# QCMD 2021 B19 virus DNA EQA Programme



Catalogue Code: QAV034116

Ref Code: B19DNA21 Challenge:

C2

Analysis Type:

Qualitative and Quantitative

**Dataset:** 460864

Report UID: 2677/460864/3563

Laboratory CZ023

### **Intended Results / Panel Composition**

Sample Code	Sample Content		Sample Status	Consensus (IU/ml) <sup>[4]</sup>		Range [5]		
						(Log <sub>10</sub> )	(n)	
B19DNA21C2-01	B19 Virus Type 1	Plasma	DS1_1	Frequently Detected	CORE	4.173	36	3.313 - 4.710
B19DNA21C2-02	B19 Virus Type 1	Plasma	DS1_2	Frequently Detected	CORE	3.246	36	2.569 - 4.072
B19DNA21C2-03	B19 Virus Negative	Plasma		Negative	CORE	N/A	N/A	N/A
B19DNA21C2-04	B19 Virus Type 1	Plasma		Detected	EDUCATIONAL	2.167	25	0.845 - 3.999

- [1] **Sample Relationships:** Indicates the relationships of the samples within this challenge. The highest titre member of dilution series DS1 is indicated by DS1\_1 and further members of the series as DS1\_2, DS1\_3 etc. in order of reducing titre. Additional dilution series are indicated by DS2 (e.g DS2\_1, DS3\_2 etc.), DS3 (e.g. DS3\_1, DS3\_2 etc.). If one duplicate pair is present this is indicated by 'D1'. Further duplicate pairs are indicated by 'D2', 'D3' etc.
- [2] **Detection Frequency:** To aid qualitative analysis each panel member is assigned a frequency of detection. This is based on the peer group consensus of all qualitative results returned from participants within the EQA challenge / distribution.
- [3] **Sample Status:** EQA samples are defined as "CORE" or "EDUCATIONAL". Core proficiency samples are reviewed by the QCMD Scientific Expert(s). This is on the basis of scientific information, clinical relevance, current literature and, where appropriate, professional clinical guidelines. Participating laboratories are expected to report core proficiency samples correctly within the EQA challenge / distribution.
- [4] **Consensus (IU/mI):** Mean consensus (Log<sub>10</sub>) calculated from data returned by participants with outliers removed and number of quantitative results (n) returned for each panel member.
- [5] Range: Maximum and minimum quantitative value (IU/ml) reported by participants within this challenge without outliers removed.

For further details please refer to the current participant manual.

### **Your Summary Results**

Units	IU/ml
EQA Assessment Group [1]	Commercial
Core Panel Detection (Qualitative) Score [2]	0
Core Panel Estimation (Quantitative) Score [3]	1

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C2

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#### **Core Panel Members Results**

Sample Code	Unitage	EQA Assessment Group Consensus [4]	SD [5]	Quantitative Result		Qualitative Result			
		Group consensus		Your Result	Estimation Score <sup>[7]</sup>	Percentage Correct (All)	Your Result	Detection Score [10]	
B19DNA21C2-01	IU/ml	4.135	0.236	4.059	0	100.0	Positive	0	
B19DNA21C2-02	IU/ml	3.234	0.319	2.914	1	96.7	Positive	0	
B19DNA21C2-03	IU/ml	N/A	-	LOD/NR	N/A	100.0	Negative	0	

All quantitative values above expressed in Log 10 IU/ml.

- [1] **EQA Assessment Group:** To aid data analysis, participant results are grouped according to the molecular amplification/detection method specified within their molecular workflow for this challenge / distribution. For further details refer to the *Additional Information: Individual Panel Member Analysis* section of this report.
- [2] Core Panel Detection (Qualitative) Score: An overall core panel detection score provided per challenge / distribution.
- [3] Core Panel Estimation (Quantitative) Score: An overall core panel estimation score provided per challenge / distribution.
- [4] EQA Assessment Group Consensus: The mean value for all results within your EQA assessment group.
- [5] SD: The standard deviation for results from your EQA assessment group.
- [6] Your Quantitative Result: The quantitative result you returned for each sample within this EQA challenge. LOD/NR (limit of detection or not reported).
- [7] **Estimation Score:** Your estimation (quantitative) scores are calculated based on your variation from the consensus for your EQA assessment group. With 0 (zero) scored if the quantitative value you reported is within one standard deviation (SD) from your EQA assessment group consensus, 1 (one) if your quantitative value is between one and two SDs, 2 (two) if your quantitative value is within two and three SDs and 3 (three) if your quantitative value is more than three SDs from the mean of your EQA assessment group.
- [8] Percentage Correct (AII): Percentage of datasets (%) reporting the correct qualitative results for each panel member.
- [9] Your Qualitative Result: The qualitative result you reported for each sample within this EQA challenge / distribution.
- [10] **Detection Score:** Your detection (qualitative) scores are based on the assigned detection frequency of each panel members, where 0 (zero) is "highly satisfactory" and 3 (three) is "highly unsatisfactory". Scores are provided for individual panel members.

