Such water may be produced by distillation or by ion exchange with relatively inexpensive equipment. Type III grade of reagent water may be prepared by distillation, ion exchange, continuous electrodeionization reverse osmosis, or a combination thereof, followed by polishing with a 0.45 µm membrane filter. Type IV grade of reagent water may be prepared by distillation, ion exchange, continuous electrodeionization reverse osmosis, electrodialysis, or a combination thereof.

The ASTM D-1193 specifications for conductivity and resistivity are noted in Table 1.

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Table I	Conductivity	y and recict	ivitv sneciti	cations for water.
I abic I.	Community	dia i coist	ivity specifi	cations for water.

	Type I	Type II	Type III	Type IV
Electrical conductivity, max, μS/cm at 298 K (25 °C)	0.056	1.0	0.25	5.0
Electrical resistivity, min, MΩ·cm at 298 K (25 °C)	18	1.0	4.0	0.20

Conductivity or resistivity (along with other water quality measurements) are often used to assess the water quality used in cooling towers, boilers, relative humidity systems, micro and nanoelectronic systems and in pharmaceuticals, to ensure water of sufficient purity and to minimize corrosion or build-up within such systems.

Conductivity and resistivity measurements do not have a direct correlation to water density, which is the critical attribute of concern for gravimetric volume calibrations. However, conductivity or resistivity measurements are a good indicator of water quality and whether the system is in good operating condition or needs service. Conductivity is simply the reciprocal of resistivity. For water quality specifications and assessment purposes, conductivity is usually measured in microSiemens per centimeter (μ S/cm) and resistivity is usually measured in megaohms-centimeter (μ C-cm), both usually at a reference temperature of 25 °C. Conductivity is greatly influenced by temperature and is not linear. However, this is not a major concern for typical laboratory applications requiring pure water. Either a meter or indicator light should be included in laboratory systems to monitor water quality output. Standard reference materials are available to test conductivity and resistivity units. But, because the measurement values are not used to perform corrections to volume calibrations, traceable calibrations of the units are not essential.

Most manufacturers of systems at this level indicate that the resulting water will meet Type I specifications. Sales literature will usually specify whether the equipment will provide water of the above quality. Additionally, many manufacturers will provide a source water quality test and recommend a system to meet the purity and volume requirements of the laboratory.

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¹ ASTM D 1193 - (06)2011, Standard Specification for REAGENT WATER, ASTM, http://www.astm.org, 1916 Race St., Philadelphia, PA 19103.

A cartridge-type ion-exchange system is recommended for its simplicity and ease of operation. It can operate intermittently (on demand) and requires little or no maintenance except for change of cartridges, the need for which will be indicated. A relatively small system (2 L/hr to 30 L/hr) is adequate for laboratories calibrating glassware and test measures up to and including 20 L (5 gal) standards. It may be used on demand or to fill a small (20 L to 40 L) storage bottle to assure a continuous supply of calibration water.

There are two broad types of ion-exchange systems. Pressure cartridge systems (PCS) operate directly from line pressure (up to 700 kPa) and need no special operation precautions. The less expensive type operates from the water line through a needle valve to produce a specified flow through the cartridge(s). In this system, the outlet must not become blocked or turned off to prevent the water pressure from building up and bursting the cartridge. It is common practice to plumb directly from the output of this cartridge to a storage tank without using a valve in between. The unit is operated by simply turning the shut-off valve located at the water supply.

Water Density Measurements

Water density may be measured with a five- or six-place oscillation-type density meter calibrated using suitable standard reference materials that are representative of the range of use. These systems typically measure the density at a specific reference temperature (generally 20 °C). However, density meters are generally not needed if an appropriate water purification system is used that includes a way to measure conductivity or resistivity and where outside validation of water quality has been obtained.

One method that may be used in the laboratory for monitoring water over time is to maintain a quality assurance reference flask (QARF) that is calibrated at any time a gravimetric calibration is performed. If the results of a proficiency test (interlaboratory comparison) that is conducted at the same time are found acceptable, one can assume that the quality of the water was acceptable on that day. In this way, the reference value for the flask may be monitored over time. Glassware is unlikely to change with use of pure water; thus, significant changes on a control chart for a QARF indicate other problems that may need to be investigated such as water quality or other changes in the calibration system or process.

Exposure of pure water in storage to air will likely cause degradation in the conductivity and/or resistivity measurements. However, pure water has been stored for over a year with little degradation in the density quality (provided that storage containers and lines are clean and that there is no bacterial growth, algae, or other contamination).

Temperature equilibrium is a critical factor in the density stability of water. This is especially important for large volumes. Water temperatures must be stable throughout the container. Temperature accuracy is as important as purity for a correct density determination. If water is coming straight from the tap through the purification system into the prover, the temperature may fluctuate appreciably. Therefore, it is important to store an adequate volume of water to complete a calibration either already purified or ready to go through the system.

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Water Density Calculations

Water density tables (see Table 9.8 in NISTIR 6969) or calculations are required for most gravimetric calculations. For use in computer programs (most often spreadsheets), the use of calculations are preferred to the use of look-up tables.

The following equation provides the density for air-free water and is recommended for use between 0 °C and 40 °C.² This equation provides results in kg/m³. Therefore, divide by 1000 to convert the value to g/cm³.

$$\rho(t_{W}) = a_{5} \left[1 - \frac{\left((t + a_{1})^{2} (t + a_{2}) \right)}{a_{3} (t + a_{4})} \right]$$
 Eqn. 1

where:

 $a_1 = -3.983~035~^{\circ}\text{C}$

 $a_2 = 301.797 \, ^{\circ}\text{C}$

 $a_3 = 522528.9 \, ^{\circ}\text{C}^2$

 $a_4 = 69.348 81 \, ^{\circ}\text{C}$

 $a_5 = 999.974 950 \text{ kg/m}^3$

 t_w is the temperature of the water in °C.

In Excel Format:

 $=999.97495*(1-(((G8-3.983035)^2*(G8+301.797))/(522528.9*(G8+69.34881))))$

where, G8 is the cell with the Celsius temperature and units are given in kg/m³.

Pure water that is used in the laboratory is generally air-saturated, so the density must be corrected. The impact is approximately between 1 and 4 parts per million (or several times the uncertainty of the water density calculation). To adjust the air-free water density in Equation 1 between 0 °C and 40 °C to air-saturated water (the standard laboratory condition), use the following equation,

$$\Delta \rho / kg \ m^{-3} = s_0 + s_1 t$$
 Eqn. 2

where,

 $S_0/(10^{-3} \text{ kg m}^{-3}) = ^-4.612$ and $S_1/(10^{-3} \text{ kg m}^{-3} \circ \text{C}^{-1}) = 0.106$.

In Excel Format:

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² M. Tanaka, G. Girard, R. Davis, A. Peuto, and N. Bignell, Recommended table for the density of water between 0 °C and 40 °C based on recent experimental reports, Metrologia, 38, 301-309 (2001).

$$=(-4.612+0.106*G8)$$

where, G8 is the cell with the Celsius temperature. (Add this value to the density. Air-saturated water density will be less than air-free water density.)

Note: The units in Equation 2 are 10^{-3} kg/m³ or "parts per million," so divide by 1 000 000 to find the change to the water density in g/cm³. Equation 1 and Equation 2 may be combined, taking care to match units.

Uncertainty in Water Density Values

The uncertainty in water density given by the references for the equation itself may be calculated using Eqn. 3. The calculated uncertainty for all of the values in the example data shown below is 0.000 000 83 g/cm³.

$$u(\rho_w) = 0.8394 - 0.00128 \times t + 0.000110 \times t^2 - 0.00000609 \times t^3 + 0.000000116 \times t^4 (10^{-3} kg/m^3)$$

Eqn. 3

Example Data

Temperature (°C)	Air-Free Water Density (g/cm³)	Correction for Air Saturation (g/cm³)	Air-Saturated Water Density (g/cm³)
10	0.999 702 70	-0.000 003 55	0.999 699 15
15	0.999 102 57	-0.000 003 02	0.999 099 55
20	0.998 206 75	-0.000 002 49	0.998 204 26
25	0.997 047 02	-0.000 001 96	0.997 045 06

Additional References:

ASTM D 1125 – (95)2009, Standard Test Method for Electrical Conductivity and Resistivity of Water, ASTM, http://www.astm.org, 1916 Race St., Philadelphia, PA 19103.

ISO 15212-1:2002, Oscillation-type density meters – Part 1: Laboratory instruments, ISO, 2002.

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GLP 13

Good Laboratory Practice for Drying "To Contain" Volume Standards

Vessels calibrated "to contain" must be dried of all measurable water in order to obtain an "empty" weight. The drying process should not contaminate the container; otherwise, it will need to be re-cleaned before further calibration or volumetric use. The following is presented as guidance when drying is required.

Drain as much of the residual water as practical before starting any drying process. If time is not a consideration, a glass tube may be inserted into the container to blow clean dry air (or nitrogen) through it to evaporate the residual water film. A filter or dust trap may be necessary to pre-clean the air used. Air lines must be selected to ensure that aging and wear do not introduce contaminants. Air may be drawn through a tube connected to a vacuum pump, but with some danger of drawing in dirty air from the surroundings. Clean absorbent cotton placed at the neck opening can minimize the entrance of foreign matter. Compressed air systems may introduce finely atomized oils or moisture into the air which may not be visible. If laboratory quality air is not available, nitrogen may be used.

Safety Note: Material Safety Data Sheets (MSDSs) must be available in the laboratory and should be reviewed to ensure safe handling and disposal of all chemicals noted in this procedure. Care must be taken to avoid mixing chemicals such as acetone and alcohols in the laboratory.

Cleaning with solvents may be performed; ethanol is the preferred medium. Preliminary rinsing with acetone will remove large amounts of water, with which it is infinitely miscible, but this solvent often contains impurities such as traces of oils that could deposit on the container walls and be a disposal problem. Thus, a final alcohol (preferably ethanol or methanol) rinse is recommended, even if acetone is used to remove most of the water. Care must be taken to ensure that the alcohol is not mixed with acetone, and that the alcohol is not denatured with oils to an extent that will leave residue on the standard. The alcohol is allowed to drain as much as possible, followed by air drying as described above.

Some metal containers may be internally coated to minimize corrosion. When present, it should be ascertained that such coatings are not affected by alcohol or acetone; otherwise, solvent treatments should not be used.

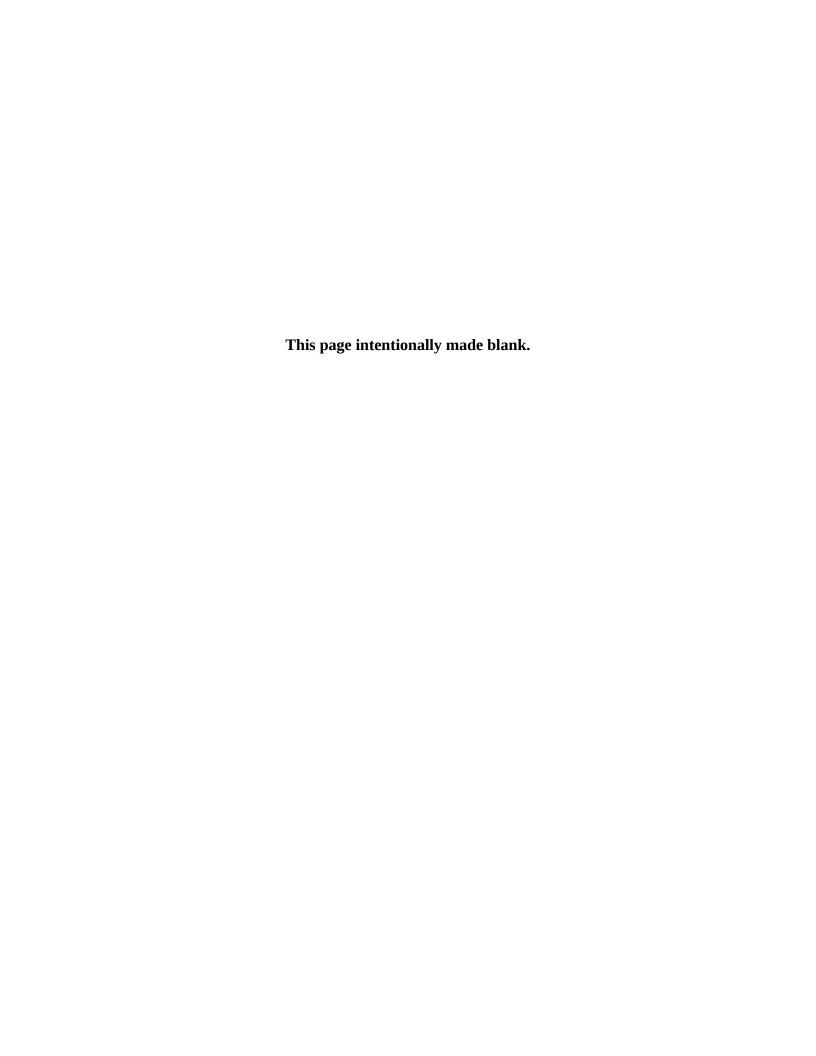
The external surfaces of all containers should be clean when gravimetric calibrations are performed. Otherwise, any removal of external dust or dirt during the measurement process could cause errors of unknown magnitude.

Analytical glassware should not be dried by heating in an oven as the glass may suffer non-elastic expansion and put the glassware out of calibration.

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Calibration Tip: Cover a vessel after it is clean and dry to minimize contaminants from collecting inside the vessel. Clean and dry the flask one to two days prior to the calibration and allow the flask to come to equilibrium with the environment. Obtain an initial baseline "dry weight" of the clean container for use as a dry reference weight on subsequent weighing and drying cycles.

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Good Measurement Practices



GMP 3

Good Measurement Practice for Method of Reading a Meniscus Using Water or Other Wetting Liquid

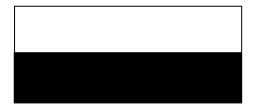
The meniscus of most liquids used in volumetric calibrations is concave-up, with the lowest point in the center that is used to determine the meniscus reading. The curvature of a meniscus is related to the surface tension of the liquid and inversely related to the diameter of the tubing in which it is formed. When reading any meniscus, it is important to ascertain that it is in an equilibrium position. Tapping of sight glasses and/or small motions of containers may be used to induce slight displacements of the meniscus. Return to the same reading is evidence of a stable meniscus.

The meniscus formed by a non-wetting liquid, such as mercury (Hg), is convex with the highest point in the center and is no longer regularly used for volumetric calibrations. In the case of a convex meniscus, the highest point is used to make the reading. The reading of a mercury barometer or thermometer is a classic example of this kind.

In all apparatus where the volume is defined by a concave-up meniscus, the setting or reading is made on the lowest point of the meniscus. In order that the lowest point may be observed, it is necessary to place a shade of some dark material immediately below the meniscus, which renders the profile of the meniscus dark and clearly visible against a light background. A convenient device for this purpose is a collar-shaped section of thick black rubber tubing, cut open at one side and of such size as to clasp the tube firmly.

Two common methods are used for setting and visually reading a meniscus: Options A and B. The method used in calibration should be consistent with the intended use of the volumetric standard. For interlaboratory comparisons, the method to be used should be defined during the planning stages of the comparison. The width of the graduation on the neck scale will affect the readability of the meniscus and should be estimated to the nearest 1/10 of a division when using either Option A or Option B.

Several types of meniscus readers are available. These include black/white meniscus card readers, magnifying glasses with cross-hairs, and collar shaped tubes that may be placed around the graduated neck of the volumetric container. Black/ white meniscus card readers are simple to create as shown in the figure below.



This type of meniscus reader is generally preferred for the Option A method of reading the

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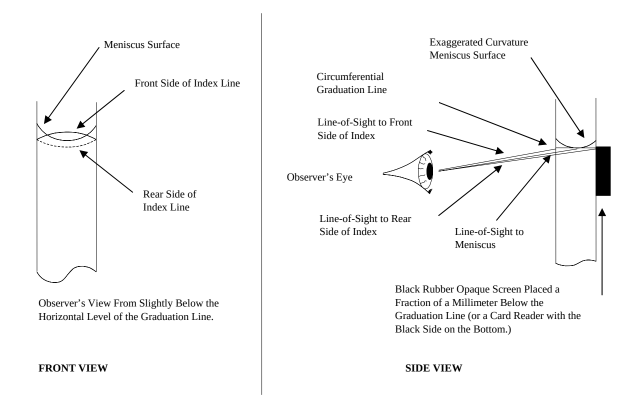
meniscus. More elaborate card readers may be purchased that allow placement around the neck of a flask.

Another type of meniscus reader consists of a clear lens (plastic or glass) with etched lines on the front and back that are aligned to prevent parallax errors in reading. This type may or may not include magnification. This type of reader is preferred for Option B.

Option A

Option A is suitable when graduation lines extend more than 75 percent of the circumference of the sight gage area, for example with graduated neck type flasks or single mark flasks.

The position of the lowest point of the meniscus with reference to the graduation line is such that it is in the plane of the middle of the graduation line. This position of the meniscus is obtained by making the setting in the center of the ellipse formed by the graduation line on the front and the back of the tube as observed by having the eye slightly below the plane of the graduation line. This is illustrated below. The setting is accurate if, as the eye is raised and the ellipse narrows, the lowest point of the meniscus remains midway between the front and rear portions of the graduation line. By this method it is possible to observe the approach of the meniscus from either above or below the line to its proper setting.

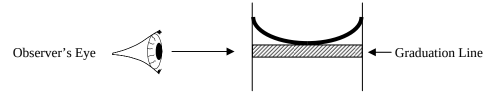


Option B

Option B, is typically used with opaque liquids or when the graduation mark does not extend around the circumference of the volumetric standard. (Option B is also referenced in a number of ISO and OIML standards for glassware, thus it is important to review meniscus reading practices with the user.)

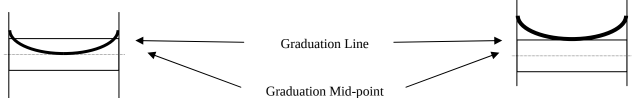
In setting the meniscus or taking a reading, the observer's eye should be normal to and in the same horizontal plane as the meniscus. The illumination is adjusted to get a sharp definition of the meniscus. Elimination of parallax error is very important and can be judged by slight fluctuations of eye level that do not affect the reading.

The position of the lowest point of the meniscus with reference to the graduation line is horizontally tangent to the plane of the upper edge of the graduation line. The position of the meniscus is obtained by having the eye in the same plane of the upper edge of the graduation line as shown below. Offsets from reading in the same plane will produce parallax errors.



Comparison of Option A and Option B Methods

For most practical applications the difference between these two methods is insignificant compared to the tolerances of the volumetric standards. However, a component for measurement uncertainty should be included as appropriate. When performing calibrations, using the glassware as precision standards with clear liquids, or when comparing results among laboratories the difference in meniscus setting is directly related to the visible thickness of the meniscus and the volume of liquid contained in the neck between the top of a graduation line and the middle of the graduation line as shown in the following diagram as 1) which results with Option B when clear liquids are used and 2) which results with Option A and is only possible when clear liquids are used.



Option A, Mid-point of the meniscus.

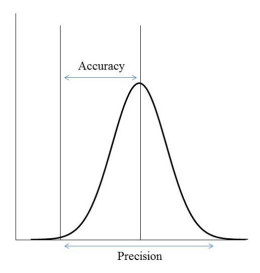
Option B, Top of the meniscus.

When opaque liquids are used, an additional error or correction factor based on the volume contained between the upper and lower edges of the meniscus may need to be considered. The curve of the meniscus and impact of this error is dependent on the diameter of the meniscus and surface tension of the liquid being used.

Uncertainty and Error Analysis in Meniscus Readings

The standard deviation of replicate meniscus readings of volumetric measures provides an estimate of precision that can be used in the uncertainty analysis. However, potential errors in reading a meniscus may be caused by poor eye alignment according to Option A or B, by lack of cleanliness on material surfaces, by variations in visual acuity, by lighting, by the use of cameras or other optical tools, and by the thickness of the graduation line and characteristics of the meniscus itself. Repeatability, or precision, provides an estimate of the process variability, but the potential errors may result in systematic inaccuracies or biases.

"Precision" in the following graphic illustrates the repeatability of the measurement process and "Accuracy" demonstrates the possible systematic inaccuracy of the mean value versus a reference value, due to uncorrected errors.



The sensitivity of the volume measurement to the meniscus reading uncertainty may be calculated using the equation for the volume of a cylinder and the neck diameter.

 $Vol = \pi r^2 h$ where r is the radius of the internal diameter and h is the height of the line or meniscus reading uncertainty. Units will be determined based on the dimensional units that are used. For example radius in cm² and height in cm, will provide results in cm³ or mL. The meniscus itself often has an observed "thickness" that is larger than the graduation lines so this should be considered as well. The potential error in meniscus reading may be treated as a triangular distribution.

Example for a Glass Flask

If the internal diameter of a flask is 1.5 cm (15 mm) and the height of the graduation line is 0.50 mm, the volume associated with the line can be calculated as follows: $Vol = \pi \left(\frac{15}{2}\right)^2 0.5 = 88.36 \text{ mm}^3 (0.088 \text{ mL})$. These values are rounded for illustrative purposes only; significance must be

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evaluated based on measuring instruments and flask resolution. Experimental data obtained by reading a meniscus among multiple laboratory staff members may also be used to incorporate an estimate of uncertainty in meniscus reading.

Example for a Test Measure or Prover

In a test measure or prover, the volume inside the gauge also represents the volume inside the neck. So, two volumes can be calculated and added together to determine the impact.

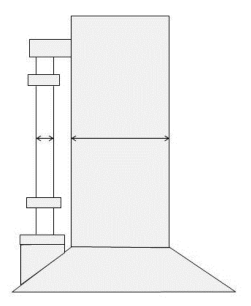
If the internal diameter of the gauge tube is 13 mm (0.5 in) and the height of the graduation line is 0.50 mm, the volume associated with the line inside the gauge tube can be calculated as follows:

$$Vol = \pi \left(\frac{13}{2}\right)^2 0.5 = 66.4 \text{ mm}^3 (0.066 \text{ mL}).$$
 This

amount alone is generally not readable on the test measure or prover. But, the equivalent amount of volume that this gauge tube volume represents in the neck (given a 3 in neck) can be calculated as

follows:
$$Vol = \pi \left(\frac{76.2}{2}\right)^2 0.5 = 2.28 \text{ mL}.$$
 These

two volumes are added together, giving a sum of 2.346 mL, or about 0.143 in³. In the case of a 5 gal test measure, the typical calibration uncertainty is about 0.35 in³, so this potential error is significant.



These values are rounded for illustrative purposes only; actual significance must be evaluated based on measuring instruments and resolutions. Experimental data obtained by reading a meniscus among multiple laboratory staff members may also be used to incorporate an estimate of uncertainty in meniscus reading.

GMP 6

Good Measurement Practice for Cleaning Metal Volumetric Measures

All volumetric test measures, including laboratory standards and those submitted for calibration must be clean at the time of measurement. Drainage is altered by greasy and oily surfaces. Corroded surfaces raise questions that impair future use, so corrosion-damaged containers may not be worth testing. Maintaining clean volumetric test measures improves reproducibility of delivered volume between calibration and usage.

Many laboratories will not accept test measures and provers for calibration unless they have been suitably cleaned to remove contents such as petroleum or other chemical residues. In those cases, all surfaces (inside and out), drain lines, pumps, and hoses that may be used during the calibration process must be cleaned so that no petroleum or other product residue remains that would otherwise spill onto the laboratory floor or drain into the sewer/waste-water system.

Metal measuring vessels are best cleaned by using a **non-foaming** dishwashing detergent. Warm water is most effective both for cleaning and rinsing, which must be thorough. If warm water cleaning and rinsing is used, the vessel must be equilibrated to calibration temperatures before use or calibration.

The cleaning operation involves the use of a liberal amount of the detergent solution and vigorous shaking, swirling, or other motion so that the solution contacts the entire inner surface of the vessel. Depending on the condition of the surface, this operation should be repeated one or more times to ensure removal of oily films and residues. As much as possible of the detergent solution should be drained prior to the water rinses to facilitate complete removal of the detergent from the prover. Ordinarily, three rinses will be sufficient, but this should be confirmed by testing the final rinse for any visual evidence of detergent or other residues.

When detergents are not sufficient to remove oily or greasy deposits, solvents may be used, provided it is ascertained that they do not affect any coatings on the container. The surfaces should be dry before any calibrations are attempted.

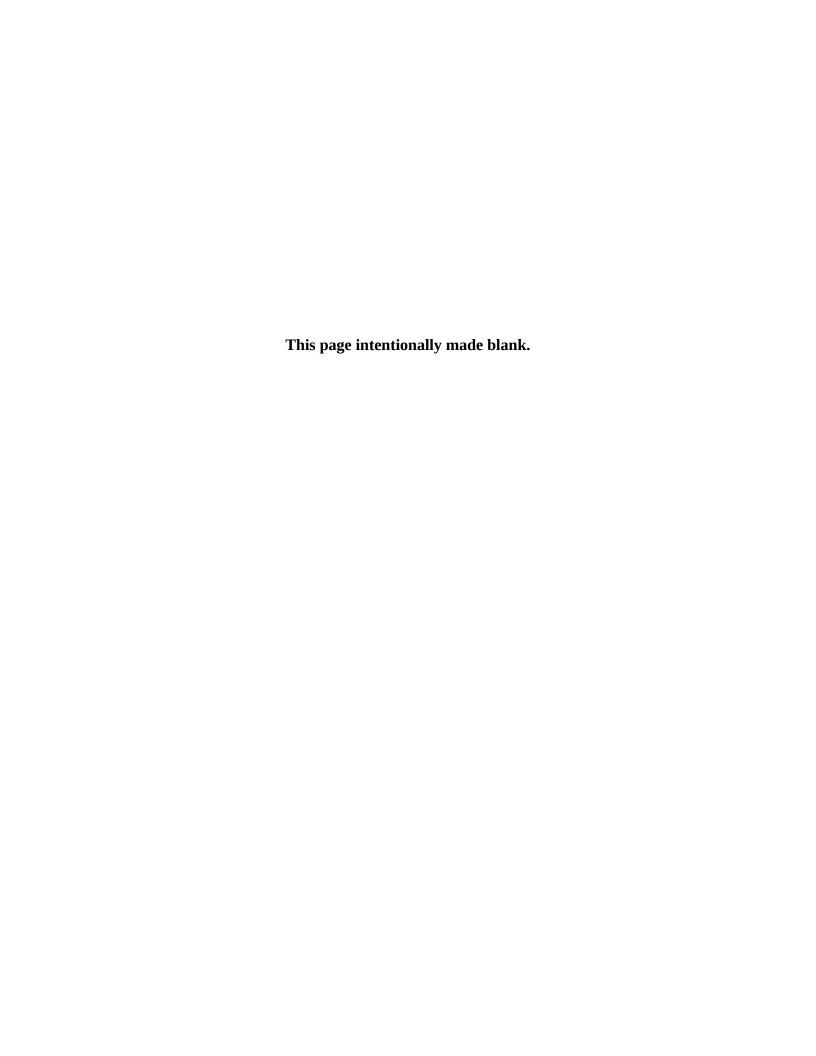
If lime or scale build-up has occurred (usually only in standards that are primarily used with water), they may be cleaned with a suitable lime or scale remover. Many of these chemicals contain acids and may react with metals, so appropriate care should be taken and these chemicals should not be allowed to soak for extended periods of time. Many lime and scale removers have safety and handling requirements and may have special disposal requirements. The material safety data sheet (MSDS) should be reviewed prior to using or disposing of scale and lime removers.

After cleaning or use, vessels should be drained completely and stored in a dry place. They should be covered to prevent entrance of dust and foreign matter. Metal containers further should be protected from damage by denting and/or dropping. Once such a vessel has been visibly dented, it

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is difficult to ascertain whether additional changes have occurred. Even when dents have been removed and the vessel has been calibrated, visual deformities can compromise future use. Accordingly, protection from damage while stored and when in use should be a major concern of the metrologist or owner of the device.

