

GC2 Calibration Verification / Linearity Test Kit

INTENDED USE

VALIDATE GC 2 Calibration Verification / Linearity Test Kit solutions are intended for *in vitro* diagnostic use in the quantitative determination of linearity, calibration verification and verification of reportable range in automated, semi-automated and manual instrument systems for the following analytes: ammonia (NH3), carbon dioxide (CO2), ethyl alcohol (ETOH) and iron (FE).

Each test kit consists of one bottle each of Levels 1 through 5. Each bottle contains 3.0 milliliters. There exists a linear relationship among Levels 1 through 5.

SUMMARY

For each VALIDATE Calibration Verification / Linearity Test Kit, multiple levels are provided to establish the relationship between theoretical and actual performance of each of the included analytes. The VALIDATE Calibration Verification / Linearity Test Kit will assist in the documentation of linearity, calibration verification and verification of linear range required by many inspection agencies. The solutions will also provide assistance when troubleshooting instrument systems, reagent problems and calibration anomalies.

REAGENTS

Reactive Ingredients:

Purified chemicals for CO2, ETOH, FE and NH3 in an aqueous base.

Nonreactive Ingredients:

Preservatives and stabilizers.

Precautions and Warnings:

For In Vitro Diagnostic Use

Disposal of all waste material should be in accordance with local guidelines.

STORAGE AND STABILITY

VALIDATE GC 2 Calibration Verification / Linearity Test Kits are stored at 2° to 8°C. **DO NOT FREEZE**. Test kits are stable until the expiration date printed on the bottle and storage container when handled according to instructions.

PREPARATION

Prior to use, remove the **VALIDATE** GC2 Calibration Verification / Linearity Test Kit from storage and allow to come to room temperature (18° to 25°C). Invert gently several times before dispensing.

To maximize stability, it is recommended that exposure to room temperature be minimized. Tightly cap opened bottles and return to 2° to 8°C immediately after dispensing.

Discard any solutions that appear to have gross bacterial contamination.

VALIDATE Calibration Verification / Linearity Test Kits should be treated in the same manner as patient samples. If dilutions or pre-treatment are required as part of the testing procedure, follow the manufacturer's instructions.

ASSAY

Analyze each level in replicates. If following the CLSI EP6 guidelines for linearity, use a random analytical sequence to assay each level.

CALCULATION OF RESULTS

VALIDATE Calibration Verification / Linearity material is prepared in a manner such that an equal distance (delta) exists between Levels 1 through 5. This dilution scheme is consistent with the CLSI EP6 recommendation for preparing linearity sets.

Two examples for calculating the theoretical values of Levels 1 through 5 are provided below.

Example 1:

Choose two consecutive levels and calculate the delta between the recovered values. The following example demonstrates the use of the delta between Levels 2 and 3 to calculate the theoretical value for Levels 1, 4, and 5.

Level 3 – Level 2 = Delta

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Level 1 Theoretical = Level 2 Recovered – Delta
Level 4 Theoretical = Level 3 Recovered + Delta
Level 5 Theoretical = Level 4 Theoretical + Delta
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NOTE: The user can select the calculated delta between any two consecutive levels to calculate the theoretical values. Typically, the user should choose an area of recovery known to be linear for the method being studied.

Example 2:

Theoretical values can be determined using the recovered values for Levels 1 and 5. Using this method, the following formulas apply:

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Level 2 Theoretical = 0.75 * (Level 1) + 0.25 * (Level 5)
Level 3 Theoretical = 0.5 * (Level 1) + 0.5 * (Level 5)
Level 4 Theoretical = 0.25 * (Level 1) + 0.75 * (Level 5)
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After theoretical values are calculated, for each analyte plot the expected (Theoretical) value on the x-axis versus the Recovered value on the y-axis using standard linear graph paper. If the system is linear, the plot should approximate a straight line. The point at which the line is no longer straight can be used to determine the limit of linearity or the reportable range.

Data reduction is available from LGC Maine Standards. Commercially available linear regression software may also be used. The software should provide data point display and x-y graphical presentation. Linear regression should be interpreted using standard statistical analysis and the results should be compared with the instrument manufacturer's claims for linearity or with individual laboratory performance requirements. The degree of acceptable nonlinearity is an individual judgment based on methodology, clinical significance and medical decision levels of the test analyte.

LIMITATIONS

VALIDATE Calibration Verification / Linearity Test Kit solutions are not intended for use as routine quality control materials or as calibration materials.

EXPECTED VALUES

VALIDATE Calibration Verification / Linearity Test Kits are manufactured such that an equal distance (delta) exists between levels as recommended by CLSI EP6 for assessing linearity. As the distance between levels is equal, any two levels can be held to be 'true' when assayed and the theoretical values for each of the other three levels can be calculated allowing this test kit to be used on multiple automated instrument systems.

The reagent manufacturer's recommended diluent can be used to make dilutions of the low level to obtain a result lower than that level, if needed.

The following three (3) analytes are inverted in GC2: CO2, ETOH, and NH3. Level 1 contains the highest concentration for these analytes and concentration decreases from Level 1 down to Level 5.

TYPICAL VALUES

Actual results obtained may vary depending on instrumentation, methodology and assay temperature. Results may also be dependent on the accuracy of the instrument / reagent system calibration. The degree of acceptable nonlinearity is an individual judgment based on methodology, clinical significance and medical decision levels of the test analyte.

Typical recovered values for Level 1 and Level 5 are presented in the table(s) provided. Typical values for mid-levels are based on an equal distance (delta) between levels.

Typical Recovered Values on Ortho Vitros®										
1200vt Lot #: 12BP012210										
Analyte	Units	Level 1	Level 2	Level 3	Level 4	Level 5				
CO2	mmol/L	40	32	24	15	7				
FE	μg/dL	10	152	294	435	577				
ETOH	mg/dL	290	222	154	85	17				
NH3	μmol/L	465	351	238	124	10				

1200vt Lot #: 12BP012210										
Analyte	SI Units	Level 1	Level 2	Level 3	Level 4	Level 5				
CO2	mmol/L	40	32	24	15	7				
FE	µmol/L	1.790	27.163	52.537	77.910	103.283				
ETOH	mmol/L	62.930	48.120	33.310	18.499	3.689				
NH3	µmol/L	465	351	238	124	10				





C€ Symbols – This product fulfills the requirements of the European Directive 98/79/EC for in vitro medical devices. The following symbols may be used where applicable in labeling for Maine Standards Company products:

LOT

Lot Number

Expiration Date



Manufacturer



Storage Temperature



In Vitro Diagnostic Medical Device



Catalog Number



Insert



Biological Risk



EC REP Wellkang Ltd (www.CE-marking.eu)
29 Harley St., London W1G 9QR, UK



For a list of countries in which VALIDATE $^{\! \otimes}$ is registered see:

http://www.mainestandards.com/Products/ce_reg.php

Rx Only

ORDERING INFORMATION

ORDER NO.: 1200vt

VALIDATE GC2

Calibration Verification / Linearity Test Kit:

5 x 3 mL

CONTACT INFORMATION:

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