For further details please refer to the current participant manual.

# QCMD 2021 B19 virus DNA EQA Programme



Catalogue Code: QAV034116

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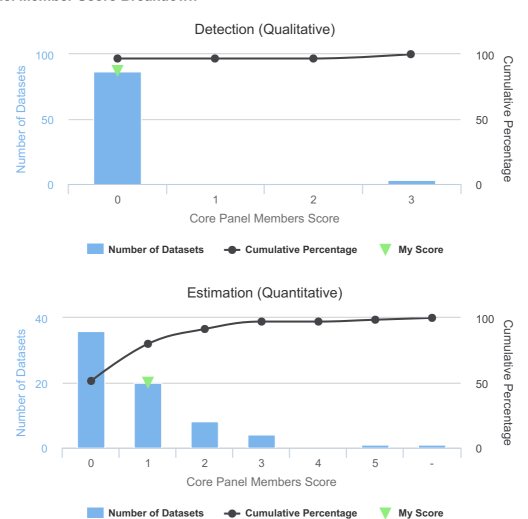
**Analysis Type:**Qualitative and Quantitative

Dataset: 460864

Report UID: 2677/460864/3563

Laboratory CZ023

### **Core Panel Member Score Breakdown**



Core Panel Member Score Breakdown - Detection: This figure gives you a breakdown of the qualitative detection scores for all qualitative datasets returned within this EQA challenge / distribution independent of the EQA assessment group. Panel detection scores are generated from only those panel members that are defined as "CORE".

Core Panel Member Score Breakdown - Estimation: This figure gives you a breakdown of the quantitative estimation scores for all quantitative datasets returned within this EQA challenge / distribution independent of the EQA assessment group. Panel estimation scores are based on positive core panel members only. Those datasets that did not return quantitative values for all core samples are represented by '-'.

For further details please refer to the current participant manual.

# QCMD 2021 B19 virus DNA EQA Programme



Catalogue Code: QAV034116

Ref Code: B19DNA21 Challenge:

C2

Analysis Type:

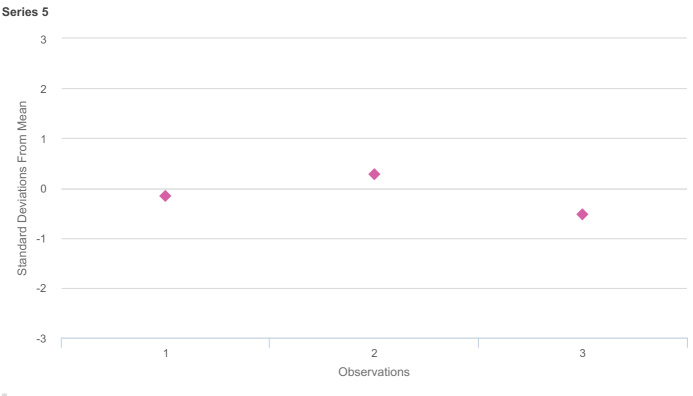
Qualitative and Quantitative

**Dataset:** 460864

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Laboratory CZ023

### **Duplicate Sample Performance Over Time**



### **Observation Details**

- 1: B19DNA101C2-03 B19DNA20C1-04: Your Variation 0.005 (IU/ml): Overall Mean Variation 0.076: Overall SD 0.453.
- 2: B19DNA21C1-03 B19DNA101C2-03: Your Variation -0.026 (IU/ml): Overall Mean Variation -0.096: Overall SD 0.250. 3: B19DNA21C2-02 - B19DNA21C1-03: Your Variation -0.169 (IU/ml): Overall Mean Variation 0.008: Overall SD 0.337.
- QCMD monitors your laboratory's performance over time based on the reported quantitative variation between duplicate panel members within the EQA

The mean variation and standard deviation are calculated from the quantitative variation reported by each participant between duplicate panel members in the same unit of measurement once outliers have been removed. (See 'Observation Details')

Previous and current observations are plotted on the chart as the number of standard deviations your variation was from the mean variation for all participants who submitted corresponding results in the same unitage.

Any reported variation greater than ±3 SD will not be shown on the graph, but your variation value will be provided in red in the *Observation Details*. When "N/A" is displayed for an observation, either no valid quantitative results were provided or there was a change in reported unitage.

challenge and, where appropriate, across EQA challenges.

# QCMD 2021 B19 virus DNA EQA Programme



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**Analysis Type:**Qualitative and Quantitative

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Laboratory CZ023

### My Workflow Details

The details of the workflow(s) used to submit your results for this challenge.