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GMP 7

Good Measurement Practice for Cleaning Precision Glassware

The volume contained in or delivered from precision glassware depends on the cleanliness of the container. Glassware must be sufficiently clean to ensure uniform wetting of the entire internal surface of the standard. When clean, the walls will be uniformly wetted and the water will adhere to the surface in a continuous film. If films of dirt or grease are present, the meniscus may crinkle at the edges and liquids will not drain properly and will leave water drops on the internal walls. Lack of cleanliness can cause errors in setting a meniscus and incomplete wetting/drainage of the calibration liquid (generally water.)

Glassware that is submitted for calibration should be cleaned (and autoclaved if needed) prior to submission to the laboratory to ensure removal of all chemical, biological, radiological, or other contaminants.

If glassware is cleaned by the laboratory, appropriate inquiries should be made to determine prior and subsequent uses along with implications of certain cleaning methods. Some methods of cleaning glassware prior to calibration or use are ineffective or detrimental and may leave chemical contamination deposits that can be detected in some chemical analysis operations.

There are a number of suitable detergents (both liquid and powder), available from laboratory supply houses that do not contain phosphates. The catalog descriptions usually indicate whether or not they contain phosphates. Laboratory detergents that contain phosphates can leave a deposit on the glass that will cause water to "bead" on the surface making it appear to be dirty, and making it difficult to properly set a meniscus. Once a phosphate deposit has occurred, it is very difficult to remove. It may be removed with hot (approx. 65 °C) sodium dichromate-sulfuric acid cleaning solution. However, use of this hot solution is hazardous, is a problem for safe disposal, and is not recommended unless proper safety equipment is available (see the next section).

If acetone is used to remove oil or grease film, always follow with alcohol (ethanol) before drying or rinsing with water. Denatured methanol can leave an oily residue due to the denaturing process. Acetone, if allowed to dry, may also leave a film deposit.

The above solvents need not be used if the glass does not have oil or grease film. Mechanical shaking of water and suitable non-phosphate detergents is usually adequate for cleaning glassware.

These cleaning agents do not preclude the use of other suitable methods, of which there are several. Some have special applications that would not normally be encountered by metrologists or field inspectors.

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See GLP 13, Good Laboratory Practice for Drying Containers for appropriate drying techniques. It is not necessary to dry any container marked "to deliver."

Cleaning Methods ¹

Safety Note: Material safety data sheets (MSDS) should be available, studied, and carefully followed before using any of the following cleaning methods. Chromic acid solutions are not recommended for routine use because it is a hazardous waste and hazardous to health! Suitable education or training in the handling of chemicals is highly recommended.

Frequently it is desirable to give glassware a preliminary rinse or soak with an organic solvent such as xylene or acetone to remove grease, followed by a water rinse. The rinsing with water must be done thoroughly if acid will be used later to clean the glassware.

Unless autoclaving is necessary, glassware should be cleaned as soon as possible after use to avoid setting and caking of residues. Pipets, for example, may be placed in a jar containing a weak antiseptic solution immediately after use. Autoclaving is necessary to disinfect glassware that may have been used to contain potentially dangerous biological fluids.

A selection of general cleaning methods follows:

1. Fill with sulfuric acid-dichromate mixture and let stand. After removal of the mixture, rinse with distilled water at least six times. To make the cleaning mixture, dissolve (60 to 65) g of sodium- or potassium-dichromate by heating in (30 to 35) mL of water, cool and slowly add concentrated sulfuric acid to make one liter of solution. This solution is available from laboratory supply companies.

Note: Extreme care should be exercised in handling acidic solutions.

- 2. Scrub with a 1 % to 2 % hot solution of a detergent. Rinse well after brushing. A number of suitable, commercial washing compounds are available.
- 3. Fuming sulfuric acid (very hazardous material) is an excellent cleaning agent. Usually, cleaning can be accomplished by use of a comparatively small amount of acid, manipulating the vessel so that the acid comes in contact with the entire surface, and immediately emptying and rinsing.

When a piece of glassware is badly contaminated with stopcock grease (except silicone grease), it may be necessary to rinse with acetone once or twice before using one of the above methods. For silicone grease, the acetone can be omitted and the piece soaked for 30 min in fuming sulfuric acid. Warm decahydronaphthalene (decalin) also has been suggested as a solvent for silicone grease. In this case, let the piece soak for 2 h, drain, and rinse once or twice with

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¹ While the metrologist is not ordinarily faced with the problems for which these procedures are necessary, they are given here in the event that they are needed.

acetone, followed by a water rinse.

Cautions

Care should be exercised when using most cleaning solutions because they can cause skin irritations or severe burns on contact. Dilute solutions become concentrated as the water evaporates; therefore, always flush the exposed area immediately with large quantities of water.

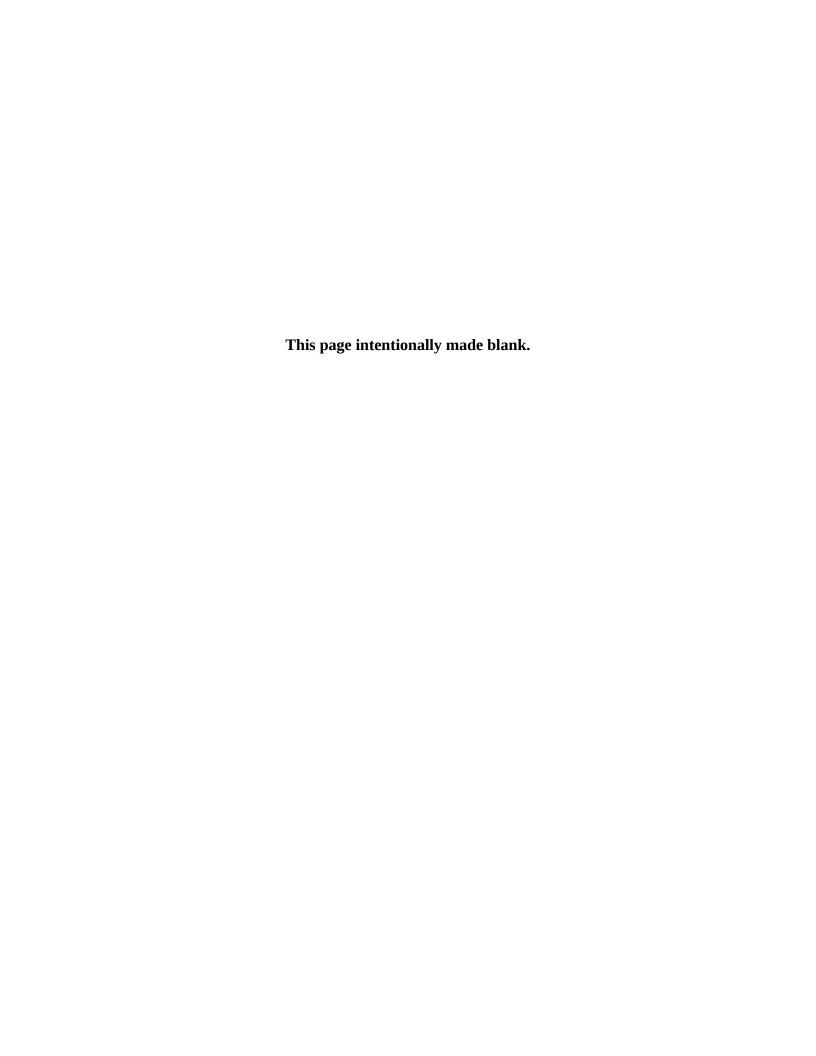
Suitable chemical-specific-resistant goggles or a face mask should be worn to protect the eyes from splashes and rubber gloves should be worn to protect the hands. It is advisable to wear an acid resistant laboratory coat or a rubber apron to protect clothing when using strong acids for cleaning. The glassware should be handled gently to avoid breakage and also to prevent spilling acids and other cleaning fluids. All cleaning should be done in a laboratory sink or on an acid-proof laboratory bench, preferably within a fume hood, to the extent possible.

Some of the cleaning materials mentioned leave minute traces or residues unless the rinsing process is carried out thoroughly. While such traces may not be harmful if the purpose of cleaning is to prepare the glassware for calibration, they can give trouble when the glassware is used in certain laboratory operations. For example, manganese and chromium compounds, even in extreme dilution, may retard or inhibit growth of micro-organisms, and traces of phosphorus may interfere with delicate tests for this element. When glassware is to be calibrated, final rinsing must be with pure distilled or deionized water.

If an article is to be dried after cleaning, as is necessary for all vessels marked "To Contain", acetone, followed by ethyl alcohol (American Chemical Society Specification) may be used. Drying may be hastened by blowing clean, dry air into the vessel (or drawing the air through the vessel). Be sure not to mix acetone with alcohol. (See GLP 13.)

Efficient air filters must be provided to remove any particles of oil or dirt from compressed air used for drying purposes.

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Standard Operating Procedures



SOP No. 14

Recommended Standard Operating Procedure for Gravimetric Calibration of Volumetric Standards Using an Electronic Balance

1 Introduction

1.1 Purpose of Test

This procedure is a precision mass calibration converting mass values to volumetric values using pure water as a standard reference material. The results provide calibration of either the "to deliver" or "to contain" volume of measuring containers that may be used as volumetric measuring standards. The procedure uses gravimetric calibration principles to minimize calibration uncertainties. Accordingly, the procedure is especially useful for high accuracy calibrations. The procedure references measurement control standards to ensure the validity of the standards and the measurement process; however, additional good measurement practices such as those required for precision mass calibrations must be used. The procedure makes use of an electronic balance and is suitable for all sizes of gravimetric calibrations only limited by the capacity and resolution of the balance and handling capabilities. Standards that are calibrated for use in legal weights and measures applications should be evaluated for conformance to the appropriate specifications and tolerances that apply. This procedure calculates the average volume based on two runs. NOTE: NIST calibrations generally make use of the average of five replicates.

1.2 Prerequisites

- 1.2.1 Verify that valid calibration certificates with appropriate values and uncertainties are available for all of the standards used in the calibration. All standards must have demonstrated metrological traceability to the international system of units (SI), which may be to the SI through a National Metrology Institute such as NIST.
- 1.2.2 Verify that the mass standards to be used have sufficiently small standard uncertainties for the level of calibration. Reference mass standards should not be used for gravimetric calibration due to the risk of water contamination. Weights of ASTM Class 2 or 3 or OIML Class F₁ or F₂ are recommended for this procedure. Working standards of other classes are generally not designed to maintain adequate stability needed; however, corrections obtained within a few days of the volume calibration may provide adequate stability and may be used if the uncertainty is sufficiently small and the density is available for performing buoyancy corrections.

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- 1.2.3 Verify that the balance used is in good operating condition with adequate capacity, sufficiently small resolution, and sufficiently small process standard deviation, as verified by a valid control chart or preliminary experiments using this procedure. NOTE: standard deviations obtained from mass calibrations do not reflect the process repeatability of gravimetric calibrations; therefore process repeatability must be obtained using this procedure! The accuracy of the balance and weighing procedures should be evaluated to minimize potential bias in the measurement process.
- 1.2.4 Verify that the operator is experienced in precision weighing techniques and has had specific training in SOP 2, SOP 4, SOP 29, GMP 3, GMP 10, and gravimetric calibrations.
- 1.2.5 Verify that an adequate quality and supply of distilled or deionized water (see GLP 10) is available. Note: Do not use tap water for this procedure!
- 1.2.6 Verify that air currents are minimized in the laboratory when weighing is performed. Verify that the laboratory facilities meet the following minimum conditions to enable meeting the expected uncertainty that is achievable with this procedure:

Table 1. Laboratory environmental conditions.

Procedure	Temperature	Relative Humidity
Gravimetric	20 °C to 23 °C Stable to \pm 1 °C / 1 h, during the calibration	40 % to 60 % Stable to ± 10 % / 4 h

2 Methodology

2.1 Scope, Precision, Accuracy

The procedure is applicable for the calibration of any size of measuring container that, when filled with water, will not overload the electronic balance used. Typical containers range in capacity from 1 mL to 20 L; however, this procedure is also applicable for larger provers provided that facility, equipment and standards meet the requirements in this SOP. When larger provers (e.g., 100 gal or more) are tested, also see the Test Notes in the Appendix. The procedure is generally only appropriate for micropipettes when additional evaporation corrections are made. The precision of calibration depends on the care exercised in adjusting the various volumes and strict adherence to the various steps of the procedure. The accuracy attainable depends on the ability of the operator to read and set the meniscus, uncertainties of the standard weights, the air buoyancy corrections, and thermal expansion corrections that are made.

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2.2 Summary

The electronic balance used is calibrated by incorporating standard masses into the procedure. The volumetric vessel to be calibrated is then weighed dry or "wetted down," depending on whether the calibration is to be made on a "To Contain" or "To Deliver" basis. The container is filled with pure water of known temperature and reweighed. The difference in mass is used to calculate the capacity of the container at the nominal capacity and at various capacities when multiple neck graduations are present. Transfer vessels may be used for all procedures except for flasks and containers marked "To Contain" (TC).

2.3 Standards and Equipment Requirements

- 2.3.1 An electronic balance having sufficient capacity to weigh the loaded vessel is required. The balance selection affects the potential measurement errors associated with this procedure. Prior to use and where available, it is recommended to use the adjust/calibrate feature on balances. Option A provides a single point calibration factor for the balance. In general, concerns are greater for balances or mass comparators larger than 5 kg due to corner loading errors, non-linearity, and repeatability over the range of use. Option B as described in the calculation section for larger volumes corrects for balance non-linearity in both the filled and empty/drained ranges of use. Option B will also likely be required on mass comparators that have multiple ranges. It also provides corrections for variations in air density that may occur over a longer weighing process on larger volumes. For all procedures in this SOP, the balance should be zeroed prior to each measurement to minimize possible effects due to balance drift. The sensitivity or resolution of the balance may be a limiting factor in the accuracy of the measurement. The resolution and repeatability must be smaller than the accepted uncertainty of the calibration. Linearity errors or additional uncertainties may need to be considered when the mass standards are slightly less than the volume or container being measured.
- 2.3.2 Sufficient quantity of calibrated mass standards. Mass standards are selected so that they are slightly larger than the combined mass of the volume and container or transfer vessel that will be weighed. When summations of masses are used, the summation mass is used, and the "effective density" must be calculated, taking care not to use "average" density values.
- 2.3.3 Thermometer with resolution and uncertainty less than 0.1 °C to determine water temperature.
- 2.3.4 Thermometer with resolution and uncertainty less than 0.50 °C to determine air temperature. ¹

- 2.3.5 Barometer with resolution and uncertainty less than 135 Pa (1 mm Hg) to determine air pressure.¹
- 2.3.6 Hygrometer with resolution and uncertainty less than 10 % to determine relative humidity. ¹
- 2.3.7 Distilled or deionized water (See GLP 10) of sufficient quality and quantity for the calibration. Note: Do not use tap water for this procedure!
- 2.3.8 Stopwatch or other suitable timing device (does not need to be calibrated.)

2.4 General Considerations

2.4.1 Cleanliness checks

Verify that all containers to be calibrated are clean as evidenced by uniform drainage of water. No water droplets should remain on any interior surface as the water drains from the container. A reproducible "wet-down" weight is evidence for cleanliness in cases where it is not possible to visually check for uniform drainage. Use GMP 6 or 7 to clean vessels as necessary. All glassware must be meticulously cleaned, prior to calibration. When clean, the walls will be uniformly wetted. Instructions for cleaning are given in GMP 6 and GMP 7. An exception is plastic ware, which will not be wetted. Follow manufacturer's instructions for cleaning such vessels. Do not use cleaning agents that will attack, discolor, or swell plastic ware.

- 2.4.2 Use water that is thermally equilibrated with the laboratory environment. Equilibration can be achieved by storing the water in clean containers in the laboratory. It is important to use water that is equilibrated in the laboratory as much as possible to minimize potential calibration errors and convection currents during the calibration. Consider water to be equilibrated when within ± 5 °C of the laboratory temperature.
- 2.4.3 Volumetric calibrations to a marked volume (graduation line) are critically dependent on the setting of a meniscus. See GMP 3 for guidance in reading a meniscus.
- 2.4.4 Use GLP 13 as the procedure to dry any container to be calibrated on a "To Contain" basis.
- 2.4.5 Wet down (not used for any container calibrated "To Contain")

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For glassware and hand-held test measures: Fill the container to capacity with distilled or deionized water, then empty over a 30 s period while avoiding splashing. Drain for 10 s unless another drain time is specified. (This is commonly called a "30 s pour, 10 s drain" emptying procedure.) A 30 s (\pm 5 s) pour followed by a 10 s drain, with the measure held between a 10 degree and 15 degree angle from vertical is required during calibration and use for glass flasks. A wet-down is not required for a transfer vessel that is used to weigh a delivered volume of water.

For stationary provers: Fill the container to capacity with distilled or deionized water, then empty. Time the drain once the cessation of the main flow is complete for 30 s and close the valve.

2.5 Calibration Procedure for Burets

- 2.5.1 Clamp the buret vertically on a support stand. Also clamp a plain glass test tube or beaker, large enough to hold a thermometer, in the vicinity of the buret.
- 2.5.2 Fill the buret with water and test for absence of leaks from the tip and stopcock. Drain and fill several times to condition the buret. Fill when ready to test.
- 2.5.3 Drain and record the delivery time, defined as the time of unrestricted flow from the zero mark to the lowest graduation mark with the stopcock fully open.
- 2.5.4 Fill the buret slightly above the zero mark with temperature-equilibrated water and also fill the test tube that holds the thermometer. Record the water temperature.
- 2.5.5 Set the meniscus on the zero mark and touch the tip with the wetted wall of a beaker to remove any excess water. The buret tip must be full.
- 2.5.6 Measure and record the air temperature, air pressure, and relative humidity.
- 2.5.7 Zero the balance and then place a known mass standard on the balance pan that is slightly larger than the filled capacity of the empty transfer vessel or flask when filled with water. Record the reading as O_1 .
- 2.5.8 Zero the balance and then weigh an empty transfer vessel or flask including the stopper or cover to be used. Record as O_2 .

- 2.5.9 Fully open the stopcock and discharge the contents of the buret into the previously weighed flask or transfer vessel. The tip of the buret should be in contact with the wall of the flask. When the level in the buret is within a few millimeters above the line being calibrated, slow the discharge and make an accurate setting. When the setting is completed, move the flask horizontally to break contact with the tip. Recheck the setting.
- 2.5.10 Stopper (or cover) the filled transfer vessel or flask. Check to make sure that the outside of the transfer vessel or flask is not wet. Zero the balance and then weigh the filled transfer vessel or flask. Record the balance reading as O₃.
- 2.5.11 Measure and record the temperature of water in the container.
- 2.5.12 Test the next interval in the same manner from the zero mark to the next interval of test.
- 2.5.13 For burets with a specified waiting time, empty as in 2.5.9 to within a few millimeters of the calibration mark. Pause for the specified waiting time (e.g., 10 s), then adjust the meniscus to the graduation line as in 2.5.9.
- 2.5.14 Measure and record the air temperature, air pressure, and relative humidity.
- 2.5.15 Make a duplicate determination for each interval (Run 2).
- 2.5.16 Calculate the volume for each interval as described in Section 3, Option A.
- 2.6 Calibration Procedure for Pipets (One-Mark)
 - 2.6.1 Measure and record the air temperature, air pressure, and relative humidity.
 - 2.6.2 Zero the balance and then place a known mass standard on the balance pan that is slightly larger than the filled capacity of the empty transfer vessel or flask when filled with water. Record the reading as O_1 .
 - 2.6.3 Zero the balance and then weigh an empty transfer vessel or flask, including the stopper or cover that will be used. Record the balance reading as O_2 .
 - 2.6.4 Fill the pipet to the index mark and measure the delivery time with the tip in contact with the internal surface of a beaker (not the transfer vessel that will be used.)
 - 2.6.5 Refill the pipet by suction, slightly above the index line. Record the water temperature. Wipe tip with filter paper, then slowly lower level to the index line,

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- using a stopcock or pinch clamp for fine control. The tip must be in contact with the wetted wall of the beaker while this setting is being made. Do not remove any water remaining on tip at this time.
- 2.6.6 Hold the pipet in a vertical position and deliver water into the previously weighed transfer vessel or flask, with the tip in contact with the inside wall or neck.
- 2.6.7 After flow has ceased, wait two seconds then remove the pipet from contact with the flask. Check to make sure that the outside of the transfer vessel or flask is not wet.
- 2.6.8 Stopper (or cover) the filled transfer vessel or flask. Zero the balance and then weigh the transfer vessel or flask. Record the balance reading as O_3 .
- 2.6.9 Measure and record the temperature of water in the container.
- 2.6.10 Measure and record the air temperature, air pressure, and relative humidity.
- 2.6.11 Make a duplicate determination for the pipet (Run 2).
- 2.6.12 Calculate the volume as described in Section 3, Option A.
- 2.7 Calibration of Flasks (To Contain) Transfer vessel must **not** be used.
 - 2.7.1 Clean and dry the flask to be calibrated as described in GLP 13. Then stopper the flask.
 - 2.7.2 Measure and record the air temperature, air pressure, and relative humidity.
 - 2.7.3 Zero the balance and then place a known mass standard on the balance pan that is slightly larger than the filled capacity of the flask when filled with water. Record the reading as O₁.
 - 2.7.4 Zero the balance and then weigh the dry flask including its stopper or cover. Record the balance reading as O_2 .
 - 2.7.5 Place an appropriate sized funnel in neck and fill the flask to just below the reference graduation while maneuvering the flask to wet the entire neck below the stopper. Let stand for two minutes then adjust the meniscus to the reference graduation line, taking care to avoid wetting the neck above the capacity graduation.
 - 2.7.6 Check that the outside surface of the flask, and its internal surface above the

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- water level are dry, and that neither bubbles or foam are present in the water. Remove visible water droplets if needed with a laboratory wipe. Stopper or cover the flask.
- 2.7.7 Zero the balance and then weigh the filled flask and its stopper or cover, and record the balance reading as O_3 .
- 2.7.8 Measure and record the temperature of the water in the flask or other container.
- 2.7.9 Measure and record the air temperature, air pressure, and relative humidity.
- 2.7.10 Make a duplicate determination (Run 2) after drying the flask per GLP 13.
- 2.7.11 Perform volume calculations as described in Section 3, Option A.
- 2.8 Calibration of Flasks (To Deliver)
 - 2.8.1 Clean but do not dry the flask to be calibrated. Condition the flask by filling and emptying the flask over a 30 s period by gradually inclining it to avoid splashing. When the main flow has ceased, hold the flask in a nearly vertical position for 10 seconds unless another drain time is specified, then touch off the drop of water adhering to the tip of the flask.
 - 2.8.2 Measure and record the air temperature, air pressure, and relative humidity.
 - 2.8.3 Zero the balance and then place a known mass standard on the balance pan that is slightly larger than the filled capacity of the flask when filled with water. Record the reading as O_1 .
 - 2.8.4 Zero the balance and then weigh the drained (empty) flask including its stopper or cover. Record the balance reading as O_2 .
 - 2.8.5 Fill the flask to just below the reference graduation. Let the flask stand for two minutes, then adjust the meniscus to the reference graduation line. Do not dry the inside of the flask above the graduation line, but take care to ensure that the outside of the flask is dry.
 - 2.8.6 Zero the balance and then weigh the filled vessel, with cap or stopper. Record the balance reading as O_3 .
 - 2.8.7 Measure and record the temperature of the water in the flask.
 - 2.8.8 Measure and record the air temperature, air pressure, and relative humidity.

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- 2.8.9 Make a duplicate determination (Run 2).
- 2.8.10 Calculate the volume of the flask as described in Section 3, Option A.

Note: The order of weighing the filled and drained flask in Section 2.8 may be reversed. In that case, the filled vessel is still weighed and recorded as O_3 , the flask is drained as in 2.8.1, and the properly drained vessel is weighed and recorded as O_2 .

2.9 Calibration of Other Volumetric Glassware

- 2.9.1 Measuring Pipets (non-single-mark) Calibrate in a manner similar to that used to calibrate burets (2.5).
- 2.9.2 Graduated Cylinders Calibrate in a manner similar to that used for flasks (To Deliver cylinders, use section 2.8; To Contain cylinders, use section 2.7).

2.10 Calibration of Slicker-Plate Type Standards

- 2.10.1 This calibration makes use of Option B where equivalent mass standards are used at both the filled and empty loads on the balance. A transfer vessel is recommended.
- 2.10.2 Condition the slicker-plate type standards with several wet-down runs to fully ensure wet down and smooth valve operation.
- 2.10.3 Measure and record the air temperature, air pressure, and relative humidity.
- 2.10.4 Zero the balance and then place a standard mass, M_{S1} , on the balance platform (M_{S1} should be slightly larger than the mass of the drained vessel, dry vessel, or empty transfer vessel.) Record reading as O_1 .
- 2.10.5 Zero the balance. Place the dry or "wet-down" transfer vessel on the balance platform, as appropriate, and record reading as O_2 . Caution: all containers must be dry on the outside for all weighing.
- 2.10.6 Fill the slicker-plate standard to just above the rim of the standard. Record the water temperature. Slide the slicker plate across the level top. Set the transfer vessel below the nozzle to ensure all volume is transferred into the vessel and that no splashing occurs.
- 2.10.7 Open the slicker-plate standard valve and remove the plate simultaneously and smoothly to deliver the water into the transfer vessel. Time the drain for 30 s after cessation of the main flow and close the valve. Cover the transfer vessel

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- and move it from beneath the standard to ensure additional drops of water are not transferred. Note: never fill the transfer vessel while it is sitting on the balance platform to ensure that the balance is not damaged with water and to avoid errors associated with balance drift and hysteresis.
- 2.10.8 Zero the balance and then place a standard mass, M_{S2} , on the balance platform (M_{S2} should be slightly larger than the mass of the filled vessel.) Record the balance reading as O_3 .
- 2.10.9 Zero the balance and then weigh the filled transfer vessel and record the balance reading as O_4 .
- 2.10.10 Immediately after weighing, check the temperature of the water in the filled container. Ensure that the water has not changed by more than $0.2~^{\circ}\text{C}$ during the measurement process or repeat the run.
- 2.10.11 Measure and record the air temperature, barometric pressure, and relative humidity.
- 2.10.12 Make a duplicate determination (Run 2).
- 2.10.13 Calculate the volume as described in Section 3, Option B.
- 2.11 Calibration of Hand-held Graduated Neck Type Provers
 - 2.11.1 A hand-held test measure is weighed empty and filled much like the procedure for To Deliver flasks (section 2.8). However, this calibration makes use of Option B where equivalent mass standards are used at both the filled and empty loads on the balance.
 - 2.11.2 Condition the test measure by filling and draining using a 30 s pour followed by a 10 s drain. Take care to avoid splashing of water on the outside of the measure. The wet-down will help equilibrate the test measure with the water temperature.
 - 2.11.3 Measure and record the air temperature, air pressure, and relative humidity.
 - 2.11.4 Zero the balance and then place standard masses, $M_{\rm S1}$, on the balance platform ($M_{\rm S1}$ should be slightly larger than the mass of the drained vessel.) Record reading as O_1

- 2.11.5 Zero the balance and then place the "wet-down" test measure on the balance platform, and record the balance reading as O_2 . Caution: all containers must be dry on the outside for all weighing.
- 2.11.6 Fill the graduated neck type test measure to just below the nominal volume mark. Record the water temperature. Adjust the meniscus to nominal.
- 2.11.7 Zero the balance and then place standard masses, M_{S2} , on the balance platform (M_{S2} should be slightly larger than the mass of the filled vessel.) Record the balance reading as O_3 .
- 2.11.8 Zero the balance and then weigh the filled transfer vessel and record the balance reading as O_4 .
- 2.11.9 Immediately after weighing, check the temperature of the water in the filled container. Ensure that the water has not changed by more than 0.2 °C since step 2.11.6 or repeat the run.
- 2.11.10 Measure and record the air temperature, barometric pressure, and relative humidity.
- 2.11.11 Make a duplicate determination (Run 2).
- 2.11.12 Calculate the volume as described in Section 3, Option B. You may use the average water temperature in calculations. If a neck scale plate verification is performed, additional weighings may be performed after setting the meniscus at each level.
- 2.12 Calibration of Large Volume Graduated Neck Type Provers
 - 2.12.1 This calibration makes use of Option B where approximately equivalent mass standards are used at both the filled and empty loads on the balance. A transfer vessel is recommended. The unknown vessel to be calibrated must be elevated to facilitate transfer of water into the transfer vessel.
 - 2.12.2 Condition the prover with a wet-down run to fully ensure smooth valve operation, temperature equilibration, and wet-down of the delivery system.
 - 2.12.3 Measure and record the air temperature, air pressure, and relative humidity.
 - 2.12.4 Zero the balance and then place standard masses, M_{S1} , on the balance platform (M_{S1} should be slightly larger than the mass of the drained vessel, dry vessel, or empty transfer vessel.) Record reading as O_1 .

