

Method Validation Worksheets

Refer to QA Manual for instructions on using the Method Validation Worksheets. Also, visit the Qualigen, Inc. website at www.qualigeninc.com for “On Q” Training.

1. Verify Accuracy and Precision – this is performed once per analyzer per assay during installation and training.

2. Verify Reportable Ranges (Calibration Verification) – this is performed initially at start up and routinely every 6 months thereafter.

3. Identify your Reference Ranges

PSA - Verify Accuracy and Precision. Run C1 and C2 five times each upon installation.

C1 (Low Control Range)

C2 (High Control Range)

1

Control C1

Write "C1", the control kit lot number, and your initials on the FastPack® IP peel-off label and place it here.

Control C2

Write "C2", the control kit lot number, and your initials on the FastPack® IP peel-off label and place it here.

2

Control C1

Write "C1", the control kit lot number, and your initials on the FastPack® IP peel-off label and place it here.

Control C2

Write "C2", the control kit lot number, and your initials on the FastPack® IP peel-off label and place it here.

3

Control C1

Write "C1", the control kit lot number, and your initials on the FastPack® IP peel-off label and place it here.

Control C2

Write "C2", the control kit lot number, and your initials on the FastPack® IP peel-off label and place it here.

4

Control C1

Write "C1", the control kit lot number, and your initials on the FastPack® IP peel-off label and place it here.

Control C2

Write "C2", the control kit lot number, and your initials on the FastPack® IP peel-off label and place it here.

5

Control C1

Write "C1", the control kit lot number, and your initials on the FastPack® IP peel-off label and place it here.

Control C2

Write "C2", the control kit lot number, and your initials on the FastPack® IP peel-off label and place it here.

Calculating Accuracy and Precision

From the Control Range Card, enter the Upper and Lower limits as well as the Mean of the control material's acceptable range into the appropriate sections below.

PSA - C1

Lower Limit

Mean

Upper Limit

Calculate the Average of all 5 values (1+2+...+5)/5:

AVG = _____

Calculate the Standard Deviation of all 5 values:

SD = _____

Calculate the Install Precision Specification:

IPS = _____

$$IPS = 0.8 \times SD \text{ Spec} \quad \left\{ \begin{array}{l} \text{If AVG} < 0.2, SD \text{ Spec} = 0.0224 \\ \text{If AVG} \geq 0.2, SD \text{ Spec} = 0.112 \times \text{AVG} \end{array} \right\}$$

Calculate the Chi-Squared Statistic¹ (χ^2_{stat}):

$\chi^2_{\text{stat}} =$ _____

$$\chi^2_{\text{stat}} = SD^2 \times \left(\frac{2.5}{IPS^2} \right)$$

PSA - C2

Lower Limit

Mean

Upper Limit

Calculate the average of all 5 values (1+2+...+5)/5:

AVG = _____

Calculate the standard deviation (SD) of all 5 values:

SD = _____

Calculate the Install Precision Specification:

IPS = _____

$$IPS = 0.8 \times SD \text{ Spec} \quad \left\{ \begin{array}{l} \text{If AVG} < 0.2, SD \text{ Spec} = 0.0224 \\ \text{If AVG} \geq 0.2, SD \text{ Spec} = 0.112 \times \text{AVG} \end{array} \right\}$$

Calculate the Chi-Squared Statistic (χ^2_{stat}):

$\chi^2_{\text{stat}} =$ _____

$$\chi^2_{\text{stat}} = SD^2 \times \left(\frac{2.5}{IPS^2} \right)$$

¹ Chi-Squared Statistic (χ^2) statistical analysis is an accepted methodology for precision performance evaluations. Refer to CLSI (Clinical and Laboratory Standards Institute) EP5 approved guideline "Evaluation of Precision Performance of Quantitative Measurement Procedures".

PSA

Accuracy and Precision

Analyzer SN:

Circle Your Response

1	Do all control testing values fall within the acceptable QC range?	Y	N	N/A
2	Is the C1 control $\chi^2_{\text{stat}} < \chi^2_{\text{crit}}?$	Y	N	N/A
3	Is the C2 control $\chi^2_{\text{stat}} < \chi^2_{\text{crit}}?$	Y	N	N/A

The critical value for the Chi-Squared Statistic (χ^2_{crit}) based on 5 measurements and a 95% confidence level is 7.81.

If you can provide a "Yes" answer in all 3 question categories above, check the box below to accept the manufacturer's claims for accuracy and precision. If the answer in any of the above question categories is No, check the box that you DO NOT accept the manufacturer's claims for accuracy and precision and contact Qualigen System Support.

- Accept the manufacturer's claims for accuracy and precision
- DO NOT accept the manufacturer's claims for accuracy and precision

PSA

Verify Reportable Ranges (Calibration Verification) every 6 months

Verify that the FastPack® IP System is accurate to the limits of the reportable range specified by Qualigen, Inc. by using the FastPack® IP Total PSA Method Verification Kit.

<p style="text-align: center;">Low</p> <p>Target <input type="text"/></p>	<p style="text-align: center;"><i>Write "Low Verifier", the lot number, and your initials on the peel-off FastPack® label and place it here</i></p>
<p style="text-align: center;">Mid</p> <p>Range <input type="text"/></p>	<p style="text-align: center;"><i>Write "Mid Verifier", the lot number, and your initials on the peel-off FastPack® label and place it here</i></p>
<p style="text-align: center;">High</p> <p>Target <input type="text"/></p>	<p style="text-align: center;"><i>Write "High Verifier", the lot number, and your initials on the peel-off FastPack® label and place it here</i></p>

The Verifier results must be within the following ranges to accept the Manufacturer's Reportable Range:

	Low Verifier	High Verifier
Hybritech Standard	< 0.04 ng/mL	> 50 ng/mL
WHO International Standard	< 0.04 ng/mL	> 40 ng/mL

If either result is different than the above, record the reportable range based on actual observed values. The mid value must be within the range designated on the range card. If it is not, repeat the test. If any value is out of range after repeating the test, contact Qualigen System Support.

- Accept Manufacturer Reportable Range
- DO NOT Accept. Derived Reportable Range: _____

Identify the Reference Ranges for your Practice

Qualigen has performed extensive clinical trials to determine the reference ranges for normal, healthy male and female subjects as well as for those with malignant and non-malignant urological diseases. This information can be found in the FastPack® IP Total PSA Immunoassay direction insert. Using this information as a guide (normal healthy males had PSA levels up to 4 ng/mL), indicate the normal ranges used in this practice which will reflect the expected ranges appropriate for your patient population.

Type of Patient	Your Reference (Normal) Ranges
Normal	
Others:	

- OK to begin testing NOT OK to begin testing

Testing Analyst/Technical Consultant

Date

Laboratory Director

Date