AccuSpan™ CMV DNA Linearity Panel 2410-0174 / Batch # 10468133



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OVERVIEW

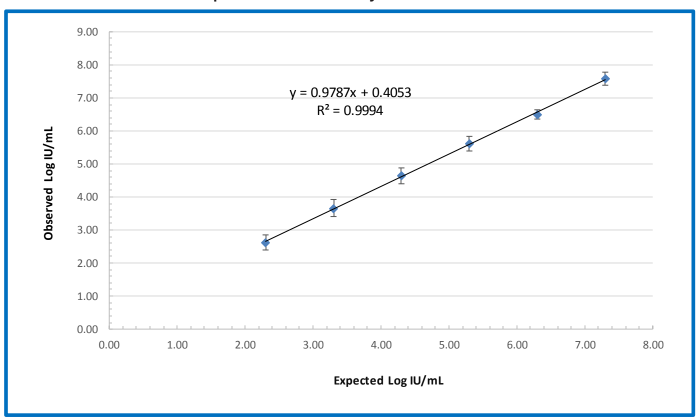
The AccuSpan™ CMV DNA Linearity Panel 2410-0174 / Batch number 10468133 is a nine-member panel consisting of seven members representing serial log dilutions of cultured CMV virus, with established reactivity for CMV (Cytomegalovirus) DNA, in CMV DNA negative diluent. This panel also consists of one negative member prepared from the diluent and one member of diluent to perform additional dilutions as desired. The diluent was prepared from normal human plasma that was 0.2 micron filtered. Sodium azide (0.09%) was added as a preservative.

Roche COBAS® AmpliPrep/COBAS® TaqMan® results are reported for each panel member. Linearity is shown graphically by plotting observed results against expected results. The WHO International Standard was tested in the same run as the AccuSpan™ CMV DNA Linearity Panel members. Both expected and observed results for the standards are reported; expected results for the WHO standards are the WHO assigned values.

For Research Use Only. Not for use in diagnostic procedures. Data are provided for informational purposes. SeraCare Life Sciences does not claim that others can duplicate test results exactly.

CAUTION: These materials have not been treated and should be considered biohazardous. Follow Universal Precautions.

AccuSpan CMV DNA Linearity Panel Members 1 – 6



CMV DNA results were obtained using the Roche COBAS® AmpliPrep/COBAS® TagMan® CMV test method. A line of best fit is shown.

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AccuSpan CMV DNA Linearity Panel

Roche COBAS®
AmpliPrep/COBAS®
TaqMan® CMV Test ¹

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Panel Member	log IU/mL	log copies/mL	
012	7.58	7.62	
02	6.50	6.54	
03	5.61	5.66	
04	4.64	4.68	
05	3.67	3.71	
06	2.62	2.67	
073	< 2.14	< 2.35	
08	TND	TND	
09	TND	TND	
Test Date	26	6 NOV 2019	
Test Site		RL	
Test Kit Range	137 to	9,100,000 IU/mL	
Test Kit Conversion Factor	1 copy/mL = 0.91 IU	J/mL, 1 IU/mL = 1.1 copies/mL	
Test Kit Part Code	04	4902025190	
Test Kit Lot No.	F17530		
Test Kit Expiration	31 JAN 2021		

¹Results are reported as the mean result of testing in six (6) replicates. Results in red are considered positive.

TND = Target Not Detected; RL = Reference Lab

²Panel Member 01 was tested at a 1:10 dilution and results were corrected for the dilution factor.

³CMV Detected below limit of quantitation.

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WHO International Standard 1st CMV DNA International Standard (NIBSC code: 09/162)

	Observed Values on Roche COBAS® AmpliPrep/COBAS®			
	Expected Values	TaqMan® CMV Test		
Sample ID	(log IU/mL)	(log IU/mL) ¹	% Difference ²	
Sample B1	5.70	5.84	2.44	
Sample B2	5.00	5.22	4.40	
Sample B3	4.70	4.81	2.42	
Sample B4	4.00	4.20	5.10	
Sample B5	3.70	3.91	5.78	
Test Date	26 NOV 2019			
Test Site	RL			
Test Kit Range	137 to 9,100,000 IU/mL			
Test Kit Conversion Factor	1 copy/mL = 0.91 IU/mL, 1 IU/mL = 1.1 copies/mL			
Kit Part Code	04902025190			
Kit Lot No.	F17530			
Test Kit Expiration	31 JAN 2021			

¹WHO panel was tested in triplicate in the same test run as the AccuSpan™ CMV DNA Linearity Panel members. Results in bold red are considered positive.

RL = Reference Lab

The package insert for this panel can be found at www.seracare.com.

A printed copy of the package insert or data sheet may be requested by email at info@seracare.com or by phone at 508.244.6400.

²Percentage difference is how much the observed concentration differs from the expected concentration. It is calculated as: $\frac{observed - Expected}{Expected}$ Values calculated for reference only.