- 2.12.5 Zero the balance. Place dry or "wet-down" transfer vessel including cover on balance platform, as appropriate, and record reading as O_2 . Caution: all containers must be dry on the outside for all weighing.
- 2.12.6 Fill the graduated neck type prover to just below the nominal volume mark. Record the water temperature. Adjust the meniscus to nominal. Set the transfer vessel below the nozzle to ensure all volume is transferred into the vessel and that no splashing occurs. Use hoses and piping of minimum length to avoid errors associated with retention during delivery. Hoses should be translucent so that cessation of the main flow is visible. Care should be taken to ensure that there are no loops or sags in the hose that may hold water. When using hoses or piping, be sure to include them as part of the wet-down process.
- 2.12.7 Open the unknown prover valve to deliver the water into the transfer vessel. Time the drain for 30 s after the cessation of the main flow and close the valve. Cover the transfer vessel and move it from beneath the prover to ensure additional drops of water are not transferred.
- 2.12.8 Zero the balance and then place standard masses, M_{S2} , on the balance platform (M_{S2} should be slightly larger than the mass of the filled vessel.) Record the balance reading as O_3 .
- 2.12.9 Zero the balance and then weigh the filled transfer vessel and its cover. Record the balance reading as O_4 .
- 2.12.10 Immediately after weighing, check the temperature of the water in the filled container. The temperature should not change by more than 0.2 °C if the prover and water have been equilibrated in the laboratory environment.
- 2.12.11 Measure and record the air temperature, barometric pressure, and relative humidity.
- 2.12.12 Make a duplicate determination (Run 2).
- 2.12.13 Calculate the volume as described in Section 3, Option B. You may use the average water temperatures for calculations.

3 Calculations

3.1 Option A – One-point balance calibration. Compute the volume at the temperature under test, V_t , for each determination using the equation:

$$V_{t} = \left(O_{3} - O_{2}\right) \left(\frac{M_{s}}{O_{1}}\right) \left(1 - \frac{\rho_{a}}{\rho_{s}}\right) \left(\frac{1}{\rho_{w} - \rho_{a}}\right)$$
 Eqn. 3.1

Note: the O_3 value is the filled volume container or transfer vessel and O_2 is the dry, empty, or drained value. If the order of any weighing operations were changed, the calculated volume may indicate a negative value.

3.2 Option B – Two point balance calibration. Compute the volume, V_t , for each determination using the equation:

$$V_{t} = \left[\frac{O_{4}}{O_{3}}M_{s2}\left(1 - \frac{\rho_{a}}{\rho_{s2}}\right) - \frac{O_{2}}{O_{1}}M_{s1}\left(1 - \frac{\rho_{a}}{\rho_{s1}}\right)\right]\left(\frac{1}{\rho_{w} - \rho_{a}}\right) \quad \text{Eqn. 3.2}$$

Table 2. Variables for volume equations.

Variable	Description
M_s, M_{s1}, M_{s2}	mass of standards (i.e., true mass, vacuum mass) (g)
$ ho_{s}$	density of M_S (g/cm ³)
$ ho_{\scriptscriptstyle W}$	density of water at the temperature of measurement (g/cm³)
$ ho_a$	density of air at the conditions of calibration (g/cm³)
V_t	represents either the "to contain" or "to deliver" volume (depending on whether O_2 represents a dry or a "wet down" container at the temperature of the measurement) (cm³ or mL)

Glassware is typically calibrated to 20 °C. Compute V_{20} , the volume at 20 °C, for each run, using the expression:

$$V_{20} = V_t \left[1 - \alpha \left(t - 20 \right) \right]$$
 Eqn. 3.3

where α is the cubical coefficient of expansion of the container being calibrated (°C⁻¹), (see NISTIR 6969, Table 9.10), and, t, is the temperature (°C) of the water. Compute the mean V_{20} for the duplicate measurements.

3.4 Test measures and provers are typically calibrated to 60 °F. Compute V_{60} , the volume at 60 °F, for each run, using the expression (taking care to use the cubical coefficient of expansion in °F⁻¹):

$$V_{60} = V_t \left[1 - \alpha \left(t - 60 \right) \right]$$
 Eqn. 3.4

3.5 If using a different reference temperature, use the following equation and take care to match the cubical coefficient of expansion units with the units of temperature:

$$V_{\text{ref}} = V_{t} \left[1 - \alpha \left(t - t_{\text{ref}} \right) \right]$$
 Eqn. 3.5

3.6. Other Reference temperatures may be used. Common reference temperatures for other liquids follow:

Commodity	Reference Temperature
Frozen food labeled by volume (e.g., fruit juice)	-18 °C (0 °F)
Beer	3.9 °C (39.1 °F)
Food that must be kept refrigerated (e.g., milk)	4.4 °C (40 °F)
Distilled spirits or petroleum	15.56 °C (60 °F)
Petroleum (International Reference)	15 °C (59 °F)
Compressed Natural Gas (CNG)	15 °C (60 °F)
Wine	20 °C (68 °F)
Unrefrigerated liquids (e.g., sold unchilled, like soft drinks)	20 °C (68 °F)
Hydrogen (H ₂)	21 °C (70 °F)
Petroleum (Hawaii)	26.67 °C (80 °F)

4 Measurement Assurance

- 4.1 Duplicate the process with a suitable check standard (See GLP 1, SOP 30, in NISTIR 6969, or Sec. 7.4, in NBS Handbook 145) or conduct replicate measurements per SOP 17 or 20. Average values of the range or standard deviation of similarly sized flasks or volumetric standards may be tracked on a single range chart or standard deviation chart according to SOP 17 or 20. A minimum of 12 replicate measurements are required to establish initial process limits and 25 to 30 points are required for reporting valid uncertainty values.
- 4.2 Plot the check standard volume and verify it is within established limits. A t-test may be incorporated to check the observed value against an accepted value.
- 4.3 The mean of the check standard is used to evaluate bias and drift over time and may be used to identify or signify problems with the volume standard or changes in water quality.
- 4.4 Check standard observations are used to calculate the standard deviation of the measurement process, s_p .

5 Assignment of Uncertainties

The limits of expanded uncertainty, U, include estimates of the standard uncertainty of the mass standards used, u_c , plus the measurement process repeatability, s_p , at approximately a 95

percent level of confidence. See SOP 29 for the complete standard operating procedure for calculating the uncertainty.

- 5.1 The standard uncertainty for the standards, u_s , is obtained from the calibration report. The combined standard uncertainty, u_c , is used and not the expanded uncertainty, U, therefore the reported uncertainty for the standard will usually need to be divided by the coverage factor k. Multiple mass standards are often used, so see SOP 29 for treatment of dependencies and correlations.
- 5.2 Standard deviation of the measurement process from control chart performance (See SOP No. 17 or 20.)

The value for s_p is obtained from the control chart data for check standards when a check standard is available using this procedure. It may also be estimated based on replicate measurements over time. Replicate measurements over time may provide a pooled standard deviation that may be used or the average range is used to estimate the standard deviation per SOP 17 or 20. This standard deviation value is a repeatability factor related to the precision of the weighings and the setting of the meniscus when present, but does not include uncertainties associated with errors in reading the meniscus (See GMP 3.)

- 5.3 Include uncertainties associated with the reading of the meniscus when one is present. See GMP 3 for details.
- 5.4 Other standard uncertainties usually included at this calibration level include uncertainties associated with water temperature measurements, thermometer accuracy, calculation of air density, standard uncertainties associated with the density of the mass standards, coefficients of expansion, viscosity or surface effects on the volume of liquid clinging to vessel walls after draining, improper observance of drainage times, and the lack of internal cleanliness.
- 5.5 Example components to be considered for an uncertainty budget table are shown in Table 3. Multiple values may need to be considered in some cases.

Table 3. Example uncertainty budget table.

Uncertainty Component Description	Symbol	Source	Typical Distribution
Uncertainty of the standard mass(es) (5.1)	u_s	Calibration report	Rectangular or Normal divided by coverage factor
Accepted standard deviation of the process (5.2)	S_p	Control chart, standard deviation chart No:	
Water temperature	u_{tx}	Consider accuracy, resolution, and gradients	Rectangular

Water density	$u_{\rho w}$	GLP 10	Rectangular	
Cubical Coefficient of Expansion	u_{CCE}	5 % to 10 % (EURAMET CG-19)	Rectangular	
Meniscus reading (when present) (5.3)	u_m	GMP 3	Triangular	
Drain time effects	u_d	From experimental data	Rectangular	
Air temperature (for air density)	u_t	SOP 2	Rectangular	
Air pressure (for air density)	u_p	SOP 2	Rectangular	
Air relative humidity (for air density)	u_{RH}	SOP 2	Rectangular	
Air density (equation)	$u_{\rho a}$	SOP 2	Rectangular	
Mass density	$u_{ ho m}$	0.05 g/cm ³ NIST Reference	Rectangular	

6 Report

- 6.1 Report results as described in SOP 1, Preparation of Calibration/Test Results, with the addition of the following:
 - 6.1.1 "To Contain" or "To Deliver" volume, reference temperature, uncertainty, material, thermal coefficient of expansion (assumed or measured), construction, any identifying markings, tolerances (if appropriate), laboratory air temperature, water temperature(s) at time of test, barometric pressure, relative humidity, and any out-of-tolerance conditions.

Additional References:

Bean, V. E., Espina, P. I., Wright, J. D., Houser, J. F., Sheckels, S. D., and Johnson, A. N., NIST Calibration Services for Liquid Volume, NIST Special Publication 250-72, National Institute of Standards and Technology, Gaithersburg, MD, (2009).

EURAMET Calibration Guide 19, Guidelines on the Determination of Uncertainty in Gravimetric Volume Calibration, (Version 2.1, 03/2012).

Appendix A (Single-Point Balance Calibration)Gravimetric Calibration Data Sheet (Option A)

Laboratory data and conditions:

Vessel ID	Operator		
M aterial	Date		
Cubical Coefficient of Expansion		Before	After
Balance Identification	Temperature		
Load	Pressure		
Standard deviation of the process, from control chart, s_p	Relative Humidity		
Degrees of Freedom	Reference water temperature		

Mass standard(s) data:

ID (Note ID and for Filled or Empty Load)	Nominal	Mass Correction*	Expanded Unc: From cal. report	Unc: k factor	Density g/cm ³
S					
S					
S					
S					
S					

^{*}Mass Correction = $True\ Mass$ values are required.

Observations:

Run 1	Weights – Zero balance before each weighing.	Balance Observations, Units	
	M_S	O_1	
	Empty or Drained	O_2	
	Filled	O_3	
		t_w	
Run 2	Weights – Zero balance before each weighing.	Balance Observations, Units	
	M_S	O_1	
	Empty or Drained	O_2	
Filled		O_3	
		t_w	

Appendix B (Two-Point Balance Calibration)Gravimetric Calibration Data Sheet (Option B)

Laboratory data and conditions:

Vessel ID	Operator		
Material	Date		
Cubical Coefficient of Expansion		Before	After
Balance	Temperature		
Load	Pressure		
Standard deviation of the process, from control chart, s_p	Relative Humidity		
Degrees of Freedom	Reference water temperature		

Mass standard(s) data:

ID (Note ID for Empty (M _{S1}) or Filled (M _{S2}) Loads)	Nominal	Mass Correction*	Expanded Unc: From cal. report	Unc: k factor	Density g/cm ³
S					
S					
S					
S					
S					
S					
S					

^{*}Mass Correction = $True\ Mass$ values are required.

Observations:

Run 1	1 Weights – Zero balance before each weighing.		Balance Observations, Units	
		M_{s1}	O_1	
		Empty or Drained	O_2	
		M_{S2}	O_3	
		Filled	O_4	
t_w in standard:		t_w in unknown:	: Temperature change is less than 0.2 °C? Y/I	
Run 2	Run 2 Weights – Zero balance before each weighing.			Balance Observations, Units
M_{s1}		O_1		
Empty or Drained		Empty or Drained	O_2	
M_{S2}		O_3		
Filled		O_4		
t_w in standard:	_	t_w in unknown:		Temperature change is less than 0.2 °C? Y/N

Appendix C

Test Notes for Large Provers

1 Pour and drain times.

It is impractical to completely drain a filled container, because some of the contents will remain as a film. By strict adherence to a specified procedure, the residual contents can be held essentially constant so that, reproducible calibration constants can be obtained. The conventionally selected conditions are as follows:

- a. For bottom-drain containers: open drain valve fully and allow contents to discharge at maximum rate. When flow ceases, wait 30 s, close valve, and touch off any drops adhering to spout.
- b. For pour-type containers: pour contents by gradually tilting container to an 85° angle, so that virtually all liquid is delivered in 30 s. Allow to drain for an additional 10 s, then touch off any drops adhering to the lip.

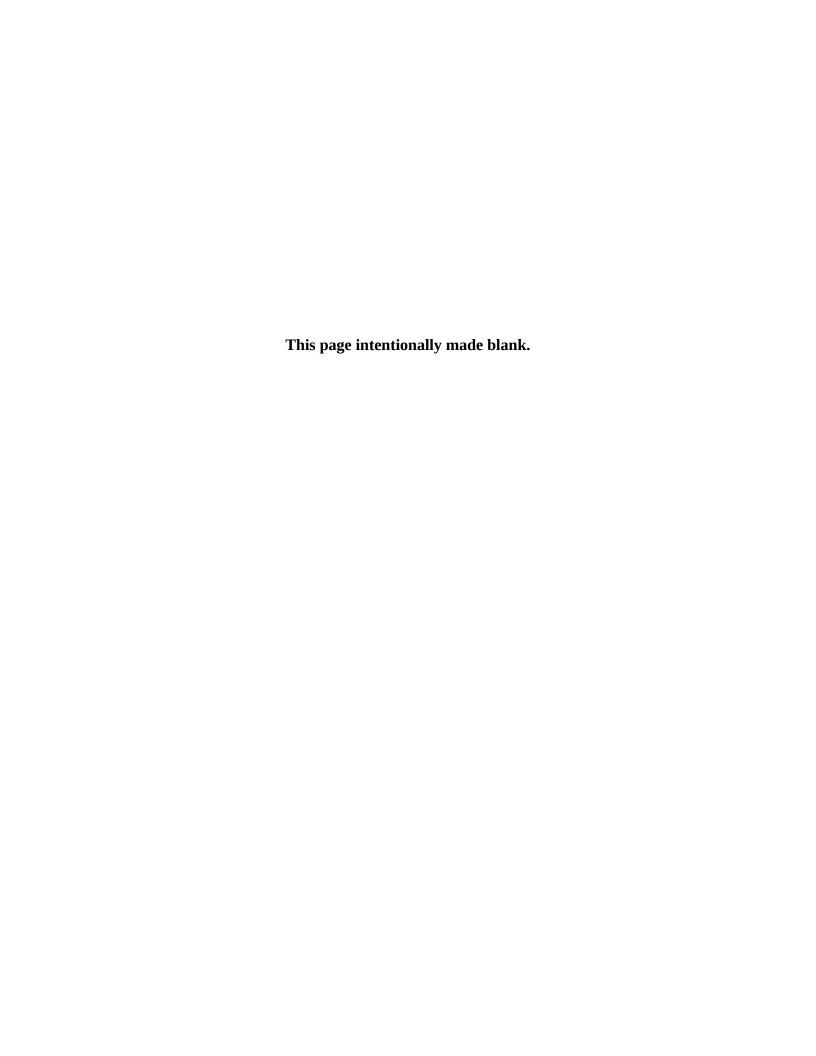
The instructions described above must be precisely followed during calibration and use of the calibrated vessels.

2 Evaporation losses.

A suitable cap should be placed on the top of open vessels to minimize evaporation losses. If used, the cap must be included in all weighings.

3 Slicker-plate.

When a slicker-plate standard is calibrated, the plate should be used to fix the water level in it. A transfer vessel is recommended for use when calibrating a slicker-plate type standard. But if the standard is weighed, the plate must be weighed with the standard during each such operation.



SOP 16

Standard Operating Procedure for Calibration of Measuring Flasks Volume Transfer Method

1 Introduction

1.1 Purpose of Test

This procedure describes a method for volume transfer calibration of the "to deliver" volume of measuring flasks from calibrated volumetric standards. The test items are then used as volumetric measurement standards, often for packaged commodity verification. If "to contain" calibrations are to be performed, appropriate modifications are needed in the procedure to ensure complete drying of the flask between calibration runs.

1.2 Prerequisites

- 1.2.1 Verify that valid calibration certificates are available for all of the standards used in the calibration. All standards must have demonstrated metrological traceability to the international system of units (SI), which may be to the SI through a National Metrology Institute such as NIST.
- 1.2.2 Verify that the standards to be used have sufficiently small standard uncertainties for the intended level of calibration.
- 1.2.3 Verify that the standard burets and pipets are clean, free of leaks, and in good operational condition.
- 1.2.4 Verify the availability of an adequate supply of pure distilled or deionized water (See GLP 10) of sufficient quality and quantity for the calibration. Do not use tap water for this procedure to ensure the reference glassware is maintained with valid reference values!
- 1.2.5 Verify that the operator has had specific training and is proficient in SOP 16, GMP 3, GMP 7, and is familiar with the operating characteristics and conditioning of pipets and burets.
- 1.2.6 Verify that the laboratory facilities meet the minimum conditions shown in Table 1 to meet the expected uncertainty that is achievable with this procedure.

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Table 1. Laboratory environmental conditions.

Procedure	Temperature	Relative Humidity
Volume	$18 ^{\circ}\text{C}$ to $27 ^{\circ}\text{C}$,	40 % to 60 %
Transfer	Stable to $\pm 1 ^{\circ}\text{C} / 1$ h during the calibration	Stable to ± 20 % / 4 h

2 Methodology

2.1 Scope, Precision, Accuracy

The method is applicable for the calibration of any size of measuring flask for which standard pipets of comparable volume are available. Typical flasks have volumes in the range of 100 mL to 5 L (1 gill to 1 gallon). The precision of calibration depends on the care exercised in adjusting the various volumes and strict adherence to the various steps of the procedure. The accuracy will depend on the accuracy of calibration of the volumetric standards used, accuracy in reading the meniscus, together with the precision of the comparison. Clean glassware and strict adherence to the drainage instructions are essential for precise and accurate results.

2.2 Summary

The flask to be calibrated is given an initial wet-down using the correct drain procedure, then nearly filled with water delivered from a calibrated pipet. (This procedure relies on calibrated pipets that deliver slightly less than the full nominal volume.) Additional water is added to the test flask from a calibrated buret until the meniscus in the test item coincides with the calibration graduation mark. The sum of the volumes delivered into the wetted flask is equivalent to its "to deliver" volume. The process described assumes that the flask is marked with a 10-second drain time. If the flask is marked with a drain time other than 10 s, the specified drain time should be used rather than the 10 s drain time described. A 30-second drain time should be used when a flask has no drain time specified.

2.3 Standards and Equipment

- 2.3.1 Calibrated standard pipet(s) of suitable volumes. These standards are typically made of borosilicate glass.
- 2.3.2 Calibrated standard buret(s) of 10 mL (or 120 minim¹) capacity. These standards are typically made of borosilicate glass.

-

 $^{^{1}}$ 1 [US] minim = 0.061615496 milliliter (mL). 120 minim standards were provided to the States as reference standards for use in volume transfer calibrations. These standards have higher resolution than metric standards provided at the same time.

- 2.3.3 Calibrated thermometers, with resolution and uncertainty to ≤ 0.1 °C to determine water temperature.
- 2.3.4 Meniscus reading device (See GMP 3).
- 2.3.5 Stopwatch or other suitable timing device (does not need to be calibrated).

2.4 Procedure

2.4.1 Cleanliness check

Verify that all glassware, including the standards used and the vessels to be calibrated, are internally clean, as evidenced by uniform drainage of water. No water droplets should remain on the internal surfaces as the water drains from the vessels. If this occurs, the glassware must be cleaned with suitable agents and rinsed with pure water until uniform drainage is obtained. (See GMP 7).

2.4.2 Wet-down of pipet and flask

- 2.4.2.1 Fill the standard pipet to until water flows from the top tip, then completely drain into an empty vessel. Check for uniform drainage. Touch off the pipet outlet tip against the container wall to remove excess droplets and to establish a constant tip retention volume. The pipet and bore of the delivery side of the stopcock should appear to be "empty" and the delivery tip should contain a small volume of water retained in it. This is the "wet-down" condition of the pipet. It must be done at the beginning of each test sequence. Ensure that the outside of the standard pipet is free of water droplets that may fall into the unknown flask.
- 2.4.2.2 Refill the pipet and use it to fill the flask to be calibrated. Empty the flask using a gentle pour in a 30 s ± 5 s period by gradually inclining the flask so as to avoid splashing of the walls as much as possible. When the main drainage stream has ceased, the flask will be nearly vertical. Allow an additional 10 second drainage after discharge of the flask contents while holding the flask at a 10° to 15° angle from vertical, then touch off the rim of the flask to remove any drops adhering to it. At the same time, check that uniform drainage has been achieved. This establishes the "to deliver" condition of the flask. Note that some small amount of water will remain in the flask. Each flask to be calibrated must be given this "wet-down" treatment immediately before the start of the test.

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2.4.3 Conditioning the buret

- 2.4.3.1 Fill the standard buret to overflow and drain several times to verify uniform drainage. Refill with water. Note that the stopcock bore and delivery tip should be filled with water at all times, in contrast to the condition for the pipet.
- 2.4.3.2 Touch off the delivery tip against the wall of the receiving vessel, to remove any droplet adhering to the external surface of the tip. This is an operation that must be done every time a measured volume of water is delivered from the buret to ensure consistent volume delivery.

2.4.4 Calibration

- 2.4.4.1 Run 1. Fill the standard pipet to overflow. The delivery bore and the delivery tip will be empty, except for the small volume retained in the tip. Place the inside of the flask neck in contact with the tip of the pipet or buret, to avoid splashing but in a manner that does not block the flow of water.
- 2.4.4.2 Deliver the contents of the pipet into the "wet-down" flask. Repeat as necessary to nearly fill the flask to the calibration line. The value recorded for the standard pipet will be the total of the deliveries used.
- 2.4.4.3 Add water to the flask from the buret until the meniscus coincides with the calibration mark. (See GMP 3 for instructions on how to read a meniscus.) Read the volume of water delivered from the buret and record on a suitable data sheet such as the one in the Appendix. Note: If multiple deliveries of the complete buret contents are required, ensure that at no time the buret water level drops below the lowest graduation mark, or the entire process must be started over. The value recorded for the standard buret will be the sum of the delivered volumes.
- 2.4.4.4 Empty the measuring flask as described in 2.4.2.2 to re-establish "wet-down."
- 2.4.4.5 Record all data using the form given in the Appendix or a similar format.
- 2.4.4.6 Run 2. Replicate Measurement. Repeat the procedure described in 2.4.4. The test measure must be capable of repeating to 0.01 % of the test volume during calibration. Repeatability problems may be due to contamination, poor meniscus readings, lack of cleanliness, or conduct

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of the calibration in an unstable environment. Repeatability problems must be corrected before calibration can be completed.

2.4.4.7 Calibration Note: If the test vessel has a graduated neck (in addition to the nominal mark) additional graduations should be inspected for proper marking and calibrated as well.

3 Calculations

3.1 Compute the individual "to deliver" volumes, V_{TD} , for Run 1 and Run 2.

$$V_{TD} = V_P + V_B$$
 Eqn. 1

Table 2. Variables for volume equation.

Variable	Description
V_{TD}	"To deliver" Volume
V_P	"To deliver" calibrated volume of the standard pipet
V_B	Volume of water delivered from the buret, corrected for any calibration values

3.2 Temperature correction

For calibration of glassware that is not made of borosilicate glass, not only should the above temperature conditions be realized but the water temperature must also be known. The water used must be stored in the laboratory until its temperature is equilibrated with its surroundings and the temperature of the discharged water is measured, using the calibrated thermometer. The "To Deliver" volume of the flask, corrected to the reference temperature of 20 °C, V_{TD20} , is computed, using the expression:

$$V_{TD\,20} = V_{TD} \left[1 + (\alpha - \beta)(t - 20) \right]$$
 Eqn. 2

Table 3. Variables for volume temperature correction equation.

Variable	Description	
V_{TD}	observed delivered volume as computed in 3.1	
V_{TD20}	"To deliver" volume at the reference temperature of 20 °C	
t	Temperature of water at time of calibration	
α	Cubical coefficient of expansion of glass of standard (e.g., 0.000010 °C ⁻¹ for borosilicate glass)	
β	Cubical coefficient of expansion of glass of flask (e.g., 0.000025 °C ⁻¹ for soda-lime glass)	

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The volumetric standards are typically made of borosilicate glass. If borosilicate flasks are calibrated, no temperature correction is required. This procedure relies on the fact that since deliveries are made over a short period of time, the temperature remains the same in the standard as in the unknown. When standards and flasks are not of the same material, the water temperature change must be less than 0.5 °C while the water is in the standards and the flask.

Report the average volume V_{TD20} as the value for the flask after the temperature correction has been applied to each Run.

4 Measurement Assurance

- 4.1 Duplicate this process with a suitable check standard (Quality Assurance Reference Flask, QARF) or have a suitable range of check standards for the laboratory with similar size graduations and neck diameters. See SOP 17 and SOP 30.
- 4.2 Plot the check standard volume and verify that it is within established limits. Alternatively a *t*-test may be incorporated into the process to check the observed value against an accepted reference value. (See NISTIR 6969, Section 8).
- 4.3 The mean of the check standard value is used to evaluate bias and drift over time.
- 4.4 Check standard observations are used to calculate the standard deviation of the measurement process.

5 Assignment of Uncertainties

The limits of expanded uncertainty, U, include estimates of the standard uncertainty of the laboratory volumetric standards used, u_s , plus the standard deviation of the process, s_p , at the 95 percent level of confidence. See SOP 29 for the complete standard operating procedure for calculating the uncertainty.

- The standard uncertainty for the standard, u_s , is obtained from the calibration reports for the pipet and buret used for the calibration. The combined standard uncertainty, u_c , from the calibration report must be used and not the expanded uncertainty, U. Therefore, the reported uncertainty for the standard will usually need to be divided by a coverage factor, k. Multiple standards are used in this procedure; however, they are typically calibrated independently. Multiple standard uncertainties are handled according to SOP 29.
- 5.2 The standard deviation of the measurement process, s_p , is obtained from control chart performance (See SOP 17 or 20, and SOP 30) for a suitable QARF using the same volume transfer procedures on flasks of comparable size and neck dimensions.
- 5.3 Other standard uncertainties usually included at this calibration level include uncertainties associated with the ability to read the meniscus, only part of which is included in the process variability (See GMP 3), the cubical coefficient of expansion for the flask under test, use of proper temperature corrections, the accuracy of temperature measurements taking into consideration potential gradients in the glassware during calibration, round robin data showing reproducibility, environmental variations over time, and bias or drift of the standard.

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6 Report

- Report results as described in SOP 1, Preparation of Calibration/Test Results, with the addition of the following items:
 - 6.1.1 "To Contain" or "To Deliver" volume, applicable reference temperature, uncertainty, material, thermal coefficient of expansion (assumed or measured), construction, any identifying markings, tolerances (if appropriate), laboratory temperature, water temperature at time of test, barometric pressure, relative humidity, and any out-of-tolerance conditions.
- 7 Alternative Reference Temperatures
 - 7.1 Reference temperatures other than 20 °C (68 °F) may occasionally be used. Common reference temperatures for other liquids follow:

Commodity	Reference Temperature
Frozen food labeled by volume (e.g., fruit juice)	- 18 °C (0 °F)
Beer	3.9 °C (39.1 °F)
Food that must be kept refrigerated (e.g., milk)	4.4 °C (40 °F)
Distilled spirits or petroleum	15.56 °C (60 °F)
Petroleum (International Reference)	15 °C (59 °F)
Wine	20 °C (68 °F)
Unrefrigerated liquids (e.g., sold unchilled, like soft drinks)	20 °C (68 °F)
Petroleum (Hawaii)	26.67 °C (80 °F)

7.2 When alternative reference temperatures are used, at the request of the user of the field standard, and when the glassware is *not* marked with a reference temperature, the calibrated value of the standard may be reported at a different reference temperature, t_{ref} , by using this equation:

$$V_{TDtref} = V_{TD} \left[1 + (\alpha - \beta) (t - t_{ref}) \right]$$
 Eqn. 3

Additional References:

Bean, V. E., Espina, P. I., Wright, J. D., Houser, J. F., Sheckels, S. D., and Johnson, A. N., NIST Calibration Services for Liquid Volume, NIST Special Publication 250-72, National Institute of Standards and Technology, Gaithersburg, MD, (2009).

Appendix

Calibration of Measuring Flasks (Volume Transfer Method)

Laboratory data and conditions:

·			
Test Number	Date		
Vessel ID	Operator		
Nominal Volume		Before	After
Coefficient of Expansion	Temperature		
Process standard deviation	Pressure		
from control chart, s_p			
Degrees of Freedom	Relative Humidity		

Volume standard(s) data:

ID (enter interval if needed)	Nominal	Calibrated Volume	Volume Correction	Coefficient of Expansion	Uncertainty	k factor
Pipet						
Buret						
Buret						
Buret						

Observations:

	Run 1	Run 2
Calibrated Pipet Volume: V_p		
Buret Volume:		
Final Reading		
Initial Reading		
Difference		
Calibration Correction (if required)		
Corrected Buret Volume, V_B		
$V_{TD} = V_P + V_B$		
Water temperature at time of test (t)		
$V_{TD20} = V_{TD} [1 + (\alpha - \beta)(t - 20)]$		
Average V_{TD20}		

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SOP 17

Standard Operating Procedure for Control Charts of Check Standards

1 Introduction

1.1 Purpose

This procedure may be used to develop and maintain control charts to monitor the statistical control of laboratory check standards when replicate measurements are made as a part of the standard operating procedure. This procedure may be used for volume, length, time, or other calibrations when replicate measurements are made.

1.2 Prerequisites

- 1.2.1 The procedure to be monitored must match the calibration procedure that is used.
- 1.2.2 Either a check standard at each nominal value is used or a set of check standards are selected to monitor the range of items that are calibrated by the laboratory.

2 Summary

A check standard is obtained and calibrated several times initially to establish a reliable mean value and to estimate the standard deviation of calibration. All such calibrations are made using the applicable procedure that is used for calibration. A reference value may be obtained from a suitable calibration laboratory or by using a more accurate and precise calibration than the one that will be monitored. Directions for preparing and using a control chart for monitoring the mean (\bar{x}) and the standard deviation chart are given. The \bar{x} control chart monitors the process with respect to both the standard and the variability, while the standard deviation chart monitors its short-term precision. When the calibration process is determined to be in a state of statistical control, the calibrations made at that time may be considered to be valid and the standard deviation of the measurement process may be included in uncertainty calculations using SOP 29.

Note: If a full evaluation of potential bias is desired, it is best if the reference value of the check standard is provided by an outside laboratory accredited to perform the applicable calibrations taking care to ensure that the source calibration has a sufficiently small uncertainty.

3 Equipment

3.1 A check standard is required, and should be constructed of similar materials and design as the standards or devices under calibration. Check standards are maintained in the laboratory as a surrogate for items submitted for routine calibration. Examples of check standards might include a quality assurance reference flask (QARF), a test measure, an extra length

SOP 17 Page 1 of 6

standard, or a stopwatch, taking care that all check standards are sufficiently similar to those submitted for calibration to obtain a suitable standard deviation of the measurement process and similar stability over time.

3.2 All equipment designated in the applicable SOP.

4 Procedure

4.1 Initial Measurements

- 4.1.1 Calibrate the check standard a minimum of 12 times to establish the baseline chart. A calibration is defined as the result of replicate measurements as required by the SOP (i.e., a complete test consists of the number of runs specified in the SOP). Calibrations may be made on successive days, but no two complete tests should be made on any single day. Note: 25 to 30 points are recommended to determine uncertainties.
- 4.1.2 Tabulate the measurement data using the notation and a form such as the one contained in the Appendix of this SOP. The appendix provides an example for duplicate runs only; modifications would be needed when other replicates are used. The data may be maintained in a spreadsheet or other electronic program in lieu of a paper form.
- 4.1.3 A standard deviation may be calculated for each set of runs according to the appropriate SOP. A pooled standard deviation is then determined for the measurement process. When all of the runs have the same number of replicates, each test will have the same number of degrees of freedom and an average of the standard deviations may be used. Use of a standard deviation of replicate measurements is preferred to the use of Range charts (see SOP 20 for guidance on Range charts.)
- 4.2 Construction of Control Charts (See also SOP 9)
 - 4.2.1 Construct the following control charts using the data of section 4.1.
 - 4.2.2Construct an x control chart for a check standard with the following control limits:

(when available)
\bar{x} (mean of the average values)
$\overline{x} - 2s_p$
$\overline{x} - 3s_p$
$\frac{1}{x} + 2s_p$
$\frac{1}{x} + 3s_p$

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4.2.3 Construct a Standard Deviation chart using the same approach. You should use 2 and 3 as the respective multipliers for the Upper Warning Limit and Upper Control Limits. Note that there will be no negative numbers when calculating standard deviations.

4.3 Use of Control Charts

- 4.3.1 An appropriate check standard is calibrated each time the laboratory performs calibrations using the SOP or on a set periodic basis if many calibrations are performed by the laboratory. If the calibrations extend over several days, the check standard is calibrated daily. The values of \bar{x} and s_p for each calibration of the check standard are plotted on the respective charts, preferably in sequential order. The limits on the charts are such that 95 % of the values should fall within the warning limits and rarely should a value fall outside of the control limits, provided that the system is in a state of statistical control.
- 4.3.2 If the plotted value of \bar{x} lies outside of the control limits and the corresponding value on the standard deviation chart is within the control limits, a source of systematic error is suspected.
- 4.3.3 If the values for the standard deviation chart fall outside of the warning limits but inside of the control limits, a decrease in precision is indicated. Other problems should be investigated.
- 4.3.4No calibration data should be accepted when the system is out of control.
- 4.3.5 If the plotted values for either x, or s_p are outside of the warning limits but inside of the control limits, a second set of duplicate calibrations should be made. If the new values are within the warning limits, the process may be considered to be in control. If they lie outside of the warning limits, lack of control is indicated. Investigation actions should be taken and attainment of control demonstrated before calibration measurements are considered to be acceptable.
- 4.3.6Even while the system is in an apparent state of control, incipient troubles may be indicated when the control data show short- or long-term trends, shifts, or runs. The *t*-test and *F*-test may be used to assess the significance of such observations (see NIST IR 6969 Sections 8.9, 8.10, and 8.11).

4.4 Interpretation of Control Chart Data

4.4.1 Demonstration of "in control" status indicates that the calibration process is consistent with the past experience of the laboratory. That is to say, there is no reason to believe that excessive systematic error or changes in precision have occurred.

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- 4.4.2To the extent appropriate, the precision of measurement of the check standard may be extended to the calibration of other standards of similar nominal size made by the same measurement method.
- 4.4.3 Extension of the s_p for the check standard to other calibrations assumes that all aspects of its calibration correspond to those for the other calibration.

Figure 1. Example Control Chart

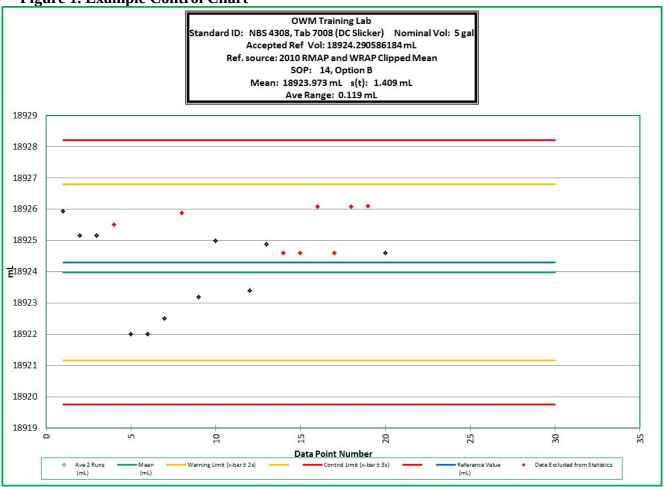
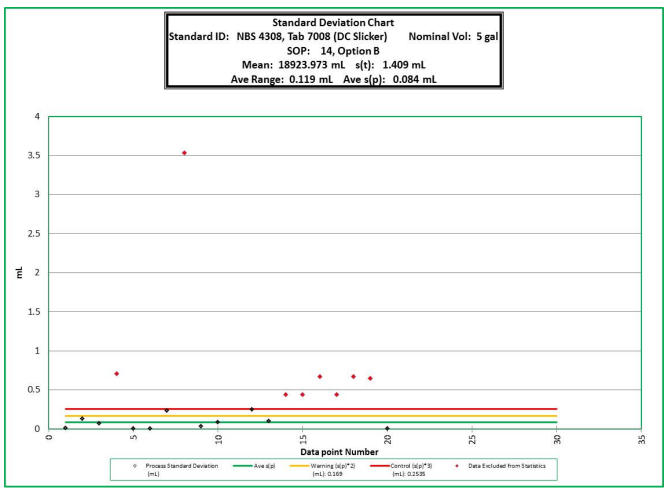


Figure 2. Example Standard Deviation Chart



Note: see point 8 that is outside of control limits.

Appendix Control Chart Check Standard Data

Check Standard Identification	Nominal Value
	Nominal value

Test Number	Date	Run 1	Run 2*	Average of Runs	Standard deviation
1					
2					
3					
4					
5					
6					
7					
8					
9					
10					
11					
12					
13					
14					
15					
16					
17					
18					
19					
20					
21					
22					
23					
24					
25					
26					
27					
28					
29					
30					

^{*} Additional columns will be needed when replicates runs beyond two are used.

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SOP 18

Standard Operating Procedure for Calibration of Graduated Neck-Type Metal Volumetric Field Standards Using a Slicker-Plate Type¹

1 Introduction

1.1 Purpose of Test

This procedure may be used to calibrate small non-pressurized, graduated necktype, metal field standards such as the 5 gal (or 20 L) standards used by weights and measures officials to test liquid dispensing equipment, gasoline pumps, for example. The test measure or prover being calibrated should be evaluated for conformance to appropriate specifications and tolerances (using the checklist provided in NIST Handbook 105-3, Specifications and Tolerances for Graduated Neck Type Volumetric Field Standards, 2010) if being used for legal weights and measures applications. (Alternatively, if requested by the customer, evaluation against OIML R 120 (2010), Standard capacity measures for testing measuring systems for liquids other than water, may be referenced.)

This procedure uses a slicker-plate type standard to fill an unknown test measure of equal nominal volume. It requires water temperature stability during the transfer from the standard to the unknown test measure within the limits shown in Table 1. SOP 19 is a more appropriate procedure when temperature corrections are needed due to lack of water equilibration, temperature differences between the standard and unknown provers, or unstable environments. You may use water that is temperature-equilibrated with the laboratory environment. Equilibration can be achieved by storing the water in clean containers in the laboratory.

Limiting factors: If the limits in Table 1 are exceeded, or materials other than stainless steel and mild steel are used, use SOP 19. These limits are to ensure that the impact of errors resulting from temperature differences on measured values is less than the resolution and repeatability on a 20 L or 5 gal test measure with 1 in³ graduations. If smaller graduations are present, potential errors due to temperature variations must be evaluated further and SOP 19 is recommended.

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¹ Non-SI units are predominately in common use in State legal metrology laboratories, and/or the petroleum industry for many volumetric measurements, therefore non-SI units have been used to reflect the practical needs of the laboratories performing these measurements as appropriate. Most laboratory standards for this calibration procedure are 5 gal "slicker-plate" type standards. Very few laboratories have 20 L "slicker-plate" type standards.

Table 1. Temperature limitations.

Temperature Limitations	Standard Material	Unknown Material	Temperature limit between standards (°C)
1	Stainless steel	Stainless steel	< 0.5 °C
2	Stainless steel	Mild steel	< 0.2 °C

1.2 Prerequisites

- 1.2.1 Verify that the unknown prover has been properly cleaned and vented with all petroleum products removed prior to submission for calibration to ensure laboratory safety.
- 1.2.2 Verify that current calibration certificates with measurement values and uncertainties are available for all of the standards used in the test. All calibration values must have demonstrated metrological traceability to the international system of units (SI). Metrological traceability may be to the SI through a National Metrology Institute such as NIST.
- 1.2.3 Verify that the slicker-plate type standard has sufficiently small standard uncertainties for the intended level of calibration (i.e., less than 0.2 in³ with an approximate 95 % level of confidence).
- 1.2.4 Verify the availability of an adequate supply of clean, preferably soft water (filtered and thermally equilibrated as appropriate) (GLP 10). Water does not need to be distilled or deionized for use in this procedure. The equations used in GLP 10 for the calculation of water density (air saturated) may be used without a significant impact on the measurement results.
- 1.2.5 Verify that the operator has had specific training and is proficient in SOP 18, GMP 3, SOP 17 and is familiar with the operating characteristics and conditioning of the standards used.
- 1.2.6 Verify that the laboratory facilities meet the following minimum conditions to enable meeting the expected uncertainty achievable with this procedure:

Table 2. Laboratory environmental conditions.

Procedure	Temperature	Relative Humidity
Volume transfer	18 °C to 27 °C Stable to ± 2.0 °C / 1 h	35 % to 65 % Stable to ± 20 % / 4 h

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1.3 Field tests

- 1.3.1 A "field" calibration is considered one in which a calibration is conducted in an uncontrolled environment, such as out-of-doors. Calibrations conducted under field and laboratory conditions are not considered equivalent and uncertainties must reflect the conditions of the calibration.
- 1.3.2 SOP 19 is a more suitable procedure for non-laboratory conditions. The care required for field calibrations includes proper safety, a clean and bubble-free water supply, measurement control programs, and a stable temperature environment shaded from direct sunshine to allow the prover, field standard, and clean test liquid (water) to reach an equilibrium temperature with minimal evaporation. Environmental conditions must be selected to be within stated laboratory conditions as shown in Table 2, during the measurements. All data and appropriate environmental conditions must be documented regardless of test location.
- 1.3.3 An increased number of check standard verifications are required to ensure continued suitability of calibration values generated in field conditions as well as to verify the validity of any standards taken out of a secure laboratory environment once the standard(s) are returned to the laboratory.

2 Methodology

2.1 Scope, Precision, Accuracy

This procedure is applicable for the calibration of a small test measure within the limitations of the standards available. The precision attainable depends on the care used in the various volumetric adjustments and readings, in the strict observance of drainage times, and the internal cleanliness of the various volumetric vessels which can influence their drainage characteristics. The accuracy depends on the uncertainties of the calibrations of the standards used.

2.2 Summary

Water is delivered from the standard to the vessel under calibration. Because the "to deliver" volume of the latter is calibrated, the delivery must be into a "wetted-down" vessel. The wet-down also ensures consistent retention in the slicker-plate type standard. The gauge scale of the test vessel is adjusted to a correct reading, as necessary, and then sealed.

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2.3 Standards and Equipment

- 2.3.1 Calibrated slicker-plate type standard made of stainless steel.
- 2.3.2 Calibrated thermometer, with a resolution and uncertainty less than 0.1 °C.
- 2.3.3 Meniscus reading device. (See GMP 3).
- 2.3.4 Timing device (Calibration is not required; uncertainty of the measurement only needs to be less than 5 s for a 30 s pour time.)

2.4 Procedure

- 2.4.1 Cleanliness Verification Fill and drain both standard and vessel to be calibrated and check for any soiling that would affect drainage, as evidenced by clinging droplets, greasy films, and the like. Clean either or both with non-foaming detergent and water, as necessary, and rinse thoroughly. NOTE: Many laboratories have a policy regarding cleanliness of submitted volumetric standards to minimize water contamination with flammable petroleum products. (See GMP 6).
- 2.4.2 Fill the unknown hand-held vessel with water to its nominal level and pour contents during a 30 s ± 5 s period then drain for a 10 s period after cessation of flow. Touch off any adhering drop from the neck. If a stationary test measure is being calibrated, the valve is opened and the measure is emptied, followed by a 30 s drain time after the cessation of the main flow. This constitutes the "wet-down" condition. Filling the vessel from the slicker-plate type standard following the instructions in steps 2.4.3 and 2.4.4 will ensure that both the standard and vessel are properly "wetted-down".
- 2.4.3 Run 1. Fill the slicker-plate standard with water by placing the fill tube or hose inside the standard near the bottom to minimize air bubbles. Fill the standard to slightly higher than the rim as the water level is raised by surface tension. Ensure that the reference standard is level to ensure consistent drain and retention characteristics. Measure and record the temperature in the standard. Use the slicker plate to strike off excess water, checking to see that no air bubbles are entrained in the water during this process.
- 2.4.4 Position the unknown vessel beneath the slicker-plate standard drain valve or piping to be sure no water is lost. Open the valve at the base of the standard at the same time as removing the slicker plate from the top of the standard to transfer water from the standard to the wet-down vessel. Allow a 30 s drain period after cessation of the main flow. Close the valve on the

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- slicker-plate standard and then move the unknown vessel from beneath the drain valve or piping to prevent additional water transfer.
- 2.4.5 Level the test measure and verify that the neck of the unknown test measure is vertical, or perpendicular to a level horizontal plane, by placing a level on the side of the neck as specified in NIST Handbook 105-3 (or suspend by its handle, if appropriate). Read and record the gauge reading. Measure and record the temperature of water in the unknown measure. If the temperature change between the standard and unknown test measure exceeds the allowable limits shown in Table 1, use SOP 19.
- 2.4.6 Adjust the graduated scale of the vessel as described in 3.3 if needed. Seal the scale adjustment device.
- 2.4.7 Run 2. Make a duplicate determination.
- 2.4.8 Replicate runs of the test measure or prover must agree within \pm 0.02 % of the test volume during calibration (\pm 0.2 in³ for 5 gal test measure). (Agreement value is calculated as the difference between Run 1 and Run 2, divided by the nominal volume, and then multiplied by 100.)
- NOTE: If excess disagreement between the runs is observed, check all vessels for cleanliness, leaks, or other damage, identify and correct any problems. Agreement problems may be due to temperature changes, contamination or lack of cleanliness, or poor field conditions, such as when calibration is conducted in an unstable environment. Agreement problems must be corrected before calibration can be completed.

3 Calculations

- 3.1 When the water temperature is reasonably close to 60 °F, the coefficients of expansion of the standard and the test vessel are sufficiently close together, and the deliveries and readings are made over a short period of time, temperature corrections are not made in this procedure. When conditions are not reasonably close to 60 °F and temperature corrections are needed, use SOP 19. Calculations provided in SOP 19 may be required for reporting volumetric results.
- 3.2 Within the accuracy requirements of Table 1, no temperature corrections arising from dissimilarities of the standard and vessel are used. If differences are suspected or observed, use SOP 19.
- 3.3 The reading of Run 1 is used to adjust the scale of the vessel, if necessary, to the correct reading, which is set at the calibrated volume of the slicker-plate standard at 60 °F. Record the adjusted value as the "as left" value. Run 2 will validate the setting. Alternatively, the average of Run 1 and Run 2 may be used with the adjustment made after Run 2. In that case, a validation run should be conducted to ensure correct setting of the gage plate.

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Note: When the accuracy requirements necessitate a temperature correction, the temperature of the water that has been recorded in both the standard and the unknown should be used to perform calculations according to the procedure given in SOP 19.

3.4 Determine and report the volume of the test vessel as follows:

Prover volume = V_s - gauge reading Eqn. 1

where: V_S = Volume of the slicker-plate standard at the appropriate reference temperature (usually 60 °F).

4 Measurement Assurance

- 4.1 Duplicate the process with a suitable check standard. See SOP 17 or SOP 30. Plot the check standard volume and verify that it is within established limits. Alternatively, a *t*-test may be incorporated to check the observed value against an accepted value. The mean of the check standard observations may be used to evaluate bias and drift over time when a reference value for the check standard is available. Check standard observations are used to calculate the standard deviation of the measurement process which contributes to the Type A uncertainty components.
- 4.2 A standard deviation chart is also used for measurement assurance through the evaluation of replicate measurements. The standard deviation of each combination of Run 1 and Run 2 is calculated and the pooled (or average) standard deviation over time may be used to estimate the short-term variability in the measurement process. The short-term standard deviation may be used to incorporate an F-test (observed vs. accepted) into the measurement process. A standard deviation chart for unknown provers represents the variability in condition of test measures submitted for calibration as well as the short-term repeatability of the measurement process.

5 Assignment of Uncertainties

The limits of expanded uncertainty, U, include estimates of the standard uncertainty of the laboratory volumetric standards used, u_s , plus the standard deviation of the process, s_p , at an approximate 95 % level of confidence. See SOP 29 for the complete standard operating procedure for calculating the uncertainty.

5.1 The standard uncertainty for the standard, u_s , is obtained from the calibration report. The combined standard uncertainty, u_c , is used and not the expanded uncertainty, U, therefore the reported uncertainty for the standard will usually need to be divided by the coverage factor k.

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- 5.2 The standard deviation of the measurement process, s_p , is obtained from a control chart for the check standard so that it reflects performance over time (See SOP 17 or 20, and SOP 30). The larger of the value from the standard deviation over time for a check standard from Section 4.1 or from the standard deviation chart taken from Section 4.2 should be used in the uncertainty calculations.
- 5.3 Other standard uncertainties usually included at this calibration level primarily include 1) uncertainties associated with the ability to read the meniscus, only part of which is included in the process variability due to parallax and visual capabilities (See GMP 3), and 2) uncorrected temperature corrections related to the cubical coefficient of expansion for the prover under test, use of proper temperature corrections, and the accuracy of temperature measurements. Additional factors that might be included are: round robin data showing bias, environmental variations over time, and drift of the standard as noted in control charts.
- 5.4 To properly evaluate uncertainties and user requirements (tolerances), assessment of additional user uncertainties may be required by laboratory staff. Through proper use of documented laboratory and field procedures, additional uncertainty factors may be minimized to a level that does not contribute significantly to the previously described factors. Additional standard uncertainties in the calibration of field standards and their use in meter verification may include: how the prover level is established, how delivery and drain times are determined, the use of a proper "wet-down" prior to calibration or use, the cleanliness of the prover and calibration medium, prover retention characteristics related to inside surface, contamination or corrosion, total drain times, and possible air entrapment in the water.
- 5.5 Example components to be considered for an uncertainty budget table are shown in Table 3.

Table 3. Example uncertainty budget table.

Uncertainty Component Description	Symbol	Source	Typical Distribution
Uncertainty of the standard (5.1)	u_s	Calibration report	Rectangular or Normal divided by coverage factor
Accepted standard deviation of the process (5.2)	S_p	Control chart, standard deviation chart	Normal
Ability to read the Meniscus in S (5.3)	u_m	None if using a slicker-plate type standard	Triangular
Ability to read the Meniscus in X (5.3)	u_m	GMP 3	Triangular
Uncorrected temperature errors (5.3)	u_{te}	Calculated based	Rectangular

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		on limits and SOP 19 equations	
Uncertainty of bias or drift of standards (5.3)	u_b	From control chart	Rectangular
Water temperature (S)	u_{ts}	Consider accuracy, resolution, and gradients	Rectangular
Water temperature (X)	u_{tx}	Consider accuracy, resolution, and gradients	Rectangular
Cubical Coefficient of Expansion on S	u_{CCE}	5 % to 10 % (EURAMET CG- 21)	Rectangular
Cubical Coefficient of Expansion on X	u_{CCE}	5 % to 10 % (EURAMET CG- 21)	Rectangular
Uncertainty of drain time	u_d	From experimental data	Rectangular

6 Report

- Report results as described in SOP 1, Preparation of Calibration/Test Results, with the addition of the following:
 - 6.1.1 "To Contain" or "To Deliver" volume, reference temperature, uncertainty, material, thermal coefficient of expansion (assumed or measured), construction, any identifying markings, tolerances (if appropriate), laboratory temperature, water temperature(s) at time of test, barometric pressure, relative humidity, and any out–of-tolerance conditions.

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Appendix AVolume Transfer Data Sheet

Laboratory data and conditions:

zanooratory anta ana contartrons.		
Vessel Owner	Operator	
Vessel ID	Date	
M aterial	Air Temperature	
Cubical Coefficient of Expansion	Relative Humidity	
Standard deviation of the process, s_p	Reference temperature of unknown prover	
s _p from Control Chart or Standard Deviation Chart		
Degrees of Freedom		

Volume standard(s) data:

ID (Note ID of Standards)	Nominal	Volume/Correction	Expanded Unc: From cal. report	Unc: k factor	Cubical Coefficient of Expansion
S					
S					
S					
S					

Observations:

Run	Standard Material (MS/SS)	Unknown Material (MS/SS)	Water Temp in Standard (°C)	Unknown Gauge Reading (in ³)	Water Temp in Unknown (°C)	Adjustment Made?	Temperature Difference Acceptable for SOP 18?
1						YesNo	YesNo
2						YesNo	YesNo
3						YesNo	YesNo

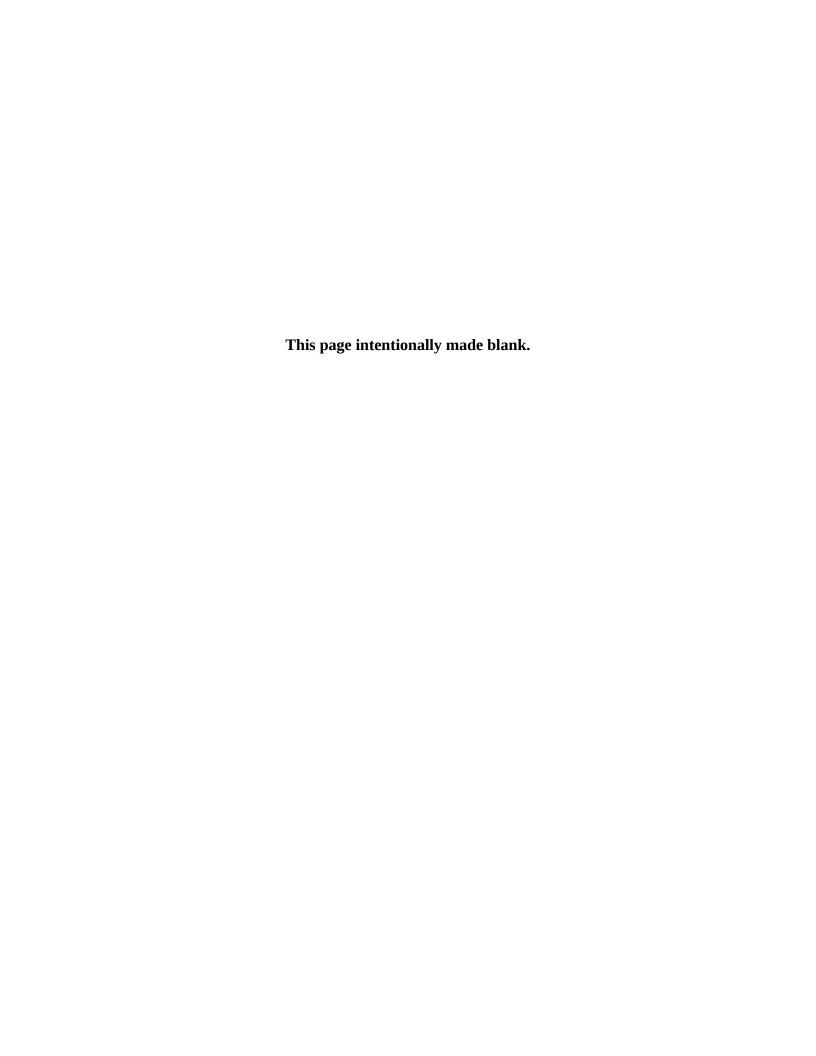
Temperature Difference Evaluation:

If Standard is stainless steel (SS) and Unknown is stainless steel, the temperature of the water between the standard and unknown must not change more than $0.5~^{\circ}$ C during the calibration.

If the standard is stainless steel and the unknown test measure is mild steel (MS), the change in water temperature between standard and unknown must be less than $0.2~\mathrm{^{\circ}C}$ during the calibration.

If these limits are exceeded, use SOP 19 calculations to determine the volume of the unknown volumetric measure.

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SOP 19

Standard Operating Procedure for Calibration of Graduated Neck-Type Metal Provers (Volume Transfer Method)¹

1 Introduction

1.1 Purpose of Test

This procedure is used to calibrate graduated neck type metal test measures and provers (20 L (5 gal) and larger) that are used in verification of petroleum, biodiesel, ethanol, milk, and/or water meters. The test measure or prover being calibrated should be evaluated for conformance to appropriate specifications and tolerances (using the checklist provided in NIST Handbook 105-3, Specifications and Tolerances for Graduated Neck Type Volumetric Field Standards, 2010) if being used for legal weights and measures applications. (Alternatively, if requested by the customer, evaluation against OIML R 120 (2010), Standard capacity measures for testing measuring systems for liquids other than water, may be referenced.)

1.2 Prerequisites

1.2.1 Verify that the unknown prover has been properly cleaned and vented, with all petroleum products removed prior to submission for calibration to ensure laboratory safety and compliance with environmental disposal requirements. The prover may be visually inspected to determine that residual liquid petroleum products are not present. Smell is not necessarily an adequate indicator of cleanliness.

NOTE: Many laboratories have a policy regarding cleanliness of submitted volumetric standards to minimize water contamination with flammable petroleum products.

- 1.2.2 Verify that valid current calibration certificates with measurement values and uncertainties are available for all of the standards used in the test. All calibration values must have demonstrated metrological traceability to the international system of units (SI). Metrological traceability may be to the SI through a National Metrology Institute such as NIST.
- 1.2.3 Verify that the standards to be used have sufficiently small standard uncertainties for the intended level of calibration.

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¹ Non-SI units are predominately in common use in State legal metrology laboratories, and/or the petroleum industry for many volumetric measurements, therefore non-SI units have been used to reflect the practical needs of the laboratories performing these measurements as appropriate.

- 1.2.4 Verify the availability of an adequate supply of clean, preferably soft water (filtered and thermally equilibrated as appropriate) water (GLP 10). Water does not need to be distilled or deionized for use in this procedure. The equations used in GLP 10 for the calculation of water density (air saturated) may be used without a significant impact on the measurement results.
- 1.2.5 Verify that the operator has had specific training and is proficient in SOP 17, SOP 19, SOP 20, GMP 3, and is familiar with the operating characteristics and conditioning of the standards used.
- 1.2.6 Verify that the laboratory facilities meet the following minimum conditions to make possible the expected uncertainty achievable with this procedure:

Table 1. Laboratory environmental conditions.

Procedure	Temperature	Relative Humidity
Volume	18 °C to 27 °C	35 % to 65 %
Transfer	Stable to ± 2.0 °C / 1 h	Stable to ± 20 % / 4 h

1.3 Field tests

- 1.3.1 A "field" calibration is considered one in which a calibration is conducted in uncontrolled environments, such as out-of-doors. Calibrations conducted under field and laboratory conditions are *not* considered equivalent.
- 1.3.2 The care required for field calibrations includes proper safety, a clean and bubble-free water supply, measurement control programs, and a stable temperature environment shaded from direct sunshine to allow the prover, field standard, and clean test liquid (water) to reach an equilibrium temperature with minimal evaporation. Environmental conditions must be selected to be within stated laboratory conditions during the measurements. All data and appropriate environmental conditions must be documented regardless of test location.
- 1.3.3 An increased number of check standard verifications are required to ensure continued suitability of calibration values generated in field conditions as well as to verify the validity of any standards taken out of a secure laboratory environment once the standard(s) are returned to the laboratory.

2 Methodology

2.1 Scope, Precision, Accuracy

This procedure is applicable for the calibration of any size metal prover within the limitations of the standards available. The repeatability attainable depends on strict adherence to the procedure, the care in volumetric adjustments, and the number of transfers, in the case of multiple transfers. The accuracy depends on the standards used.

2.2 Summary

Water is delivered from calibrated volumetric standard(s) into the unknown test measure or prover being calibrated. Depending on the respective volumes, multiple transfers may be required. While these should be minimized, a maximum number of 15 transfers are permitted to ensure that final uncertainties and systematic errors are sufficiently small for the intended applications. The temperature of the calibration medium (water) cannot be considered to be constant during transfers; hence, the temperature of the water for each transfer must be measured. Because of the large volumes, the difference in thermal expansion of the respective vessels must be considered.

2.3 Standards and Equipment

- 2.3.1 Calibrated volumetric standards of suitable sizes.
- 2.3.2 Calibrated flask(s) of suitable sizes to calibrate neck of prover.
- 2.3.3 A funnel for transferring water from the flask into the prover or test measure.
- 2.3.4 Meniscus reading device (See GMP 3).
- 2.3.5 Calibrated thermometer, with resolution and uncertainty less than 0.1 °C.
- 2.3.6 Timing device (calibration is not required; uncertainty of the measurement only needs to be less than 5 s for a 30 s pour time.)
- 2.3.7 Sturdy platform, with appropriate safety conditions, with sufficient height to hold standard and to permit transfer of water from it to the prover by gravity flow.
- 2.3.8 Clean pipe or tubing (hoses) to facilitate transfer of water from the laboratory standard to prover. Pipe and hose lengths must be minimized to reduce water retention errors. Care must be taken during wet-downs and runs to ensure complete drainage and consistent retention in all hoses or pipes.

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2.4 Procedure

2.4.1 Cleanliness verification

Fill and drain both the standard and unknown test measure or prover to be calibrated and check for visual evidence of soiling and of improper drainage. If necessary, clean with non-foaming detergent and water and rinse thoroughly (see GMP 6).

- 2.4.2 Neck scale plate verification. (This is generally only conducted for new or damaged measures or those that have not been calibrated by the laboratory in the past.)
 - 2.4.2.1 Fill the unknown prover with water from the standard. Adjust and check the prover level by placing a precision spirit or electronic digital level vertically on the neck on at least two locations, 90 degrees apart around the circumference of the neck and adjust the orientation of the standard until the neck is as close to vertical (plumb or perpendicular to the horizontal plane) as possible. Verify and adjust any mounted levels that are on the prover to agree (when present and when possible). Check the prover system for leaks. This neck scale plate verification may be used as a wetted-down run.
 - 2.4.2.2 Bleed the liquid level down to a graduation near the bottom of the upper neck. "rock" the prover to "bounce" the liquid level, momentarily, to ensure that it has reached an equilibrium level. Read and record this setting to be used as the initial scale reading sr_i for verification of the neck scale plate.
 - 2.4.2.3 Recheck the scale reading, and then add water from calibrated standards equal to approximately $^{1}/_{4}$ or $^{1}/_{5}$ of the graduated neck volume and record the scale reading.
 - 2.4.2.4 Repeat 2.4.2.3 by successive additions until water is near the top of the scale. Record scale readings after each addition. The last reading will be the final scale reading, sr_f . The closer the water is to the top of the neck, the harder it may be to "bounce" the liquid in the gauge.
 - 2.4.2.5 A plot of scale readings with respect to the total volume of water that is added V_w should be linear and will be a gross check of the validity of this calibration.
 - 2.4.2.6 Calculate and assess the accuracy of the neck scale for each interval. The error should be less than 0.5 % of the graduated neck

volume or $^{1}/_{4}$ of a graduation (whichever is smaller) per NIST Handbook 105-3. If the error is greater than this, the scale should be replaced. Uncertainties associated with the neck scale calibration are primarily related to the setting of the meniscus on the flask used (with each drop), the reading of the meniscus on the unknown prover (at each volume reading), and the calibration uncertainty of the flask. These uncertainties must be considered when evaluating the acceptability of the neck scale. Alternatively, a Neck Scale Correction Value (NSCV) may be issued with instructions to the user if it is anticipated that this correction value will be used

- 2.4.2.7 Neck scale errors less than 0.5 % of the graduated neck volume should be included in the uncertainty estimate associated with the prover calibration.
- 2.4.2.8 The neck scale correction value is calculated as follows:

$$NSCV = \frac{V_{w}}{\left(sr_{f} - sr_{i}\right)}$$
 Eqn. 1

Table 2. Variables for neck scale correction value equation.

NSCV	Neck scale correction value
Vw	Total volume of water added to neck
sr_f	Scale reading, final
Sr_i	Scale reading, initial

2.4.3 Body Calibration

2.4.3.1 Fill the standard with water and then transfer the water into the unknown prover establishing a wet down of the standard. Wait 30 s after the cessation of the main flow before closing the drain valve on the standard. Level the unknown prover, and drain the water, again waiting 30 s after cessation of the main flow, before closing the drain valve. This establishes a "wetted-down" condition for provers with no bottom zero. If the unknown test vessel is a 5 gal (20 L) hand-held test measure, fill the unknown vessel with water to its nominal mark and empty it using a 30 s (± 5 s) pour followed by a 10 s drain after cessation of the main flow to establish the wetted-down condition.

If a bottom zero is present, follow the guidance provided in SOP 21 for LPG provers as follows: When the liquid reaches the top of the lower gauge glass, close the valve and allow the water to drain

from the interior of the prover into the lower neck for 30 s. Then bleed slowly with the bleed valve (4) until the bottom of the liquid meniscus reaches the zero graduation. (This step should be started during the 30 s drain period but should not be completed before the end of the drain period.)

Alternatively, the prover may be completely drained with a 30 s drain time after the cessation of the main flow, and then refilled with a funnel and small volume of water to set the zero mark (which will add to the prover calibration uncertainty due to variable retention characteristics).

- 2.4.3.2 Run 1. Fill the standard and measure and record the temperature.
 - 2.4.3.2.1 Measure and record the temperature of the water in the standard, t_1 , then adjust the standard prover to its reference mark or record the neck reading, and then discharge into the unknown prover. Wait 30 s after cessation of the main flow to attain specified drainage, and then close the delivery valve. Remove any hoses or pipes to prevent additional water transfer.
 - 2.4.3.2.2 Repeat step 2.4.3.2.1 as many times as necessary (note the 15-drop limit) to fill the unknown prover to its nominal volume. Verify the level condition and record the temperature of water in the standard for each drop, t_1 to t_N .
 - 2.4.3.2.3 Level the filled unknown prover. Check the prover level by placing a precision spirit or electronic digital level vertically on the neck on at least two locations, 90 degrees apart around the circumference of the neck and adjust the orientation of the unknown measure until the neck is as close to vertical (plumb or perpendicular to the horizontal plane) as possible. Verify and adjust any mounted levels that are on the prover to agree (when present and when possible). Read and record the scale plate (gauge) reading.
 - 2.4.3.2.4 Measure the temperature of the water in the unknown prover, t_x , and record. For test measures without thermometer wells, the temperature should be taken as close as possible to the center (vertically and horizontally) of the cylinder of the test measure main body (and not in the neck). For larger provers, and when thermometer wells are present, the average temperature, calculated from temperatures taken at

multiple locations from within the unknown prover should be used. (Alternatively, if a prover has mounted thermometers, the internal temperature from multiple locations within the prover may be used).

- 2.4.3.2.5 Perform the calculations described in section 3 to determine the prover volume at the appropriate reference temperature.
- 2.4.3.4 Adjust the scale as needed. If adjusted, record the adjusted prover gauge reading for determining the "as left" value for Run 1. Run 2 will validate the setting. Alternatively, the average of Run 1 and Run 2 may be used with the adjustment made after Run 2. In that case, a validation run is required to ensure correct setting of the scale plate.
- 2.4.3.5 Run 2 Repeat the process described in 2.4.3.2. Replicate runs of the test measure or prover must agree within \pm 0.02 % of the test volume during calibration. (Agreement value is calculated as the difference between Run 1 and Run 2, divided by the nominal volume, and then multiplied by 100.)
- NOTE: If excess disagreement between replicated measurements is observed, check all vessels for cleanliness, leaks, drain lines, additional valve, or damage, identifying and correcting any problems. Lack of measurement agreement may be due to contamination, lack of cleanliness, excessive temperature changes, poor laboratory conditions, or poor field conditions, such as when calibration is conducted in an unstable environment. Repeatability problems must be corrected before calibration can be completed.
- 2.4.3.7 Seal the equipment as specified in the laboratory policy.

3 Calculations

The following calculations assume that the standard was calibrated using a reference temperature of 60 °F (15.56 °C) and that you are calibrating a field standard to a reference temperature of 60 °F (15.56 °C). Equations for situations where different reference temperatures are involved follows in Sections 3.4 to 3.6.

3.1 Single Delivery

3.1.1 Calculate V_{X60} , the volume of the unknown prover at 60 °F, using the following equation:

$$V_{X60} = \frac{\rho_1 \left\{ \left(V_{S60} + \Delta_1 \right) \left[1 + \alpha \left(t_1 - 60 \, ^{\circ} \text{F} \right) \right] \right\}}{\rho_x \left[1 + \beta \left(t_x - 60 \, ^{\circ} \text{F} \right) \right]}$$
Eqn. 2

3.2 Multiple Deliveries

3.2.1 Calculate V_{X60} , the volume of the unknown prover at 60 °F, using the following equation:

$$V_{x60} = \frac{\rho_1 \left\{ \left(V_{s60} + \Delta_1 \right) \left[1 + \alpha \left(t_1 - 60 \, ^{\circ} \mathrm{F} \right) \right] \right\} + \rho_2 \left\{ \left(V_{s60} + \Delta_2 \right) \left[1 + \alpha \left(t_2 - 60 \, ^{\circ} \mathrm{F} \right) \right] \right\} + \dots + \rho_N \left\{ \left(V_{s60} + \Delta_N \right) \left[1 + \alpha \left(t_N - 60 \, ^{\circ} \mathrm{F} \right) \right] \right\}}{\rho_x \left[1 + \beta \left(t_x - 60 \, ^{\circ} \mathrm{F} \right) \right]} \quad \text{Eqn. 3}$$

Table 3. Variables for V_{X60} equations.

Tuble 5. Variation	Symbols Used in Equations				
	Symbol Cott in Equations				
V_{X60}	volume of the unknown vessel at 60 °F				
V_{S60}	volume of the standard vessel at 60 °F				
$ ho_1, ho_2,, ho_N$	density of the water in the standard prover where ρ_I is the density of the water for the first delivery, ρ_2 is the density of the water for the second delivery, and so on until all N deliveries are completed				
$\Delta_1,\Delta_2,,\Delta_N$	volume difference between water level and the reference mark on the standard where the subscripts $1, 2,, N$, represent each delivery as above. If the water level is below the reference line, Δ is negative. If the water level is above the reference line, Δ is positive. If the water level is at the reference line, Δ is zero NOTE: units must match volume units for the standard. The Δ is zero for slicker-plate type standards.				
$t_1, t_2,, t_N$	temperature of water for each delivery with the subscripts as above				
α	coefficient of cubical expansion for the standard in units / °F				
β	coefficient of cubical expansion for the prover in units / °F				
t_{x}	temperature of the water in the filled unknown vessel in units °F				
$\rho_{\scriptscriptstyle X}$	density of the water in the unknown vessel in g/cm ³				
Note: Values f	Note: Values for the density of water may be calculated from the equations given in GLP 10.				

3.3 Prover Error/Correction or Deviation From Nominal

The total calculated volume of the prover at its reference temperature should be reported on the calibration report. The SI unit of volume is m³, so a conversion factor is to be included on the report in the notes section when other volume units are used.

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The prover volume for an open neck prover equals the V_{x60} value minus the gauge reading that is the difference from the nominal volume (with matched units).

Prover volume = V_{x60} - gauge reading Eqn. 4

 $Prover\ error = Prover\ volume\ -V_{Nom}$ Eqn. 5

Prover error = V_{X60} - gauge reading - V_{Nom} Eqn. 6

where:

 V_{Nom} = Nominal Volume (taking care to match units)

 V_{X60} is the calculated volume of water that should be observed in the prover. A positive prover error means that the prover is larger than nominal. A negative prover error means that the prover is smaller than nominal.

- Example 1: If V_{X60} is 100.02 gal and gauge reading is 0.02 gal (above nominal); then the prover volume at nominal is 100.00 gal; and the prover error and correction are 0; and no adjustment is needed.
- Example 2: If V_{X60} is 100.02 gal and gauge reading is -0.02 gal (below nominal); then the prover volume at nominal is 100.04 gal; the prover error is + 0.04 gal; and to adjust the prover, set the gauge to read 0.02 gal (the volume level will show a gauge reading of 0.02 gal, which is 4.62 in³ or about 5 in³, above nominal.)

3.4 Alternative Reference Temperatures

3.4.1 Reference temperatures other than 60 °F (15.56 °C) may occasionally be used. Common reference temperatures for other liquids follow, however not all of them may be suitable for use with standards calibrated using this procedure:

Commodity	Reference Temperature
Frozen food labeled by volume (e.g., fruit juice)	-18 °C (0 °F)
Beer	3.9 °C (39.1 °F)
Food that must be kept refrigerated (e.g., milk)	4.4 °C (40 °F)
Distilled spirits or petroleum	15.56 °C (60 °F)
Petroleum (International Reference)	15 °C (59 °F)
Compressed Natural Gas (CNG)	15 °C (60 °F)
Wine	20 °C (68 °F)
Unrefrigerated liquids (e.g., sold unchilled, like soft drinks)	20 °C (68 °F)
Hydrogen (H ₂)	21 °C (70 °F)
Petroleum (Hawaii)	26.67 °C (80 °F)

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Equations for calculations when using alternative reference temperatures follow:

3.5 Single Delivery

3.5.1 Calculate V_{Xtref} , the volume of the unknown prover at its designated reference temperature (°F), using the following equation:

$$V_{Xtref} = \frac{\rho_1 \left\{ \left(V_{Stref} + \Delta_1 \right) \left[1 + \alpha \left(t_1 - t_{ref S} \right) \right] \right\}}{\rho_x \left[1 + \beta \left(t_x - t_{ref X} \right) \right]}$$
Eqn. 6

3.6 Multiple Deliveries

3.5.1 Calculate V_{Xtref} , the volume of the unknown prover at its designated reference temperature, using the following equation:

$$V_{Xtref_{X}} = \frac{\rho_{1} \left\{ \left(V_{Stref_{S}} + \Delta_{1}\right) \left[1 + \alpha_{1} \left(t_{1} - t_{\text{ref S}}\right)\right] \right\} + \rho_{2} \left\{ \left(V_{Stref_{S}} + \Delta_{2}\right) \left[1 + \alpha_{2} \left(t_{2} - t_{\text{ref S}}\right)\right] \right\} + \dots + \rho_{N} \left\{ \left(V_{Stref_{S}} + \Delta_{N}\right) \left[1 + \alpha_{N} \left(t_{N} - t_{\text{ref S}}\right)\right] \right\} + \dots + \rho_{N} \left\{ \left(V_{Stref_{S}} + \Delta_{N}\right) \left[1 + \alpha_{N} \left(t_{N} - t_{\text{ref S}}\right)\right] \right\} + \dots + \rho_{N} \left\{ \left(V_{Stref_{S}} + \Delta_{N}\right) \left[1 + \alpha_{N} \left(t_{N} - t_{\text{ref S}}\right)\right] \right\} + \dots + \rho_{N} \left\{ \left(V_{Stref_{S}} + \Delta_{N}\right) \left[1 + \alpha_{N} \left(t_{N} - t_{\text{ref S}}\right)\right] \right\} + \dots + \rho_{N} \left\{ \left(V_{Stref_{S}} + \Delta_{N}\right) \left[1 + \alpha_{N} \left(t_{N} - t_{\text{ref S}}\right)\right] \right\} + \dots + \rho_{N} \left\{ \left(V_{Stref_{S}} + \Delta_{N}\right) \left[1 + \alpha_{N} \left(t_{N} - t_{\text{ref S}}\right)\right] \right\} + \dots + \rho_{N} \left\{ \left(V_{Stref_{S}} + \Delta_{N}\right) \left[1 + \alpha_{N} \left(t_{N} - t_{\text{ref S}}\right)\right] \right\} + \dots + \rho_{N} \left\{ \left(V_{Stref_{S}} + \Delta_{N}\right) \left[1 + \alpha_{N} \left(t_{N} - t_{\text{ref S}}\right)\right] \right\} + \dots + \rho_{N} \left\{ \left(V_{Stref_{S}} + \Delta_{N}\right) \left[1 + \alpha_{N} \left(t_{N} - t_{\text{ref S}}\right)\right] \right\} + \dots + \rho_{N} \left\{ \left(V_{Stref_{S}} + \Delta_{N}\right) \left[1 + \alpha_{N} \left(t_{N} - t_{\text{ref S}}\right)\right] \right\} + \dots + \rho_{N} \left\{ \left(V_{Stref_{S}} + \Delta_{N}\right) \left[1 + \alpha_{N} \left(t_{N} - t_{\text{ref S}}\right)\right] \right\} + \dots + \rho_{N} \left\{ \left(V_{Stref_{S}} + \Delta_{N}\right) \left[1 + \alpha_{N} \left(t_{N} - t_{\text{ref S}}\right)\right] \right\} + \dots + \rho_{N} \left\{ \left(V_{Stref_{S}} + \Delta_{N}\right) \left[1 + \alpha_{N} \left(t_{N} - t_{\text{ref S}}\right)\right] \right\} + \dots + \rho_{N} \left\{ \left(V_{Stref_{S}} + \Delta_{N}\right) \left[1 + \alpha_{N} \left(t_{N} - t_{\text{ref S}}\right)\right] \right\} + \dots + \rho_{N} \left\{ \left(V_{Stref_{S}} + \Delta_{N}\right) \left[1 + \alpha_{N} \left(t_{N} - t_{\text{ref S}}\right)\right] \right\} + \dots + \rho_{N} \left\{ \left(V_{Stref_{S}} + \Delta_{N}\right) \left[1 + \alpha_{N} \left(t_{N} - t_{\text{ref S}}\right)\right] \right\} + \dots + \rho_{N} \left\{ \left(V_{Stref_{S}} + \Delta_{N}\right) \left[1 + \alpha_{N} \left(t_{N} - t_{\text{ref S}}\right)\right] \right\} + \dots + \rho_{N} \left\{ \left(V_{Stref_{S}} + \Delta_{N}\right) \left[1 + \alpha_{N} \left(t_{N} - t_{\text{ref S}}\right)\right] \right\} + \dots + \rho_{N} \left\{ \left(V_{Stref_{S}} + \Delta_{N}\right) \left[1 + \alpha_{N} \left(t_{N} - t_{\text{ref S}}\right)\right] \right\} + \dots + \rho_{N} \left\{ \left(V_{Stref_{S}} + \Delta_{N}\right) \left[1 + \alpha_{N} \left(t_{N} - t_{\text{ref S}}\right)\right] \right\} + \dots + \rho_{N} \left\{ \left(V_{Stref_{S}} + \Delta_{N}\right) \left[1 + \alpha_{N} \left(t_{N} - t_{\text{ref S}}\right)\right] \right\} + \dots + \rho_{N} \left\{ \left(V_{Stref_{S}} + \Delta_{N}\right) \left[1 + \alpha_{N} \left(t_{N} - t_{\text{ref S}}\right)\right] \right\} + \dots + \rho_{N} \left\{ \left(V_{Stref_{S}} + \Delta_{N}\right) \left[1$$

Eqn. 7

Table 3A. Variables for V_{XtrefX} equations.

	Symbols Used in Equations			
V_{XtrefX}	volume of the unknown vessel, V_X at its designated reference temperature, t_{ref}			
V_{StrefS}	volume of the standard vessel, V_S at its designated reference temperature, t_{refS}			
$\rho_1, \rho_2,, \rho_N$	density of the water in the standard where ρ_I is the density of the water for the first delivery, ρ_2 is the density of the water for the second delivery, and so on until all N deliveries are completed			
$\Delta_1, \Delta_2,, \Delta_N$	volume difference between water level and the reference mark on the standard where the subscripts $1, 2,, N$, represent each delivery as above. If the water level is below the reference line, Δ is negative. If the water level is above the reference line, Δ is positive. If the water level is at the reference line, Δ is zero. NOTE: units must match volume units for the standard. The Δ is zero for slicker-plate type standards.			
$t_1, t_2,, t_N$	temperature of water for each delivery with the subscripts as above			
α	coefficient of cubical expansion for the standard in its designated units			
β	coefficient of cubical expansion for the prover in its designated units			
$t_{\scriptscriptstyle X}$	temperature of the water in the filled unknown vessel in designated units			
$\rho_{\scriptscriptstyle X}$	density of the water in the prover in g/cm ³			

Notes: Values for the density of water may be calculated from the equations given in GLP 10. The cubical coefficient of the materials must match the unit assigned to the temperature measurement.

4 Measurement Assurance

- 4.1. Duplicate the process with a suitable check standard or have a suitable range of check standards for the laboratory. See SOP 17, SOP 20 and SOP 30. Plot the check standard volume and verify it is within established limits OR a *t*-test may be incorporated to check the observed value against an accepted value. The mean of the check standard observations is used to evaluate bias and drift over time. Check standard observations can be used to calculate the standard deviation of the measurement process which contributes to the Type A uncertainty components.
- 4.2. If a standard deviation chart is used for measurement assurance, the standard deviation of each combination of Run 1 and Run 2 is calculated and the pooled (or average) standard deviation is used as the estimate of variability in the measurement process. Note: the pooled or average standard deviation over time reflects varying conditions of test items that are submitted to the laboratory, but may not reflect potential meniscus reading errors (See GMP 3.)

5 Assignment of Uncertainties

- 5.1 The limits of expanded uncertainty, U, include estimates of the standard uncertainty of the laboratory volumetric standards used, u_s , plus the standard deviation of the process, s_p , at the 95 % level of confidence. See SOP 29 for the complete standard operating procedure for calculating the uncertainty.
 - 5.1.1 The standard uncertainty for the standard, u_s , is obtained from the calibration report. The combined standard uncertainty, u_c , is used and not the expanded uncertainty, U, therefore the reported uncertainty for the standard will usually need to be divided by the coverage factor k. See SOP 29 for the complete standard operating procedure for calculating the uncertainty when multiple deliveries or multiple standards are used to ensure correct handling of correlated uncertainties. Fifteen is the maximum recommended number of deliveries from a laboratory standard to a prover under test to minimize calibration uncertainties to the levels identified previously.
- 5.2 The standard deviation of the measurement process, s_p, is taken from a control chart for a check standard or from standard deviation charts from provers of similar size (See SOP 17, SOP 20 and SOP 30). The larger of the value from the standard deviation over time for a check standard or from the standard deviation chart should be used in the uncertainty calculations. If a check standard is available, it is possible to evaluate the presence of bias in the measurement process.
- 5.3 Neck calibration uncertainty should be estimated based on the uncertainty of standards used, errors observed during calibration, ability to read the meniscus of all standards involved (see GMP 3) and the repeatability of the neck calibration.

- 5.4 Other standard uncertainties usually included at this calibration level primarily include 1) uncertainties associated with the ability to read the meniscus, only part of which is included in the process variability due to parallax and visual capabilities, and 2) uncertainties associated with temperature corrections that include values for the cubical coefficient of expansion for the prover under test, the accuracy and gradients associated with temperature measurements in test measures or provers. Additional factors that might be included are: round robin data showing reproducibility, environmental variations over time, and bias or drift of the standard as noted in control charts.
- 5.5 To properly evaluate uncertainties and user requirements (tolerances), assessment of additional user uncertainties may be required by laboratory staff. Through proper use of documented laboratory and field procedures, additional uncertainty factors may be minimized to a level that does not contribute significantly to the previously described factors. Additional standard uncertainties in the calibration of field standards and their use in meter verification may include: how the prover level is established, how delivery and drain times are determined, the use of a proper "wetted-down" prior to calibration or use, whether gravity drain is used during calibration or whether the volume of water is eliminated by pumping, the cleanliness of the prover and calibration medium, prover retention characteristics related to inside surface, contamination or corrosion, and total drain times, and possible air entrapment in the water, and connecting pipes. Systematic errors may be observed between laboratory calibration practices where a gravity drain is used and field use where the pumping system is used.
- 5.6 Example components to be considered for an uncertainty budget table are shown in Table 4. Multiple values of some items may need to be considered (e.g., multiple drops from the standard, multiple meniscus readings, and multiple temperature readings.)

Table 4. Example Uncertainty Budget Table.

Uncertainty Component Description	Symbol	Source	Typical Distribution
Uncertainty of the standard (5.1)	u_s	Calibration report; may be multiplied or added based on dependencies	Rectangular or Normal divided by coverage factor
Accepted standard deviation of the process (5.2)	S_p	Control chart, standard deviation chart	Normal
Uncertainty or uncorrected error associated with a neck calibration (5.3)	u_n	From experimental data	Rectangular
Ability to read the Meniscus in S (5.4)	u_m	None if using a slicker- plate type standard; GMP 3	Triangular
Ability to read the Meniscus in X (5.4)	u_m	GMP 3	Triangular

Water temperature (S) (5.4)	u_{ts}	Consider accuracy, resolution, and gradients	Rectangular
Water temperature (X) (5.4)	u_{tx}	Consider accuracy, resolution, and gradients	Rectangular
Cubical Coefficient of Expansion on S (5.4)	u_{CCE}	5 % to 10 % (EURAMET CG-21)	Rectangular
Cubical Coefficient of Expansion on X (5.4)	u_{CCE}	5 % to 10 % (EURAMET CG-21)	Rectangular
Uncertainty of bias or drift of standards (5.2)	u_b	From control chart	Rectangular
Uncertainty of drain time	u_d	From experimental data	Normal

6 Report

Report results as described in SOP 1, Preparation of Calibration/Test Results, with the addition of the following:

"To Contain" or "To Deliver" prover volume, reference temperature, uncertainty, material, thermal coefficient of expansion (assumed or measured), construction, any identifying markings, tolerances (if appropriate), laboratory temperature, water temperature(s) at time of test, barometric pressure, relative humidity, and any out—of-tolerance conditions. Appendix B, Example Temperature Correction Chart may be provided as a supplement to the calibration report to encourage appropriate volumetric corrections of the provers during routine use.

Additional References:

Bean, V. E., Espina, P. I., Wright, J. D., Houser, J. F., Sheckels, S. D., and Johnson, A. N., NIST Calibration Services for Liquid Volume, NIST Special Publication 250-72, National Institute of Standards and Technology, Gaithersburg, MD, (2009).

EURAMET Calibration Guide 21, Guidelines on the Calibration of Standard Capacity Measures Using the Volumetric Method (Version 1.0, 04/2013).

Appendix AVolume Transfer Data Sheet

Laboratory data and conditions:

Vessel Owner	Operator	
Vessel ID	Date	
Nominal Volume	Air Temperature	
M aterial	Relative Humidity	
Cubical Coefficient of Expansion	Standard deviation of the process, s_p from Control Chart	
Reference temperature of unknown prover	$\begin{array}{c} \text{Standard deviation of the process, } s_p \\ \text{from the Standard Deviation Chart} \end{array}$	
Unknown prover graduations	Degrees of Freedom	
Applicable specifications and tolerances		

Volume standard(s) data:

ID (Note ID of Standards)	Nominal Volume	Volume/Correction	Expanded Unc: From cal. report	Unc: k factor	Cubical Coefficient of Expansion
S					
S					
S					
S					

Run 1: Data for volumes delivered from the standards:

DROP #	Reported Volume (gal)	M aterial (M S/SS/BS/SL)	Water Temp (°C) (Must be ≥ 0.5 °C and < 40 °C)	Gauge Delta (in³)
1				
2				
3				
4				
5				
6				
7				
8				
9				
10				
11				
12				
13				
14				
15				

Run 1: Data from the filled unknown field standard:

Material for the Unknown:	MS, SS, TP, AL, PVLC
Final Gauge Reading:	in^3
Final Water Temperature:	°C

Run 2: Data for volumes delivered from the standards:

DROP #	Reported Volume (gal)	Material (MS/SS/BS/SL)	Water Temp (°C) $(Must be \ge 0.5 °C and < 40 °C)$	Gauge Delta (in³)
1				
2				
3				
4				
5				
6				
7				
8				
9				
10				
11				
12				
13				
14				
15				

Run 2: Data from the filled unknown field standard:

Material for the Unknown:	MS, SS, TP, AL, PVLC
Final Gauge Reading:	in^3
Final Water Temperature:	°C

Appendix B
Example Temperature Correction Table*

	Example for a 100 gallo	on prover and 60 °F ref				
CCF	CCE Mild Steel Stainless Steel					
CCL	1.86 x 10-5/°F			10-5/°F		
Temperature °F	in ³	gal	in ³	gal		
-20	-34	-0.149	-49	-0.212		
-15	-32	-0.139	-46	-0.199		
-10	-30	-0.130	-43	-0.186		
-5	-28	-0.121	-40	-0.172		
0	-26	-0.112	-37	-0.159		
5	-24	-0.102	-34	-0.146		
10	-21	-0.093	-31	-0.133		
15	-19	-0.084	-28	-0.119		
20	-17	-0.074	-24	-0.106		
25	-15	-0.065	-21	-0.093		
30	-13	-0.056	-18	-0.079		
35	-11	-0.047	-15	-0.066		
40	-9	-0.037	-12	-0.053		
45	-6	-0.028	-9	-0.040		
50	-4	-0.019	-6	-0.026		
55	-2	-0.009	-3	-0.013		
60	0	0.000	0	0.000		
65	2	0.009	3	0.013		
70	4	0.019	6	0.026		
75	6	0.028	9	0.040		
80	9	0.037	12	0.053		
85	11	0.046	15	0.066		
90	13	0.056	18	0.080		
95	15	0.065	21	0.093		
100	17	0.074	24	0.106		
105	19	0.084	28	0.119		
110	21	0.093	31	0.133		
115	24	0.102	34	0.146		
120	26	0.112	37	0.159		
CCE = coefficient of cul	bical expansion					

^{*}Provide only the applicable coefficient of cubical expansion for the prover under test.

SOP 20

Standard Operating Procedure for Standard Deviation and Range Charts

1 Introduction

1.1 Purpose

This procedure describes a process to be followed to monitor the statistical control of a measurement process using standard deviation charts or range charts for any calibration method where replicate measurements are performed and where it is not practical or feasible to maintain laboratory check standards. Standard deviation charts are preferred to the use of range charts for monitoring and evaluation process standard deviations when replicate measurements are made.

1.2 Prerequisites

The procedure must match the calibration procedure that is being monitored.

2 Summary

Because of the size and cost of some laboratory standards, it is not always practical to have check standards remain in the laboratory for the purposes of measurement control. However, it is practical to maintain standard deviation (s_p) charts or range (R) charts for each nominal size standard to evaluate the standard deviation of replicate runs. Directions for preparing and using an R control chart that monitors the precision of the test procedure are given. It is assumed that standards of the same nominal capacity and design will have similar characteristics with respect to the repeatability of tests. Since it is not practical to run a sufficient number of tests on each unknown standard to determine the repeatability, the absolute difference between replicate tests are graphed on the same s_p or R chart to reflect the repeatability of measurement of the unknown standards tested in the laboratory. Note: Provers of similar readability may be grouped together on the same chart as long as results from different sized provers pass an F-test (e.g., provers greater than and equal to 500 gal generally have 20 in 3 graduations and equivalent repeatability.)

3 Equipment

All equipment is designated in the respective calibration SOP.

4 Procedure

4.1 Data Collection

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- 4.1.1 Conduct the required number of replicates per test on the unknown test item per the designated SOP. A minimum of 12 sets of data must be available before a reasonably adequate data base is established. Note: 25 to 30 replicate sets of data are recommended to determine valid uncertainties.
- 4.1.2 Tabulate the measured errors as determined by each of the two trials using a form such as the one contained in the Appendix. The data may be maintained in a spreadsheet or other electronic program in lieu of a paper form. (If the unknown item is adjusted after the first run, the first trial reading is evaluated after the adjustment.)
- 4.2 Calculate the standard deviation (preferred)
 - 4.2.1. A standard deviation of the replicate runs, s_p, may be calculated for each set of runs according to the appropriate SOP. An accepted standard deviation over time can be calculated as the pooled standard deviation to represent the accepted standard deviation of the measurement process.
- 4.3 Calculate the range (optional)
 - 4.3.1. Where the range is used, it represents is the absolute difference of two (or more) replicate runs. Be sure that only absolute values are used in the determination of the range and average range!

$$R = \left| x_1 - x_2 \right|$$
 Eqn. 1

4.3.2. Calculate the average range of the trials, \overline{R} , for the *n* tests as follows:

$$\overline{R} = \frac{\sum |R|}{n}$$
 Eqn. 2

4.3.3. The estimated standard deviation may be calculated using the average range as follows (obtain values for d_2^* from NISTIR 6969, Table 9.1 and see NISTIR 6969, Section 8.3 for additional notes). For two runs, the d_2^* value is 1.414:

$$s_p = \frac{\overline{R}}{d_2^*}$$
 Eqn. 3

- 4.4 Construct Appropriate Charts
 - 4.4.1. Construct Standard Deviation Charts and Range Charts using the same approach. You should use 2 and 3 as the respective multipliers for the Upper Warning Limit and Upper Control Limits on standard deviation charts. Note that there will be no negative numbers when calculating standard deviations or ranges.

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4.4.2. Construct an R control chart having the following limits:

Central Line	\overline{R}
Lower warning and control limits	0
Upper warning limit (UWL)	$2.512\overline{R}$
Upper control (or action) limit (UCL)	$3.267 \overline{R}$

These limits are for duplicate measurements. For triplicate measurements see Handbook 145, Section 7.4.

4.2.3 The recommended format for construction of *R* control charts is given in NBS Handbook, Section 7.4.

4.3 Use of Control Charts

- 4.3.1 Replicate trials are run on each standard submitted to the laboratory for calibration according to the specific SOP. The values for standard deviation or range are plotted on the appropriate control chart, preferably in sequential order. The limits of the charts are such that 95 % of the values should fall within the warning limits and rarely should a value fall outside the control limits, provided that the system is in a state of statistical control.
- 4.3.2 If the values plotted on the standard deviation or range chart fall outside of the control limits, a decrease in precision is indicated. Problems with the standards or process will need to be investigated.
- 4.3.3 No calibration data should be accepted when the system is out of control.
- 4.3.4 If a plotted value for standard deviation or range is outside of the warning limit but inside the control limit, a second set of replicate calibrations should be made. If the new value for the standard deviation or range is within the warning limit, the process may be considered in control. If it lies outside of the warning limit, lack of control is indicated. Investigation should be made, with suitable corrective action taken with demonstrated attainment of control before calibration measurements are considered to be acceptable.
- 4.3.5 Even while the system is in an apparent state of control, incipient troubles may be indicated when the control data show short- or long-term trends, shifts, or runs.

5 Interpretation of Control Chart Data

5.1 Demonstration of "in control" indicates that the calibration process is consistent with the past experience of the laboratory. That is to say, there is no reason to believe that excessive changes in precision have occurred.

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- 5.2 The accuracy is inferred from a consideration of control of the sources of bias.
- 5.3 To the extent appropriate, the precision of measurement of standards may be extended to the calibration of other standards of similar type, capacity and design.

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Appendix

Standard Deviation and Range Data

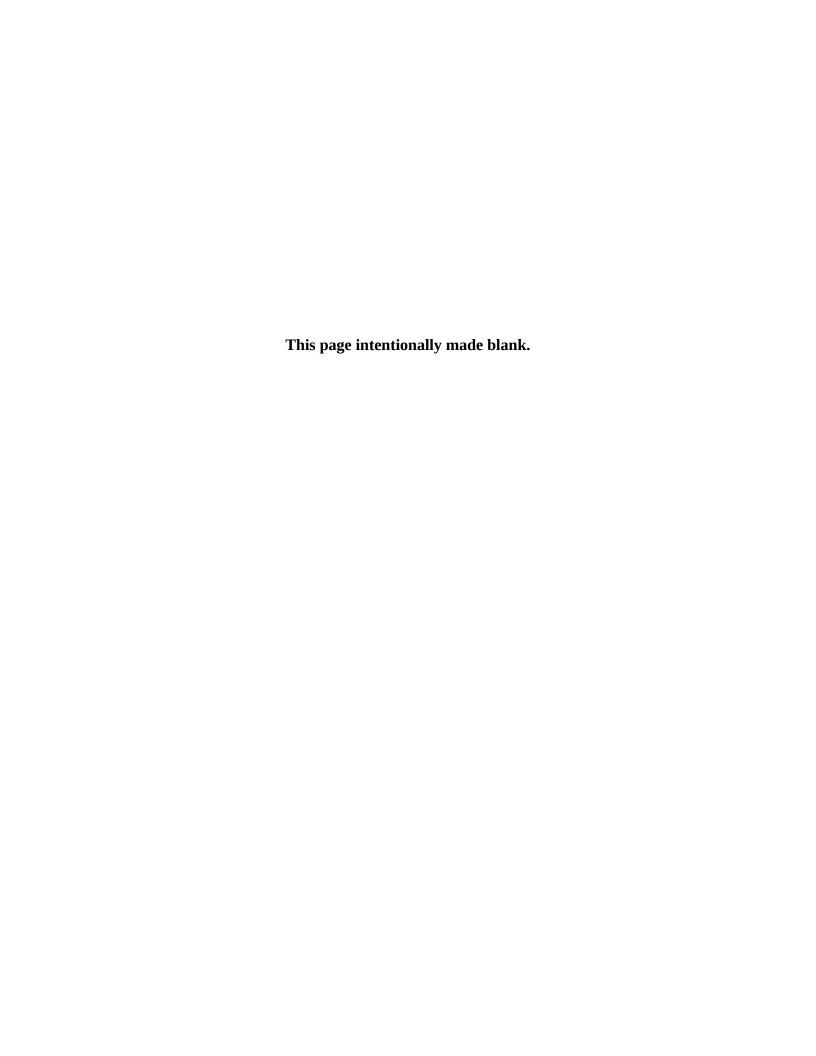
Standard Capacity: Laboratory:

Standard	Сарасну.			aboratory.		
Test Number	Date	Run 1	Run 2	Average of Runs	Standard deviation*	Range** Run 1 - Run 2 (Max - Min)
1						
2						
3						
4						
5						
6						
7						
8						
9						
10						
11						
12						
13						
14						
15						
16						
17						
18						
19						
20						

^{*} Use of the standard deviation and pooled standard deviations are preferred to the use of Range as an estimate of the standard deviation.

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^{**} This is the range, R, of the two trials and is actually the larger value minus the smaller value.



SOP 21

Standard Operating Procedure for Calibration of LPG Provers¹

1 Introduction

1.1 Purpose of Test

This procedure may be used to calibrate a volume standard used to test systems designed to measure and deliver liquefied petroleum gas (LPG) in the liquid state by definite volume, whether installed in a permanent location or mounted on a vehicle. A schematic diagram of such a prover is shown in Figure 1, together with numbers, e.g., 1, 2, 3, and 4 to clarify the various operations described in the procedure. The parts labeled A, B, and C are hose connections used in meter testing (versus prover calibrations). The prover being calibrated should be evaluated for conformance to appropriate specifications (using the checklist provided in NIST Handbook 105-4, Specifications and Tolerances for Liquefied Petroleum Gas and Anhydrous Ammonia Liquid Volumetric Provers, 2010) if being used for legal weights and measures applications. You should also review the schematic provided in Handbook 105-4.

1.2 Prerequisites

1.2.1 Verify the unknown prover has been properly cleaned and vented with all petroleum products removed prior to submission for calibration to ensure laboratory safety. The prover and/or pressure gauge may be visually inspected to determine that residual products are not present. Smell is not necessarily an adequate indicator of cleanliness.

NOTES: Many laboratories have a policy regarding cleanliness of submitted volumetric standards to minimize water contamination with flammable petroleum products. Be sure to follow laboratory safety policies for dealing with pressurized vessels.

1.2.2 Verify that valid current calibration certificates with measurement values and uncertainties are available for all of the standards used in the test. All calibration values must have demonstrated metrological traceability to the International System of Units (SI). Metrological traceability may be to the SI through a National Metrology Institute (for example, NIST.)

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¹ Non-SI units are predominately in common use in State legal metrology laboratories, and/or the petroleum industry for many volumetric measurements, therefore non-SI units have been used to reflect the practical needs of the laboratories performing these measurements as appropriate. The majority of LPG provers in use are 20 gal, 25 gal, and 100 gal nominal sizes. The volume of LPG provers is established at 60 °F and 100 psig.

- 1.2.3 Verify that the standards to be used have sufficiently small standard uncertainties for the intended level of the calibration.
- 1.2.4 Verify the availability of an adequate supply of clean, preferably soft water (filtered and thermally equilibrated as appropriate) water (GLP 10). Water does not need to be distilled or deionized for use in this procedure. The equations used in GLP 10 for the calculation of water density (air saturated) may be used without a significant impact on the measurement results.
- 1.2.5 Verify that the operator has had specific training in SOP 19, SOP 20, SOP 21, and GMP 3 and is familiar with the operating characteristics and conditioning of the standards and unknown test items being used and calibrated.
- 1.2.6 Cylinder of nitrogen or compressed air, or a regulated air source of at least 200 psig, and a proper pressure regulator.
- 1.2.7 Verify that the laboratory facilities meet the following minimum conditions to make possible the expected uncertainty achievable with this procedure:

Table 1. Laboratory environmental conditions.

Procedure	Temperature	Relative Humidity
Volume	18 °C to 27 °C	35 % to 65 %
Transfer	Stable to ± 2.0 °C / 1 h	Stable to ± 20 % / 4 h

1.3 Field tests

- 1.3.1 A "field" calibration is considered one in which a calibration is conducted in an uncontrolled environment, such as out-of-doors. Calibrations conducted under field and laboratory conditions are not considered equivalent.
- 1.3.2 The care required for field calibrations includes proper safety, a clean and bubble-free water supply, measurement control programs, and a stable temperature environment shaded from direct sunshine to allow the prover, field standard, and clean test liquid (water) to reach an equilibrium temperature with minimal evaporation. Environmental conditions must be selected to be within stated laboratory conditions during the measurements. All data and appropriate environmental conditions must be documented regardless of test location.

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2 Methodology

2.1 Scope, Precision, Accuracy

This procedure is applicable for the calibration of LPG provers with capacities of 100 L to 500 L (20 gal to 100 gal) or larger when appropriate. Provers of 20 gal, 25 gal, and 100 gal (with gal and in³ units) are encountered most frequently, hence the procedure is written with that in mind. The changes necessary for testing provers of other capacities will be obvious and are not described in this document. The agreement of duplicate measurements made within a short period of time on a given 100 gal LPG prover must be within 5 in³ (0.02 gal). For any nominal volume, replicate values must agree within 0.02 % of the volume. The accuracy will depend on the uncertainty in the volume of the standard, on the care exercised in making the various measurements and temperature readings, and on correct application of the corresponding corrections.

2.2 Summary

The procedure is a modification of one described by M.W. Jensen in NBS Handbook 99, "Examination of Liquefied Petroleum Gas Liquid- Measuring Devices." The LPG prover is calibrated with a known volume of water delivered into it from a standard prover of calibrated volume. Depending on the respective volumes, multiple transfers may be required. While multiple transfers should be minimized, a maximum number of 15 transfers are permitted to ensure that final calibration uncertainties are sufficiently small to meet user applications. The temperature of the calibration medium (water) cannot be considered to be constant during transfers; hence, the temperature of the water for each transfer must be measured. Because of the large volumes, the difference in thermal expansion of the respective vessels must be considered. The LPG prover is pressurized and the liquid level is measured at each of several values of applied pressure. The calibration thus defines the capacity of the prover over its expected range of operational pressures.

2.3 Standards and Equipment

- 2.3.1 Calibrated standard prover of minimum volume of 5 gal for 20 gal and 25 gal LPG provers is acceptable. A 10 gal standard is acceptable for calibrating a 100 gal LPG prover, but a standard that is of the same volume as the LPG prover is preferable. NOTE: standard provers used for calibration may need to have an alternative calibration value based on restricted flow delivery as the opening for many LPG provers may not be adequate to receive the full flow delivery from the reference standard.
- 2.3.2 A funnel for transferring water from calibrated flasks into the unknown prover.

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- 2.3.3 Calibrated flasks of suitable sizes to calibrate the neck of prover.
- 2.3.4 Thermometers (2) with resolution and uncertainty less than 0.1 °C.
- 2.3.5 Meniscus reading device (See GMP 3).
- 2.3.6 Timing device (calibration is not required; uncertainty of the measurement only needs to be less than 5 s for a 30 s pour time.)
- 2.3.7 Sturdy platform, with appropriate safety conditions, with sufficient height to hold standard and to permit transfer of water from it to the prover by gravity flow.
- 2.3.8 Clean the pipe or tubing (hoses) to facilitate transfer of water from the laboratory standard to prover. Nearly all LPG provers require reducers to be used between normal laboratory piping and the top hole on the prover. Pipe and hose lengths must be minimized to reduce water retention errors. Care must be taken during wet-downs and runs to ensure complete drainage and consistent retention in all hoses or pipes. Arrange all hoses so that there are no loops or low spots that can hold water.
- 2.3.9 Compressed nitrogen cylinder or air, suitable regulator, and an appropriate pressure gauge. The calibration relies on the accuracy of the pressure gauge on the prover. It assumes that systematic errors in the prover pressure gauge will be present in field applications as well, thus calibration of the laboratory pressure gauge is *not* essential. However, the prover must be recalibrated if pressure gauges are changed in the field. When a laboratory pressure gauge is used during the calibration, the unknown pressure gauge and laboratory gauge must both have current calibration reports.

2.4 Procedure

2.4.1 Preliminary Operations

- 2.4.1.1 Install and level the standard(s) on a raised platform with appropriate security and safety ensured for the prover(s) and operator(s). Provide pipe or tubing for delivery of water by most direct route to prover.
- 2.4.1.2 Position and level the unknown prover where it can be reached from the elevated standard by the shortest feasible delivery system.
- 2.4.1.3 It is recommended that all valves, hoses, and piping be removed from ports below the nominal volume line to eliminate errors that may result from leakage or expansion while the prover is under

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pressure. These potential errors typically only affect the calibration but not during normal use. Plug one port with a fitting and valve of suitable pressure rating for draining the prover. Plug all other ports with sealed plugs of suitable pressure ratings.

2.4.1.4 Remove the plug and relief valve (1) from the top, and extend the pipe into the hole. This may require the use of a reducer and a short length of hose (about 1 inch in diameter). If this is a tight fit, open the vapor return line valve (2) to provide an air bleed.

Use the prover inlet line (3) as a gravity drain. If necessary, remove the fitting on the end and connect a hose or pipe to make the necessary drain line.

Warning: ensure that a check valve is not plumbed into the prover inlet line. If it is, remove the check valve, otherwise the prover will need to be drained via the plug in the bottom of the lower neck.

2.4.2 Cleanliness Check

Both the standard and the unknown prover must be internally clean. This should be verified by checking that water drains properly from them. If necessary, either or both should be cleaned with water and non-foaming detergent (see GMP No. 6) to attain good drainage characteristics. Additional effort may be required to eliminate scaling and contamination build-up from the inside of LPG provers.

- 2.4.3 Neck scale plate verification (This is generally only conducted for new or damaged measures or those that have not been calibrated by the laboratory in the past.)
 - 2.4.3.1 Fill the unknown prover with water from the standard. Check the prover level condition in the same way in which it will be used and adjust if necessary by placing a precision spirit or electronic digital level vertically on the neck on at least two locations, 90 degrees apart around the circumference of the neck and adjust the orientation of the standard until the neck is as close to vertical (plumb or perpendicular to the horizontal plane) as possible. Verify and adjust any mounted levels that are on the prover to agree (when present and when possible). Check the prover system for leaks. This is a wet-down run.
 - 2.4.3.2 Bleed the liquid level down to a graduation near the bottom of the upper neck. "Rock" the prover to "bounce" the liquid level, momentarily, to ensure that it has reached an equilibrium level.

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Read and record this setting to be used as the initial scale reading sr_i . This is in preparation for calibration of the neck scale.

- 2.4.3.3 Remove the fill hose or pipe from the top and insert a funnel.
- 2.4.3.4 Recheck the scale reading, then add water in a quantity equal to approximately $\frac{1}{4}$ or $\frac{1}{5}$ of the graduated neck volume from a suitable standard; record the scale reading.
- 2.4.3.5 Repeat 2.4.3.4 by successive additions until water is near the top of the scale (the neck capacity is usually about 5 gal on a 100 gal LPG prover). Record scale readings after each addition. The last reading will be the final scale reading, *sr_f*. The closer the water is to the top of the neck, the harder it may be to "bounce" the liquid in the gauge.

A plot of scale readings with respect to the total volume of water that is added V_w should be linear and will be a gross check of the validity of this calibration.

- 2.4.3.6 Calculate and assess the accuracy of the neck scale for each interval. The error should be less than 0.5 % of the graduated neck volume. If more than this, the scale should ideally be replaced. Alternatively, a Neck Scale Correction Value (NSCV) may be issued with instructions to the user if it is anticipated that this correction value will be used.
- 2.4.3.7 Neck scale errors less than 0.5 % of the graduated neck volume should be included in the uncertainty estimate associated with the prover calibration. Uncertainties associated with the neck scale calibration are primarily related to the setting of the meniscus on the flask used (with each drop), the reading of the meniscus on the unknown prover (at each volume reading), and the calibration uncertainty of the flask. These uncertainties must be considered when evaluating the acceptability of the neck scale.
- 2.4.3.8 The neck scale correction value is calculated as follows:

$$NSCV = \frac{V_w}{\left(sr_f - sr_i\right)}$$
 Eqn. 1

Table 2. Variables for neck scale correction value equation.

NSCV	Neck scale correction value	
V_{w}	Total volume of water added to neck	
sr_f	Scale reading, final	

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Ī	Sr.	Scale reading, initial
I	s_{i}	2 0000 1 00000005, 11100001

2.4.4 Body Calibration

2.4.4.1 Drain the unknown LPG prover through its inlet valve and the liquid bleeder valve. When the liquid reaches the top of the lower gauge glass, close the inlet valve and allow the water to drain from the interior of the prover into the lower neck for 30 s, while controlling the water flow and level with the bleeder valve until the liquid meniscus reaches the zero graduation. The liquid level should be exactly at the zero graduation and the bleeder valve closed simultaneously at the 30 s drain time. (Draining with the bleeder valve close to the zero mark should be started during the 30 s drain period but should not be completed before the end of the drain period.)

Alternatively, though not recommended, the prover may be completely drained with a 30 s drain time and then refilled with a funnel that has been wet down, and adding a small volume of water to set the zero mark. Errors from this process will result due to the additional drain and retention time and other factors associated with the process and should be considered in the uncertainty assessment.

2.4.4.2 Run 1. Transfer the volume from the standard in the usual manner, and record the standard and prover temperature readings. If multiple transfers are required, record temperature of the standard at the time of each transfer, but that of the prover only after the final transfer. Check the prover level by placing a precision spirit or electronic digital level vertically on the neck on at least two locations, 90 degrees apart around the circumference of the neck and adjust the orientation of the standard until the neck is as close to vertical (plumb or perpendicular to the horizontal plane) as possible. Verify and adjust any mounted levels that are on the prover to agree (when present and when possible). "Rock" the prover to "bounce" the liquid in the upper gauge glass before reading. Record the final scale reading after the nominal volume has been transferred into the unknown LPG prover with no pressure applied.

Note regarding temperature measurements: A digital temperature sensing device with a long cable can allow insertion of the probe into the standard and the unknown to enable direct liquid temperature measurements at the bottom, middle, and top of the

provers. If the prover thermometer wells are used, ensure that the prover has equalized with the temperature of the water.

- Run 2. Drain the LPG prover as described in 2.4.4.1 and make another test run. Record the temperatures of the standard(s) and the unknown and final scale reading. Calculate the prover error at 0 psig (no pressure applied) for each run using the appropriate equations in Section 3; these values are used to evaluate repeatability of the test only. The error at 0 psig for the test runs must agree within 0.02 % of the prover volume or approximately one-half the prover tolerance (i.e., 5 in³ on a 100 gal LPG prover). If the two runs fail to agree within 0.02 % of the prover volume, identify and correct problems with the prover or set-up. Then, continue until replicates agree within these limits, taking care to ensure that poor cleanliness, contamination, bubbles in hoses, leaking valves or seals are not contributing to poor repeatability. Lack of measurement agreement may also be due to poor field conditions, such as when calibration is conducted in unstable environments. Poor agreement must be corrected before calibration can be completed.
- 2.4.4.4 While the LPG prover remains full from Run 2, replace the relief valve and plug in the top of the prover using suitable pipe joint compound or tape.

2.4.5 Prover Adjustments

2.4.5.1 The internal pressure and hence the volume of the prover may vary during use. Accordingly, a pressure correction must be made using the data of steps 2.4.6.

To minimize the amount of correction needed when the prover is in use, the prover should be adjusted to indicate its nominal capacity when 100 psig is applied. (An internal pressure of 100 psig is suggested as being convenient.) If the actual volume of the prover is not near a convenient whole gallon value and cannot easily be adjusted to a whole gallon value, a prover correction value can be computed (see 3.3) and added to the pressure correction values to obtain a set of combined prover and pressure correction values to be computed. The pressure correction is computed in 3.3. The pressure gauge that is mounted to the LPG prover may be used for setting calibration pressures, but, if following NIST Handbook 105-4, it is required to be calibrated. Since the correction values assigned to the prover during calibration will match those during use, the

prover corrections obtained using a mounted pressure gauge are generally acceptable. Calibration status of the pressure gauge mounted on the prover should be assessed and included on the calibration report.

2.4.5.2 Use a cylinder of nitrogen or compressed air and a proper pressure regulator with an integrated pressure gauge.

Connect the cylinder regulator output to the vapor return fitting (2) near the top of the neck. This may require fashioning a connection with steel pipe nipples or other appropriate materials and the existing fittings.

Caution: Ensure that all piping and fittings are rated for the pressures to which they will be exposed.

Make sure all valves are closed except the vapor return valve. Verify that the final scale reading has not changed since it was recorded (if it has changed it may signal a leak in one of the valves or fittings), and then slowly introduce pressure until the installed prover gauge reads 100 psig. Lightly tap the gauge to ensure that the gauge needle is not sticking. "Bounce" the liquid in the neck, then read and record the liquid level at this applied pressure.

Return pressure to 0 psig and reapply pressure as above. This reading at 100 psig should agree with the above recorded reading within 0.02 % of the nominal volume of the prover. Leaking seals or valves may cause problems with repeatability of the gauge readings under pressure.

2.4.5.3 With the pressure in the prover at 100 psig, adjust the upper scale to read the nominal volume. This is accomplished by adjusting the upper scale so that the water level reading is:

Desired scale reading = $V_{X60} - (V_{NOM} * 1.00032)$ Eqn. 2

Take care to use like units in the calculation. The calculation for V_{X60} is given in 3.1, Eqn. 5 or Eqn. 6.

Note: The correction factor 1.00032 corrects for the compressibility of the water at 60 °F and 100 psig. If the upper scale is not adjustable, see 2.4.5.4.

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2.4.5.4 For provers with only an adjustable lower scale (or one in which the upper scale is not adjustable), a prover correction, L_C , may be calculated at 100 psig as follows:

$$L_C = V_{X60} - (V_{NOM} *1.00032) - s r_u$$
 Eqn. 3

where:

L_{C}	Prover correction at 100 psig
V_{X60}	Volume of unknown prover at 60 °F
V_{NOM}	Nominal volume of prover
sr_u	Upper scale reading at 100 psig
1.00032	Correction factor for the compressibility of
	water at 100 psig

Take care to use like units in the calculation.

If the prover correction is negative, move the bottom scale down to increase the prover volume. If the prover correction is positive, move the bottom scale up to decrease the prover volume. The distance h that the bottom scale is to be moved is:

$$h = \frac{4 \mid L_C \mid}{\pi d^2}$$
 Eqn. 4

where:

h	Distance in inches the bottom scale is to be
	moved, up or down
L_{C}	Prover correction at 100 psig in cubic inches
d	Inside diameter of the lower neck of the prover in inches (as noted on identification plate)

2.4.6 Pressure Correction

- 2.4.6.1 Return pressure to 0 psig. Record the reading. Slowly introduce pressure until the installed prover gauge reads 50 psig. Lightly tap the gauge to ensure that the gauge needle is not sticking. "Bounce" the liquid in the neck, then read and record the liquid level at this applied pressure.
- 2.4.6.2 Repeat step 2.4.6.1 at 100, 150, and 200 psig. Other pressure points in between those listed may be tested if so desired. (The water level should decrease 10 in³ to 15 in³ for each 50 psig

increase in pressure on a 100 gal prover, although this varies depending on the geometry and age of the LPG prover. Water level changes significantly greater than indicated may be due to leaking seals and/or valves.) Erratic pressure readings may also be due to air entrapment based on prover design; repeated pressurizing of the prover should eliminate entrapped air. Air entrapment problems due to design may need to be investigated and corrected.

2.4.6.3 Repeat step 2.4.6.1 as the pressure is bled down to 150, 100, 50, and 0 psig (atmospheric pressure). The readings must agree with those previously obtained within approximately 0.02 % of the nominal prover volume. If the data are not linear with respect to pressure, repeat the series of measurements above to verify the nonlinearity of the readings. Leaking seals or valves may cause problems with repeatability of the gauge readings under pressure. The cause of poor agreement must be identified and corrected before continuing the calibration.

2.4.7 Final Operations

- 2.4.7.1 Seal the bottom and top scales as specified by laboratory policy and as appropriate.
- 2.4.7.2 Drain prover, then remove plug (5) at the lower neck to facilitate drainage below the lower gauge. If time permits, let the prover drain overnight.
- 2.4.7.3 With the nitrogen cylinder or compressed air connected, blow nitrogen or air through the prover to remove remaining moisture. Be sure to blow out the drain line and any other portions of the system that may have become contaminated with water.
- 2.4.7.4 If water has entered the pump-off system, pour several gallons of alcohol into the prover and pump the alcohol through the system to remove the water to prevent it from freezing in the pump when LP gas is used.

3 Calculations

3.1 Single Delivery

3.1.1 Calculate V_{X60} , the volume of the unknown prover at 60 °F, using the following equation:

$$V_{X60} = \frac{\rho_1 \left\{ \left(V_{S60} + \Delta_1 \right) \left[1 + \alpha \left(t_1 - 60 \, ^{\circ} \text{F} \right) \right] \right\}}{\rho_x \left[1 + \beta \left(t_x - 60 \, ^{\circ} \text{F} \right) \right]} \quad \text{Eqn. 5}$$

3.2 Multiple Deliveries

3.2.1 Calculate V_{X60} , the volume of the unknown prover at 60 °F, using the following equation:

$$V_{X60} = \frac{\rho_1 \left\{ \left(V_{S60} + \Delta_1 \right) \left[1 + \alpha \left(t_1 - 60 \, ^{\circ} \mathrm{F} \right) \right] \right\} + \rho_2 \left\{ \left(V_{S60} + \Delta_2 \right) \left[1 + \alpha \left(t_2 - 60 \, ^{\circ} \mathrm{F} \right) \right] \right\} + \dots + \rho_N \left\{ \left(V_{S60} + \Delta_N \right) \left[1 + \alpha \left(t_N - 60 \, ^{\circ} \mathrm{F} \right) \right] \right\}}{\rho_x \left[1 + \beta \left(t_x - 60 \, ^{\circ} \mathrm{F} \right) \right]} \quad \text{Eqn. } 6$$

Table 3. Variables for V_{X60} equations.

	Symbols Used in Equations				
V_{X60}	volume of the unknown vessel at 60 °F				
V_{S60}	volume of the standard vessel at 60 °F				
$\rho_1, \rho_2,, \rho_N$	density of the water in the standard prover where ρ_1 is the density of the water for the first delivery, ρ_2 is the density of the water for the second delivery, and so on until all N deliveries are completed				
$\Delta_1, \Delta_2,, \Delta_N$	volume difference between water level and the reference mark on the standard where the subscripts $1, 2,, N$, represent each delivery as above. If the water level is below the reference line, Δ is negative. If the water level is above the reference line, Δ is positive. If the water level is at the reference line, Δ is zero NOTE: units must match volume units for the standard. For slicker-plate type standards, the Δ is zero.				
$t_1, t_2,, t_N$	temperature of water for each delivery with the subscripts as above				
α	coefficient of cubical expansion for the standard in units / °F				
β	coefficient of cubical expansion for the prover in units / °F				
t_x	temperature of the water in the filled unknown vessel in units °F				
$\rho_{\scriptscriptstyle X}$	density of the water in the unknown vessel in g/cm ³				
Note: Values fo GLP 10 for air-s	r the density of water at the respective temperatures may be calculated using the equations given in aturated water.				

3.3 Pressure Corrections

Compute the pressure correction, P_{corr} , at each pressure that the prover was read, after any adjustments, by correcting for the compressibility of the water. The equation is:

$$\begin{split} P_{corr} &= Scale \ reading \ @ \ 100 \ psig - scale \ reading \ @ \ X \ psig \\ &+ \left(water \ compressiblity \ factor \right) \left(\frac{100 \ psig - X \ psig}{100}\right) \end{split}$$
 Eqn. 7

where the water correction factor due to compressibility is 7.4 in³ or 0.032 gal for a 100 gallon prover (1.8 in³ (0.008 gal) for a 25 gallon prover; 1.5 in³ (0.006 gal) for a 20 gal prover). The compressibility value is 0.00032 gal/al at 100 psig. The compressibility factor is given in both cubic inches and gallons so the proper unit can be selected depending upon the unit used for the scale readings.

Plot the pressure corrections. If the corrections versus the pressure are linear, make a straight line best fit of the data and interpolate to obtain the pressure corrections for any desired pressure. If the data is nonlinear, then perform a straight line interpolation between adjacent pressure readings to obtain pressure corrections at any desired intermediate pressures. Alternatively, a best fit curve can be drawn for the nonlinear data and the pressure corrections interpolated from the graph for intermediate pressures. See Appendix A for an example of the pressure corrections.

3.4 Prover Volume

LPG provers are generally adjusted to the nominal value using 100 psig as the reference pressure and a 60 °F reference temperature. A gauge reading on a prover that indicates a nominal volume at 0 psig will be lower when pressurized to 100 psig due to both the compressibility of water under pressure and the expansion of the prover. The expansion of the prover is consistent whether there is water in the prover or LPG. However, a correction must be made for the compressibility of water as noted in 3.3. Thus, to adjust a 100 gal prover to its nominal volume at 100 psig, the gauge reading should be set to the V_{X60} value minus 7.4 in³. (The water level on the prover's upper scale will be below a calculated V_{X60} value by 7.4 in³ when pressurized to 100 psig.)

Another possible way to handle the compressibility of water is to add the volume of water (e.g., 7.4 in^3 for a 100 gal prover) to the volume of water in the prover so that it can be set to the V_{X60} value at 100 psig.

The total calculated volume of the prover at its reference temperature and pressures at 0 psig, 50 psig, 100 psig, and 150 psig and 200 psig, should be reported on the calibration report.

4 Measurement Assurance

4.1 Duplicate the process with a suitable check standard or have a suitable range of check standards for the laboratory. See SOP 17, SOP 20 and SOP 30. Plot the

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check standard volume and verify it is within established limits OR a *t*-test may be incorporated to check the observed value against an accepted value. The mean of the check standard observations is used to evaluate bias and drift over time. Check standard observations are used to calculate the standard deviation of the measurement process which contributes to the Type A uncertainty components.

4.2 If a standard deviation chart is used for measurement assurance, the standard deviation of each combination of Run 1 and Run 2 is calculated and the pooled (or average) standard deviation is used as the estimate of variability in the measurement process. Note: the pooled or average standard deviation over time will reflect varying conditions of LPG provers that are submitted to the laboratory but do not reflect potential meniscus reading errors (See GMP 3.)

5 Assignment of Uncertainties

- 5.1 The limits of expanded uncertainty, U, include estimates of the standard uncertainty of the laboratory volumetric standards used, u_s , plus the uncertainty of measurement, u_m , at the 95 % level of confidence. See SOP 29 for the complete standard operating procedure for calculating the uncertainty.
 - 5.1.1 The standard uncertainty for the standard, u_s , is obtained from the calibration report. The combined standard uncertainty, u_c , is used and not the expanded uncertainty, U, therefore the reported uncertainty for the standard will usually need to be divided by the coverage factor k.

Note: See SOP 29 for the complete standard operating procedure for calculating the uncertainty when multiple deliveries or multiple standards are used to correctly calculate correlated uncertainties. Fifteen is the maximum recommended number of deliveries from a laboratory standard to a prover under test to minimize calibration uncertainties to the levels identified previously.

- 5.2 The standard deviation of the measurement process s_p , is taken from a control chart for an LPG check standard (not from charts for a refined fuel check standard) or standard deviation charts for LPG calibrations (See SOP 17, SOP 20 and SOP 30). Check standards are not normally available.
- 5.3 Neck calibration uncertainty should be estimated based on the uncertainty of standards used, errors observed during calibration, ability to read the meniscus of all standards involved (see GMP 3), and the repeatability of the neck calibration.
- 5.4 Other standard uncertainties usually included at this calibration level are 1) uncertainties associated with the ability to read the meniscus, only part of which is included in the process variability due to parallax and visual capabilities, and 2) uncertainties associated with temperature corrections that include values for the

- cubical coefficient of expansion for the prover under test, the accuracy and gradients associated with temperature measurements in the provers. Additional factors that might be included are: data showing reproducibility, environmental variations over time, and bias or drift of the standard as noted in control charts.
- To properly evaluate uncertainties and user requirements (tolerances), assessment of additional user uncertainties may be required by laboratory staff. Through proper use of documented laboratory and field procedures, additional uncertainty factors may be minimized to a level that does not contribute significantly to the previously described factors. Additional standard uncertainties in the calibration of field standards and their use in meter verification may include: how the prover level is established, how delivery and drain times are determined, the use of a proper "wet-down" prior to calibration or use, whether gravity drain is used during calibration or whether the volume of water is eliminated by pumping, the cleanliness of the prover and calibration medium, prover retention characteristics related to inside surface, contamination or corrosion, and total drain times, and possible air entrapment in the water. Systematic errors may be observed between laboratory calibration practices where a gravity drain is used and field use where the pumping system is used.
- 5.1 Example components to be considered for an uncertainty budget table are shown in Table 4. Multiple values of some items may need to be considered (e.g., multiple drops from the standard, multiple meniscus readings, and multiple temperature readings.)

Table 4. Example Uncertainty Budget Table.

Uncertainty Component Description	Symbol	Source	Typical Distribution
Uncertainty of the standard (5.1)	u_s	Calibration report; may be multiplied or added based on dependencies	Rectangular or Normal divided by coverage factor
Accepted standard deviation of the process (5.2)	S_p	Control chart, standard deviation chart	Normal
Uncertainty or uncorrected error associated with a neck calibration (5.3)	u_n	From experimental data	Rectangular
Ability to read the Meniscus in S (5.4)	u_m	None if using a slicker- plate type standard; GMP 3	Triangular
Ability to read the Meniscus in X (5.4)	u_m	GMP 3	Triangular
Water temperature (S) (5.4)	u_{ts}	Consider accuracy, resolution, and gradients	Rectangular
Water temperature (X) (5.4)	u_{tx}	Consider accuracy, resolution, and gradients	Rectangular
Cubical Coefficient of	u_{CCE}	5 % to 10 %	Rectangular

Expansion on S (5.4)		(EURAMET CG-21)	
Cubical Coefficient of Expansion on X (5.4)	u_{CCE}	5 % to 10 % (EURAMET CG-21)	Rectangular
Uncertainty associated with the pressure gauge	u_{psig}	Handbook 105-4, 5 psig increments; use 2.5 psig	Rectangular
Uncertainty of bias or drift of standards (5.2)	u_b	From control chart	Rectangular
Uncertainty of drain time	u_d	From experimental data	Normal

6 Report

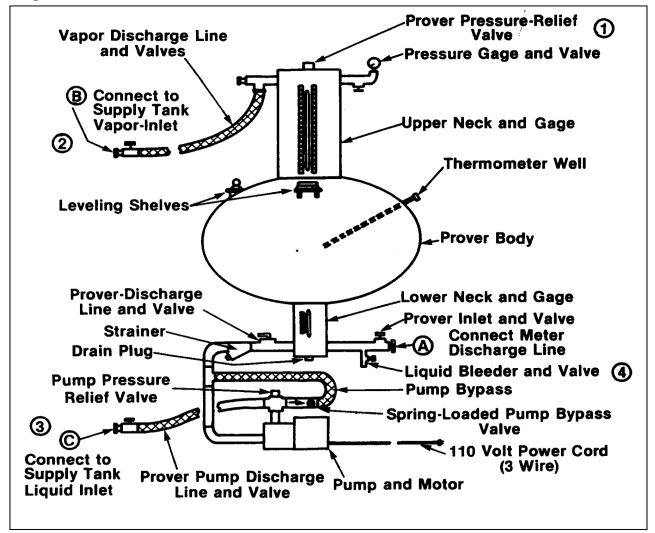
Report results as described in SOP No. 1, Preparation of Calibration/Test Results, with the addition of the following:

For LPG provers, the total prover volume and uncertainty, reference temperature, material, coefficient of expansion (assumed or measured), any identifying markings, tolerances (if appropriate), laboratory temperature, water temperature, barometric pressure, relative humidity, out-of-tolerance conditions, and the total drain time from opening of the valve, including the 30 s drain after cessation of flow.

The report should also include temperature and pressure correction tables or chart, along with a note regarding possible differences in retention characteristics between water, the calibration medium, and LPG products.

The SI unit of volume is m³, so a conversion factor is to be included on the report in the notes section when other volumes are used.

Figure 1. LPG Prover Schematic.



Appendix AVolume Transfer Data Sheet for LPG Provers

Laboratory data and conditions:

Vessel Owner	Operator	
Vessel ID	Date	
Nominal Volume	Air Temperature	
M aterial	Relative Humidity	
Cubical Coefficient of Expansion	Standard deviation of the process, s_p from the Standard Deviation Chart	
Reference temperature of unknown prover	Degrees of Freedom	
Unknown prover graduations	Applicable specifications and tolerances	

Volume standard(s) data:

ID (Note ID of Standards)	Nominal Volume	Volume/Correction	Expanded Unc: From cal. report	Unc: k factor	Cubical Coefficient of Expansion
S					
S					

Run 1: Measurement Observations

DROP #	Reported Volume (gal)	Material (MS	/SS/BS/SL)	Water Temp (°C (Must be ≥ 0.5 °C and <	•	Gauge Delta (in³)
1						
	Material for the Unknown:				MS, SS, T	P, AL, PVLC
Final Gauge Reading at 0 psig:				in ³		
	Final Water Temperature:				°C	

Run 2: Measurement Observations

DROP #	Reported Volume (gal)	Material (MS/SS/BS/SL)		Water Temp (°C) (Must be ≥ 0.5 °C and < 40 °C)		Gauge Delta (in³)
1						
	Material for the Unknown:				MS, SS, T	P, AL, PVLC
	Final Gauge Reading at 0 psig:			in ³		
	Final Water Temperature:				°C	

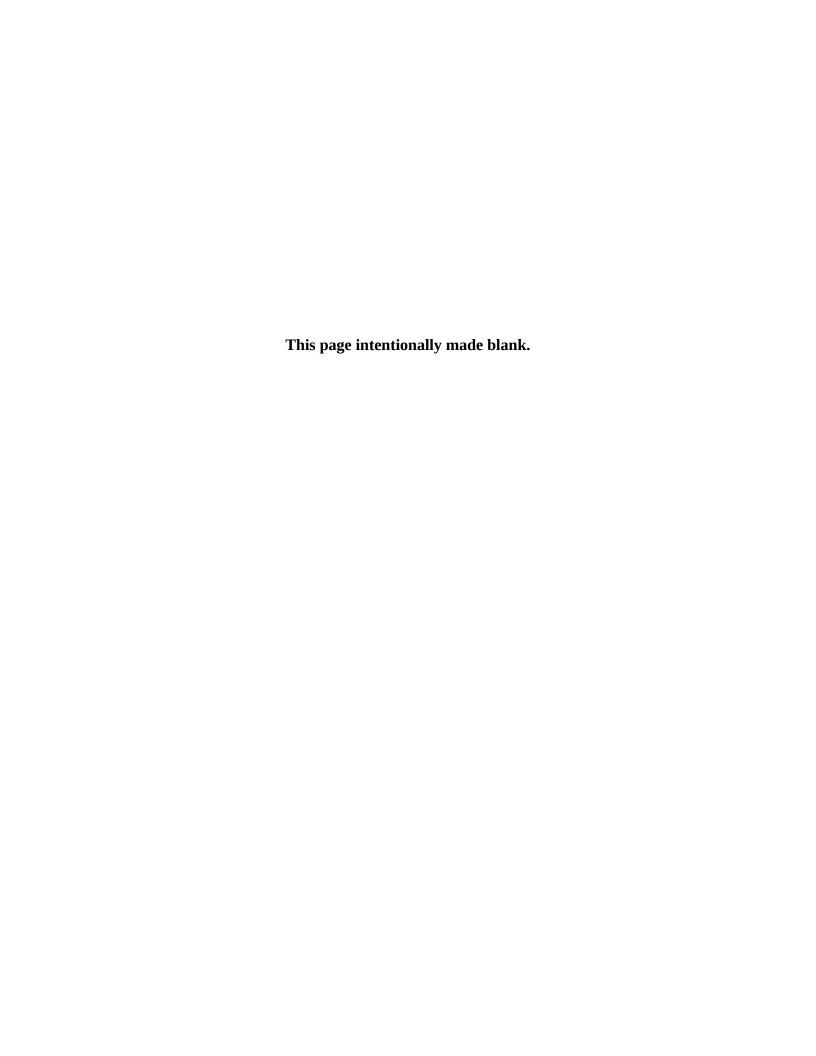
Run 2: Pressure Observations

Gauge Reading at 100 psig	in ³		
Gauge Reading at 0 psig:	in ³	Acceptable r	reccure
Repeated Gauge Reading at 100 psig	in ³	repeatability	
Increasing Pressure Readings, 0 psig	Decreasing Pressure Readings, 150 psig		in^3
Increasing Pressure Readings, 50 psig	Decreasing Pressure Readings, 100 psig		in ³
Increasing Pressure Readings, 100 psig	Decreasing Pressure Readings, 50 psig		in^3
Increasing Pressure Readings, 150 psig	Decreasing Pressure Readings, 0 psig		in^3
Increasing Pressure Readings, 200 psig			in^3

Appendix B
Example Temperature Correction Table*

Example for a 100 gallon prover and 60 °F reference temperature.						
CCE	Mild	Steel	Stainles	ss Steel		
CCE	1.86 x 1	10-5/°F	2.65 x 10-5/°F			
Temperature °F	in ³	gal	in ³	gal		
-20	-34	-0.149	-49	-0.212		
-15	-32	-0.139	-46	-0.199		
-10	-30	-0.130	-43	-0.186		
-5	-28	-0.121	-40	-0.172		
0	-26	-0.112	-37	-0.159		
5	-24	-0.102	-34	-0.146		
10	-21	-0.093	-31	-0.133		
15	-19	-0.084	-28	-0.119		
20	-17	-0.074	-24	-0.106		
25	-15	-0.065	-21	-0.093		
30	-13	-0.056	-18	-0.079		
35	-11	-0.047	-15	-0.066		
40	-9	-0.037	-12	-0.053		
45	-6	-0.028	-9	-0.040		
50	-4	-0.019	-6	-0.026		
55	-2	-0.009	-3	-0.013		
60	0	0.000	0	0.000		
65	2	0.009	3	0.013		
70	4	0.019	6	0.026		
75	6	0.028	9	0.040		
80	9	0.037	12	0.053		
85	11	0.046	15	0.066		
90	13	0.056	18	0.080		
95	15	0.065	21	0.093		
100	17	0.074	24	0.106		
105	19	0.084	28	0.119		
110	21	0.093	31	0.133		
115	24	0.102	34	0.146		
120	26	0.112	37	0.159		
CCE = coefficient of cul	CCE = coefficient of cubical expansion					

^{*}Provide only the applicable coefficient of cubical expansion for the prover under test.



SOP 26

Standard Operating Procedure For Gravimetric Calibration of Dynamic Volumetric Systems used as Standards¹

1 Introduction

1.1 Purpose of Test

This procedure describes the gravimetric calibration of volumetric systems that may be used as volumetric measuring standards. (Gravimetric calibration of fixed volume graduated-neck or slicker-plate type standards should be completed using SOP 14.) This procedure uses gravimetric calibration principles with mass values and pure water sources to obtain calibration uncertainties that are generally less than those obtained during volume transfer procedures of similar standards. Alternative test liquids are not considered in this procedure. Accordingly, the procedure is especially useful for high accuracy calibrations. The procedure uses replicate testing to incorporate measurement assurance steps to ensure the validity of the measurement process; additional analyses must be performed to assess for bias (e.g., gravimetric calibration of other similar sized standards using the same methods, or interlaboratory comparisons with similar standards or provers). The procedure makes use of an electronic balance and is suitable for all sizes of gravimetric calibrations as limited by the capacity and resolution of the balance, available water source, mass standards, and handling capacity of the laboratory.

This procedure can be used for single direction and bi-directional calibration of small volume provers, for calibration of volumes delivered through a master meter, for fixed volume delivery from alternative volumetric measuring systems, or other alternative test methods and standards where the gravimetric calibration provides a useful calibration result and measurement uncertainty. All steps in the procedure and calculations must be thoroughly reviewed and evaluated for appropriateness for the specific standards under test to ensure accurate measurement results are reported to the user.

1.2 Prerequisites

- 1.2.1 Verify that valid calibration certificates with appropriate values and uncertainties are available for all standards, instruments and sensors used in this procedure. Calibration values must have demonstrated metrological traceability to the International System of Units (SI), which may be to the SI, through a National Metrology Institute (for example, NIST.)
- 1.2.2 Verify that all standards and instruments (e.g., mass standards, thermometer, barometer, and hygrometer) have sufficiently small standard uncertainties for the level of calibration. Reference standards that need special care and protection should not be used for gravimetric calibration where there is risk for contamination or damage from water.

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¹ Non-SI units are predominately in common use in State legal metrology laboratories, and/or the petroleum industry for many volumetric measurements, therefore non-SI units are used to reflect the practical needs of the laboratories performing these measurements when appropriate.

- 1.2.3 Verify that the balance has sufficient capacity and is in good operating condition with sufficiently small process standard deviation. Process standard deviations need to be verified by a valid control chart or preliminary experiments to ascertain performance quality. The accuracy of the balance and weighing procedures should also be evaluated to minimize potential bias in the measurement process.
- 1.2.4 Verify that the operator is experienced and has demonstrated proficiency in precision weighing techniques and volumetric measurements and has had specific training in SOP 2, SOP 4, SOP 19, SOP 29, GMP 10, and especially in gravimetric calibrations such as those used in SOP 14.
- 1.2.5 Verify that an adequate quality and supply of distilled or de-ionized water (see GLP 10) is available. Do not use tap or other impure water sources for this procedure! The laboratory must have a continuous flow of water available and may thermally equilibrate water in a large volumetric prover or a water tank provided purity is ensured. Note: Calibration of some systems may require other liquids due to viscosity or lubricity requirements and alternative procedures should be considered. Use of calibration liquids other than water is not considered in this SOP.
- 1.2.6 Verify that the laboratory facility is free of air currents and/or temperature fluctuations where the mass measurements will be performed. The volumetric proving system must also be calibrated in a thermally stable environment. Where the volume and weighing areas are not the same, movement of standards and transfer vessels must not negatively affect environmental stability. Stable temperatures, and thermal equilibration of water, are especially important to minimize possible convection currents on precision balances. Humidity control is essential for minimizing possible evaporation and/or condensation during the measurement process.
- 1.2.7 Verify that the laboratory facilities meet the following minimum conditions to meet the expected uncertainty possible with this procedure:

Table 1. Laboratory environmental conditions.

Procedure	Temperature	Relative Humid ity
Gravimetric	20 °C to 23 °C Stable to \pm 1 °C / 1 h during the calibration.	40 % to 60 % Stable to ± 10 % / 4 h

1.2.8 All necessary maintenance to the unknown volumetric standard must be completed by the device owner prior to starting the gravimetric calibration. It is advisable to perform a static leak test prior to performing the test on systems that allow such testing. Replace any seals that are present on the prover if there is any doubt as to their integrity. If "as found" data is required, it must be obtained before any seal replacements. A second calibration will be required after any maintenance is completed to obtain an "as left" calibration value. Note: "as found" data is essential for setting, monitoring, or adjusting calibration intervals.

- 1.2.9 Any standard inlet(s) and outlet(s) that are not used in the calibration process must be blocked by the device owner before calibration by using a blind flange, double-block and bleed valve, or other appropriate hardware. All extra valves, piping must be evaluated to ensure that air is not entrapped in the system.
- 1.2.10 The device owner must provide all necessary manufacturers' manuals and hardware (e.g., including pipe fittings, valves, solenoids, wiring, etc.) as well as trained personnel to operate the device during the calibration to ensure that the device is calibrated in the same manner in which it will be used

2 Methodology

2.1 Scope, Precision, Accuracy

The procedure is applicable for the calibration of any size of volumetric measuring system that, when filled with water, will not overload the electronic balance used. Typical volumetric proving systems ranging in capacity from 5 gal to 120 gal may be calibrated. Continuous systems that measure these quantities that are controllable in the laboratory may be considered as well (e.g., a metering system delivering 40 gallons per minute to 100 gallons per minute may be possible.) Flow rates that will be assessed need to be determined between the laboratory and the standard owner prior to calibration and should consider the practical range of use, the laboratory capability, and any documentary or legal requirements of applicable specifications and tolerances.

The precision of this calibration depends on strict adherence to the various steps of the procedure. The accuracy attainable will depend on the uncertainties of the standard weights, the uncertainties associated with the air buoyancy corrections, the accuracy of temperature measurements, the purity and density of the water, and the thermal expansion corrections that are made. Another important factor to consider is the repeatability of the procedure and the repeatability of the standard under test.

2.2 Summary

The mass of the water delivered by the volumetric proving system is determined by weighing the water on an electronic balance and comparing it to mass standards. Volume is derived from the mass of the mass standards, the density of pure water, appropriate air buoyancy corrections, and thermal and pressure expansion corrections, using the equations provided in Section 3.

2.3 Standards, Equipment, and Calibration Notes

2.3.1 An electronic balance having sufficient capacity to weigh the loaded vessel is required. The repeatability of the balance, measurement process, and standard under test are limiting factors in the accuracy of the measurement. The resolution of the balance and repeatability of the measurement process must be smaller than the expected uncertainty of the calibration. NOTE: standard deviations obtained from precision mass

- calibrations generally do not reflect the process repeatability of this procedure; therefore repeatability must be assessed using this procedure.
- 2.3.2 Mass standards of sufficient quantity and capacity are required. Ordinarily, standards of ASTM Class 2 or 3 or OIML Class F₁ or F₂ weight specifications are required. Working standards of other classes are generally not designed to maintain adequate stability needed; however, corrections obtained within a few days of the volume calibration may be used if the uncertainty is sufficiently small. Mass standards are selected so that they are slightly larger than the volume of water and container or transfer vessel to be weighed. Linearity errors or additional uncertainties may need to be considered when the mass standards are slightly less than the volume or container being measured. When summations of masses are used, the summation mass is used, and the "effective density" must be calculated, taking care not to use "average" density values.
- 2.3.3 Filter paper and padding for protection of mass standards when placed on the balance pan, stacked, or placed on the calibration item may be used. When all such materials are used, care must be taken to determine when and how the mass of these items are incorporated into the process (as tare weight, or as tared/zeroed off the balance readings).
- 2.3.4 A transfer vessel is required that is made of some inert and stable material with a volumetric capacity larger than the volume to be measured and a port large enough to fill the vessel on the top surface. The transfer vessel should be as small as possible to minimize its influence on the balance, but approximately 25 % larger than the volume to be measured in order to prevent water loss due to splashing out of the top fill port. The transfer vessel may need to be suspended with a hoist or mounted to a pallet to be moved with a forklift or pallet jack.
- 2.3.5 Weight handling equipment is required for this procedure that has adequate capacity for handling the filled and empty transfer vessel. It must have the ability to be moved slowly and gently to minimize water sloshing and to gently set the empty or filled vessel on the balance pan.
- 2.3.6 Thermometers, with resolution and uncertainty less than 0.1 °C to determine water and volumetric proving system temperatures. Submersible thermometer probes and cables must be long enough to measure the temperature of the water near the center of the transfer vessel. Therefore, liquid-in-glass thermometers are not adequate.
- 2.3.7 Barometer with resolution and uncertainty less than 135 Pa (1 mm Hg to determine barometric pressure (absolute).
- 2.3.8 Hygrometer with resolution and uncertainty less than 10 % to determine relative humidity.

- 2.3.9 Pressure gauge(s) with resolution and uncertainty less than 1 psig for determining the volumetric proving system pressure (where applicable).
- 2.3.10 If a proving system does not have operational temperature sensors and pressure gauges installed, the initial inspection should reveal that the proving system is not adequately prepared for calibration. If the calibration relies on the accuracy of the temperature sensors and pressure gauges installed on the prover, they must be installed prior to the volume calibration and have current calibration reports from an accredited calibration laboratory that are readily available for inspection. When laboratory pressure and temperature sensors are used during the calibration all laboratory sensors must all have current calibration reports.
- 2.3.11 Mechanical problems and sources of errors that may be encountered during a calibration and affect the ability of the system to repeat may include many possibilities, several of which are listed below. Mechanical problems with the volumetric standard must be corrected before calibration. Quantification of errors that cannot be eliminated must be evaluated and included in the uncertainty budget table.
 - Proper operation of the system;
 - Air in the system;
 - Instability of the temperature of water or ambient conditions
 - Pressure fluctuations in the system;
 - Error in reading the test measure(s);
 - Incorrect drain time for reading test measure(s);
 - Contamination of the systems;
 - Damaged measures/system;
 - Error in reading temperature or pressure;
 - Displacer problems (when present);
 - Malfunction of switches (when present);
 - Damage, worn, or badly fitted seals;
 - Damage, scoring, or pitting on the inside of the proving system;
 - Damaged or misaligned detector switches (when present);
 - Light interference in optical switches (when present);
 - Malfunction of solenoid valve (when present);
 - Leaks in prover/system or bypass valves.
- 2.3.12 Errors that may be encountered in weighing steps of the calibration may include any of the items listed below. Quantification of errors that cannot be eliminated must be evaluated and included in the uncertainty budget table.
 - Balance drift;
 - Off center loading of the balance;
 - Non-linearity of the balance and improper selection of the mass standards to calibrate the weighing instrument in the range of use;
 - Air currents or convection currents when weighing is performed in a large mass laboratory or when the temperature of the water, system, or laboratory are not adequately equilibrated;
 - Care in placing items on the weighing instrument;

- Inadequate water quality.

2.4 Procedure

- 2.4.1 Inspect the volumetric proving system to verify that it and associated piping are drained, clean, and safe for placement in the laboratory. For large volumetric proving systems, ensure that there is adequate space in the laboratory to safely position the system near the pure water source and that a drain is accessible and that the proximity of the mass calibration area is convenient. Direct the driver to position to system appropriately, taking appropriate precautions to prevent injury to anyone or damage to the laboratory and/or equipment and standards. Allow adequate time for the system under test to equilibrate with the laboratory environment. The time required will be dependent on the difference between inside and outside temperature and relative humidity.
- 2.4.2 Initial inspection. Record the data about the system being calibrated, taking care to record all applicable serial numbers, coefficients of cubical expansion, reference temperatures, flow rates, presence or lack of the calibration status of all auxiliary standards associated with the standard (such as pressure gauges and temperature sensors), and means for sealing the system after calibration.
- 2.4.3 Install, or have the owner install, volumetric proving system hardware per manufacturer's instructions or have the owner provide this service. (Hardware must be provided by device owner.)
- 2.4.4 Connect the water supply to the volumetric proving system.
 - 2.4.4.1 In general, ensure that all air is bled from the system, suitable water supply is provided and the device temperature is in equilibrium with that of the water supply (several runs are generally required prior to the calibration to ensure that all air is out of the system and that all components are in thermal equilibrium with the water.) Set device valves as appropriate for the test to be conducted.
 - 2.4.4.2 Follow the operator's guidance (and per manufacturer's instructions) to become familiar with and to establish correct valve positions for each step of the calibration when applicable.
- 2.4.5 Install and/or use calibrated thermometers with evidence of current metrological traceability to record air temperature, water temperature, detector bar temperature (T_d) (where present), volumetric proving system temperatures (T_p) .
- 2.4.6 Install and/or use calibrated pressure gauge(s) to record prover pressure(s) (P_p) when present and appropriate.
- 2.4.7 Analyze the piping and valves to determine if the configuration needs to be changed to deliver water into the transfer vessel.
- 2.4.8 Record all of the prover-specific variables on the data entry form.

Note: All references in the following weighing options to the empty transfer vessel assume that the empty transfer vessel has been placed in a "wetted-down" condition. This wet down condition is achieved when the vessel has been filled with water and then drained using the same process that will be used throughout this calibration. Several runs may be needed on the volumetric proving system to ensure proper wetting of all lines and drains before the measurement data is recorded.

2.4.9 Weighings (Option A)

- 2.4.9.1 Zero the balance. Place a standard mass, M_{s1} , on the balance platform (M_{s1} should be slightly larger than the mass of the empty transfer vessel.) Record the balance indication as O_1 .
- 2.4.9.2 Zero the balance and then place the empty transfer vessel on balance platform and record the balance indication as O_2 . Caution: the transfer vessel must be dry on the outside for all weighings.
- 2.4.9.3 Record air temperature, barometric pressure and relative humidity at the time of these measurements. Due to timing of this type of calibration, environmental conditions must be recorded for both the empty and full transfer vessel.
- 2.4.9.4 Zero the balance and then place a standard mass, $M_{\rm s2}$, on the balance platform ($M_{\rm s2}$ should be slightly larger than the mass of the filled vessel.) Record the balance indication as O_3 . (This observation may be made before or after the weighing of the filled transfer vessel, depending on laboratory configurations and convenience.)
- 2.4.9.5 Fill the transfer vessel through the volumetric proving system. Do not fill the transfer vessel while it is on a laboratory balance. Ensure that all of the water from the volumetric standard or registered on a meter is captured in the transfer vessel. Verify that the water temperature in the system is uniform and stable throughout the filling process (± 0.2 °C). Read and record all pertinent volumetric standard temperatures and pressures during the calibration sequence. Place the lid on the transfer vessel to limit evaporation.
- 2.4.9.6 Zero the balance and then place the filled transfer vessel on the balance and record the balance indication as O_4 .
- 2.4.9.7 Read and record the temperature of the water in the transfer vessel immediately after weighing. It is critical to measure the temperature after weighing to ensure that water is not removed from the transfer vessel prior to obtaining the mass value.
- 2.4.9.8 Record the air temperature, barometric pressure and relative humidity at the time of these measurements.
- 2.4.9.9 Make at least five replicate runs with at least two runs at a flow rate differing from the others by at least 25 % The measured

- result (V_{60}) of each of these calibration runs must agree within 0.02 % of the average volume of all volume calibration runs (i.e., the maximum result minus the minimum result must be less than 0.02 % of the average measured volume).
- 2.4.9.10 Calculate the volume using Equation 1 in Section 3.1.1., (Option A).

2.4.10 Weighings (Option B)

- 2.4.10.1 Place mass standards that approximate the mass of the empty transfer vessel (with a lid) and adequate filter paper or other clean, lint-free padding material (to protect the standards being used) on the electronic balance. With these items on the balance, zero the balance. Note: the padding materials must be included on all measurements or their mass treated as tare.
- 2.4.10.2 Place additional mass standards approximating the nominal mass of the water volume to be measured on the balance. Take care to place the weights on filter paper or other appropriate protective padding. Record this balance indication as O_1 .
- 2.4.10.3 Remove all mass standards. Record the air temperature, barometric pressure, and relative humidity readings.
- 2.4.10.4 Place the empty transfer vessel on the balance, with the padding material and zero the balance indication. Remove the transfer vessel, lid, and padding material and record the base empty vessel reading (B).
- 2.4.10.5 Fill the transfer vessel from the volumetric proving system. Read and record all pertinent volumetric standard temperatures and pressures during the calibration sequence. Place the lid on the transfer vessel to limit evaporation.
- 2.4.10.6 Immediately prior to weighing full vessel record balance reading as (d_1) . This reading will be used to calculate any balance drift that occurred while the transfer vessel was being filled.
- 2.4.10.7 Place the filled vessel and all padding material on the balance. If needed, add known mass standards as tare weights to bring the water mass up to the mass of the standards used for O_1 . This balance reading is recorded as O_2 . Record the mass of all tare weights.
- 2.4.10.8 Remove the filled transfer vessel and record the balance indication as (d_2) . This reading is used to calculate balance drift over the entire calibration process.

- 2.4.10.9 Immediately after removing the filled transfer vessel from the balance, record the temperature of water in the vessel.
- 2.4.10.10 Make at least five replicate runs with at least two runs at a flow rate differing from the others by at least 25 % when the system allows for varying flow rates. The range measured result (V_{60}) of each of these calibration runs must agree within 0.02 % of the volume (i.e., the maximum result minus the minimum result must be less than 0.02 % of the volume being measured).
- 2.4.10.11 Calculate the volume using Equations 2 and 3 in Section 3.1.1., (Option B).

3 Calculations

- 3.1 Compute the volume at the temperature of the measurement, V_t , for each determination using the equation:
 - 3.1.1 Option A.

$$V_{t} = \left[\frac{\mathcal{O}_{4}}{\mathcal{O}_{3}} M_{s2} \left(1 - \frac{\rho_{a}}{\rho_{s2}} \right) - \frac{\mathcal{O}_{2}}{\mathcal{O}_{1}} M_{s1} \left(1 - \frac{\rho_{a}}{\rho_{s1}} \right) \right] \left(\frac{1}{\rho_{w} - \rho_{a}} \right)$$
Eqn. 1

Table 2. Variables for volume equation.

Variable	Description
M_s , M_{s1} , M_{s2}	mass of standards M_{s1} , is the mass of the standards used with the empty transfer vessel M_{s2} , is the mass of the standards used with the transfer vessel after filling with delivered water
$ ho_{\scriptscriptstyle S}$	density of M_S standards
$ ho_{\scriptscriptstyle W}$	density of water at the temperature of measurement
$ ho_a$	density of air at the conditions of calibration – may be different for empty and filled weighing of the transfer vessel
V_t	Volume at the temperature of the test for the unknown under test

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3.1.1 Option B.

$$V_{t} = \left\{ \left[O_{2c} \left(\frac{M_{s}}{O_{1}} \right) \left(1 - \frac{\rho_{a}}{\rho_{s}} \right) \right] - M_{t} \left(1 - \frac{\rho_{a}}{\rho_{t}} \right) \right\} \left(\frac{1}{\rho_{w} - \rho_{a}} \right) \text{ Eqn. 2}$$

$$O_{2c} = O_{2} + \left\{ \left[\frac{\left(d_{1} + d_{2} \right)}{2} \right] - B \right\} \text{ Eqn. 3}$$

Table 3. Variables for volume equations.

Variable	Description
O_1	Observation 1, balance reading for mass standard
O_2	Observation 2, balance reading for water delivered from prover
O_{2c}	Drift-compensated O_2 reading
d_1	Drift while filling the vessel
d_2	Drift over entire process
В	Base balance reading drift is based on
M_S	Mass of mass standards (true mass, vacuum mass)
M_t	Mass of tare weights
$ ho_a$	Air density
$ ho_s$	Density of M_S
$ ho_t$	Density of M_t
$ ho_{\scriptscriptstyle w}$	Water density (calculated using GLP 10 equations)

3.2 Compute V_{60} , the volume at 60 °F, for each run, using the equations noted below:

$$V_{60} = \frac{V_t}{CCF}$$
 where
$$CPL_p = \frac{1}{\left[1 - \left((0.0000032 * P_p)\right)\right]}$$

$$CTS_{p1} = \left\{\left(1 + \left[\left(T_p - T_b\right) * G_a\right]\right) * \left(1 + \left[\left(T_d - T_b\right) * G_t\right]\right)\right\}$$

$$CTS_{p2} = \left\{\left(1 + \left[\left(T_p - T_b\right) * G_c\right]\right)\right\}$$

$$CPS_p = 1 + \left(\frac{\left(P_p * ID\right)}{E * WT}\right)$$

$$CCF = CPL_p * CTS_p * CPS_p$$

Table 4. Additional Variables for Volume Equations.

Variable	Description
T_b	Reference (Base) temperature
T_d	Detector bar temperature (Temperature, detector) – where present
T_p	Prover temperature (Temperature, prover)
P_p	Prover pressure (Pressure, prover) in psig
E	Modulus of elasticity (flow tube)
WT	Wall Thickness of flow tube
ID	Inside Diameter of flow tube
G_l	Coefficient of linear thermal expansion (switch bar) – where present (either linear and area values are used in combination or the cubical value is used)
G_a	Coefficient of flow tube area thermal expansion (either linear and area values are used in combination or the cubical value is used)
G_c	Coefficient of cubical thermal expansion (either linear and area values are used in combination or the cubical value is used)
CPL	Correction for the pressure on the liquid
CTS	Correction for the temperature on the standard prover CTS _{p1} , based on linear and area coefficients of expansion or CTS _{p2} , based on cubical coefficients of expansion
CPS	Correction for the pressure on the standard prover
CCF	Combined correction factor

- 3.3 Calculate water density for air-saturated water, using the equations provided in GLP 10.
- 3.4 Calculate the air density per NISTIR 6969, Selected Mass Calibration Procedures, SOP 2, Option B.
- 3.5 Calculate the uncertainty for the calibration using Section 5.

4 Measurement Assurance

- 4.1 Record and plot standard deviations of each five run calibration to determine short-term repeatability of the measurement process.
- 4.2 Perform applicable calibrations of check standards where available.

5 Assignment of Uncertainties

The limits of expanded uncertainty, U, include estimates of the standard uncertainty of the mass standards used, u_s , plus the uncertainty of measurement, s_p , at the 95 % level of confidence. See SOP 29 for the complete standard operating procedure for calculating the uncertainty.

- 5.1 The standard uncertainty for the mass standards, u_s , is obtained from the calibration reports for the standards used. The combined standard uncertainty, u_c , is used and not the expanded uncertainty, U, therefore the reported uncertainty for the standard will usually need to be divided by the coverage factor k.
- 5.2 The standard deviation of the measurement process is obtained from replicate measurements and standard deviation chart performance data (See SOP 17 or 20.)

The value for s_p is obtained from the standard deviation chart data and results for the current run must be compared to the standard deviations observed from replicate measurements over time. The standard deviation of the process, determined from this approach incorporates a repeatability factor related to the precision of all weighings and need not be duplicated. An F-test may be incorporated to compare observed standard deviations from the current calibration with the accepted standard deviations of the process for similar type proving systems used as standards.

- 5.3 Other standard uncertainties usually included for this type of calibration level include uncertainties associated with water temperature measurements, thermometer accuracy, calculation of air density, and standard uncertainties associated with the density of the standards used or the lack of internal cleanliness.
- An example uncertainty budget table is shown in Table 4. Additional components may need to be considered depending on the technology being calibrated.

Table 4. Example uncertainty budget table.

Uncertainty Component Description	Symbol	Source	Typical Distribution	
Mass Standards (filled; empty)	u_s	Calibration Report	Normal, 1s	
Mass Density	$u_{ ho s}$	NIST	Rectangular	
Standard deviation of the process	s_p	Control chart, standard deviation chart	Normal, 1s	
Prover Temperature	u_{tp}	Ability to measure and possibility of gradients	Rectangular	
Water Temperature	u_{tw}	Ability to measure and possibility of gradients	Rectangular	
Water Density Equation	$u_{\rho w}$	GLP 10, CIPM Reference	Rectangular	
Water Compressibility	u_K	Kell, CIPM Reference	Rectangular	
Cubical Coefficient of Thermal Expansion	u_{Gc}	5 % to 10 % (EURAMET CG-19)	Rectangular	
Area Coefficient of Thermal		5 % to 10 %		
Expansion	u_{Ga}	(EURAMET CG-19)	Rectangular	
Linear Coefficient of Thermal Expansion	u_{Gl}	5 % to 10 % (EURAMET CG-19)	Rectangular	

Modulus of Elasticity (E)	u_E	Estimate 1 %	Rectangular
Prover Pressure	u_{psig}	Calibration report	Rectangular
		Estimate by one division in last	
Flow Tube Diameter (ID)	u_{ID}	decimal place	Rectangular
		Estimate by one division in last	
Tube Wall Thick (WT)	u_{WT}	decimal place	Rectangular
		NISTIR 6969, SOP 2, CIPM	
Air Temperature	u_{ta}	Reference, Calibration report	Rectangular
		NISTIR 6969, SOP 2, CIPM	
Barometric pressure (absolute)	u_p	Reference, Calibration report	Rectangular
		NISTIR 6969, SOP 2, CIPM	
Relative Humidity	u_{RH}	Reference, Calibration report	Rectangular
		NISTIR 6969, SOP 2, CIPM	
Air Density Equation	$u_{\rho a \ eqn}$	Reference, Calibration report	Rectangular
Viscosity	u_V	NIST SP 250-72	Rectangular

6 Report

- 6.1 Report results as described in SOP 1, Preparation of Calibration/Test Results, with the addition of the following:
 - 6.1.1 Volume, reference temperature, uncertainty, material, thermal coefficient of expansion (assumed or measured), construction, any identifying markings, tolerances (if appropriate), laboratory temperature, water temperature(s) at time of test, barometric pressure, relative humidity, and any out—oftolerance conditions. Volume may need to be reported as "up-stream" and "down-stream" volumes for bi-directional provers.
 - 6.1.2 The status of items that are not calibrated by the laboratory but that may be used to perform measurements must be noted on the calibration report (e.g., temperature sensors, pressure gauges, or full flow rates that may be used in the field but that are not calibrated by the laboratory.) A statement such as the following must be included on calibration reports: "Only items presented on this calibration report were tested under the conditions shown". Any sensors or gauges that were not calibrated by the laboratory should be clearly identified on the report (e.g., "Calibration reports for the system included temperature sensors and pressure gauges that were provided to the laboratory for evaluation prior to calibration and were found to be acceptable.")
 - 6.1.3 The volumetric proving system being calibrated as a standard should be evaluated for conformance to applicable specifications and tolerances if being used for legal weights and measures applications; if no specific standards apply, or a laboratory does not perform an assessment, a notice should be included on the calibration report to the effect that a legal metrology assessment was not performed. (For example, "Conformity assessment was not completed as a part of this calibration. Accurate use in field testing requires assessment of repeatability in field applications as well as proper training of the operator, maintenance of the proving system, integrity of all seals and sensors, and any other auxiliary mechanical or electronic components. The entire system has not been assessed for suitability as a field standard for weights and measures applications.")

Additional References

Chapter 4, Manual of Petroleum Measurement Standards, and the following Sections as applicable:

Section 1 – Introduction

Section 2 – Conventional Pipe Provers

Section 3 – Small Volume Provers

Section 4 – Tank Provers

Section 5 – Master-Meter Provers

Section 6 – Pulse Interpolation

Section 7 – Field Standard Test Measures

Section 8 – Operation of Proving Systems

NIST Handbook 105-7: Specifications and Tolerances for Reference Standards and Field Standard Weights and Measures, 7. Specifications and Tolerances for Dynamic Small Volume Provers, 1996.

Bean, V. E., Espina, P. I., Wright, J. D., Houser, J. F., Sheckels, S. D., and Johnson, A. N., NIST Calibration Services for Liquid Volume, NIST Special Publication 250-72, National Institute of Standards and Technology, Gaithersburg, MD, (2009).

EURAMET Calibration Guide 19, Guidelines on the Determination of Uncertainty in Gravimetric Volume Calibration, (Version 2.1, 03/2012).

Appendix A – Gravimetric Calibration Data Sheet

Company Name:			Test No:	
Address:			Purchase Order No:	
City, State, Zip:			_ Date Scheduled:	
Representative:			_ Date Received:	
Phone:	Fax:		Date Tested:	
Email:			_ Date Returned:	
Company URL:			_ Next Appointment:	
General Description			_	
Model Number:	Serial Numb	oer:	_	
Condition:			_	
Compressibility factor for water	3.2000E-06	Cubical The	rmal Expansion Coefficient (G_c)	
Reference Temperature(Tb) (°F)	5.20002 00	_	Il Expansion Coefficient (G_a)	
Reference Temperature (°C)		_	ermal Expansion Coefficient (G_l))
Reference Pressure (psig)			Elasticity (flow tube) (E)	
* 0		Flow Tube In	nside Diameter (inches) (<i>ID</i>)	
		Flow Tube V	Vall Thickness (inches) (WT)	-
Mass standard(s) data:		-		

ID (Note ID and whether for Filled or Empty Transfer Vessel)	Nominal Mass	Mass Correction*	Expanded Unc: From cal. report	Unc: k factor	Density g/cm ³
S					
S					
S					
S					
S					
S					
S					
S					

^{*}Mass Correction = *Mass* values are required for this procedure (i.e., Conventional Mass must *not* be used).

Observations (Option A):

Fill Number	Fill # 1	Fill # 2	Fill #3	Fill #4	Fill # 5
Flow Rate	Fast	Fast	Fast	Slow	Slow
Run Time (sec)					
Balance Reading O ₁ (Standards for Empty Transfer Vessel)					
Balance Reading O ₂ (Empty Transfer Vessel)					
Balance Reading O ₃ (Standards for Filled Transfer Vessel)					
Balance Reading O ₄ (Filled Transfer Vessel)					
Air T emperature (°C)					
Barometric Pressure (mm Hg)					
Relative Humidity (% RH)					
Water Temperature (°C) – record all values					
Detector Bar (where present) Temperature (T _d) (°C)					
Prover T emperature (T_p) (°C)					
Prover pressure (P_p) (psig)					

Observations (Option B):

Fill Number	Fill # 1	Fill # 2	Fill #3	Fill #4	Fill # 5
Flow Rate	Fast	Fast	Fast	Slow	Slow
Run Time (sec)					
Balance Reading O ₁ (standards)					
Balance Reading Empty Transfer Vessel T ared & Removed (B))					
Balance Reading Immediately Before O_2 Reading (d_1)					
Balance Reading O ₂ (water)					
Balance Reading Full Transfer Vessel Removed (d ₂)					
Validity Check (Compares d_1 -B, and d_2 - d_1)					
Drift Compensated O2 (O _{2c})					
Air Temperature (°C)					
Barometric Pressure (mm Hg)					
Relative Humidity (% RH)					
Water Temperature (°C) – record all values					
Water Temperature (°F)					
Detector Bar (where present) T emperature (Td) (°C)					
Detector Bar (where present) Temperature (Td) (°F)					
Prover T emperature (Tp) (°C)					
Prover Temperature (Tp) (°F)					
Prover pressure (Pp) (psig)					