Name	GeneProof Parvovirus B19 PCR Kit (v4)
Description	
Targets	V B19 virus
Assays	<ul> <li>Extraction - Manual Extraction Process</li> <li>Commercial         <ul> <li>Kit Manufacturer: GeneProof</li> <li>Kit Type: PathogenFree DNA Isolation Kit</li> </ul> </li> <li>Amplification - GeneProof - croBEE Real-Time PCR System</li> <li>Commercial         <ul> <li>Kit Manufacturer: GeneProof</li> <li>Kit Type: Parvovirus B19 PCR Kit</li> <li>Kit Version: ISEX</li> </ul> </li> </ul>

### **Educational Panel Members Results**

Sample Code	e Code Unitage EQA Assessment Group Consensus [1]		<b>SD</b> [2]	Quantitative Result		Qualitative Result			
				Your Result	Estimation Score <sup>[4]</sup>	Percentage Correct (All)	Your Result	Detection Score <sup>[7]</sup>	
B19DNA21C2-04	IU/ml	2.009	0.570	1.342	1	87.8	Positive	0	

All quantitative values above expressed in Log 10 IU/ml.

- [1] EQA Assessment Group Consensus: The mean value for all results within your EQA assessment group.
- [2] SD: The standard deviation for results from your EQA assessment group.
- [3] Your Quantitative Result: The quantitative result you returned for each sample within this EQA challenge. LOD/NR (limit of detection or not reported).
- [4] **Estimation Score:** Your estimation (quantitative) scores are calculated based on your variation from the consensus for your EQA assessment group. With 0 (zero) scored if the quantitative value you reported is within one standard deviation (SD) from your EQA assessment group consensus, 1 (one) if your quantitative value is between one and two SDs, 2 (two) if your quantitative value is within two and three SDs and 3 (three) if your quantitative value is more than three SDs from the mean of your EQA assessment group.
- [5] Percentage Correct (All): Percentage of datasets (%) reporting the correct qualitative results for each panel member.
- [6] Your Qualitative Result: The qualitative result you reported for each sample within this EQA challenge / distribution.
- [7] **Detection Score:** Your detection (qualitative) scores are based on the assigned detection frequency of each panel members, where 0 (zero) is "highly satisfactory" and 3 (three) is "highly unsatisfactory". Scores are provided for individual panel members.

For further details please refer to the current participant manual.

# QCMD 2021 B19 virus DNA EQA Programme



Catalogue Code: QAV034116

Ref Code: B19DNA21 Challenge: C2 Analysis Type:
Qualitative and Quantitative

**Dataset:** 460864

**Report UID:** 2677/460864/3563

Laboratory CZ023

### **Further Programme Details**

Number of Participants	84
Number of Countries	25
Number of Respondents	79
Number of Datasets Submitted	96
Quantitative Results Returned (All)	70 (72.9%)
- Quantitative Results Returned (IU/ml)	36 (51.4%)
- Quantitative Results Returned (Copies/ml)	33 (47.1%)
- Quantitative Results Returned (Copies/5ul)	1 (1.4%)
Qualitative Results Returned	90 (93.8%)

### **EQA Programme Aims**

To assess the proficiency of laboratories in detection and quantitation of B19 virus.

### Feedback and Enquiries

Participants are encouraged to read the QCMD Participants' Manual, which can be downloaded from the QCMD website.

Any enquiries should be submitted through the 'Contact Us' form that you can find in the 'Help' section of your QCMD (ITEMS) Participant Profile Area.

## QCMD 2021 B19 virus DNA EQA **Programme**



**Catalogue Code:** QAV034116

Ref Code: B19DNA21 Challenge:

C2

Analysis Type: Qualitative and Quantitative

Dataset: 460864

Report UID: 2677/460864/3563 Laboratory CZ023

Panel member analysis is separated into CORE samples followed by EDUCATIONAL samples.

### **Additional Core Samples Information**

The following section has been categorised as shown below:

Core ► Quantitative ► IU/ml, Copies/ml ► Qualitative

### **Individual Panel Member Analysis (Quantitative)**

Quantitative analysis for each panel member is provided in relation to your EQA assessment group. EQA assessment groups are established using the molecular workflow information reported by all participants within this EQA challenge / distribution. The principal level of assessment is at the individual method level which is defined based on your reported "amplification/detection method" and other laboratories using the same or similar amplification/detection methods.

To allow meaningful assessment at the individual method level the EQA assessment group must consist of 5 or more datasets. If there are not sufficient datasets at the individual method level then your results will be included within a higher EQA assessment group based on whether it is a commercial or in house technology/method. The highest level assessment grouping is all reported results using the same unit of measurement (i.e. Copies/ml or IU/ml).

The results below provide a breakdown of participant reported values on each of the panel members within this EQA challenge / distribution. Your result for each panel member is indicated by "your value". You can compare your value to the "mean" within your EQA assessment group and the overall consensus for each sample within this EQA challenge / distribution.

Key



# QCMD 2021 B19 virus DNA EQA Programme



Catalogue Code: QAV034116

Ref Code: B19DNA21 Challenge: C2 Analysis Type:
Qualitative and Quantitative

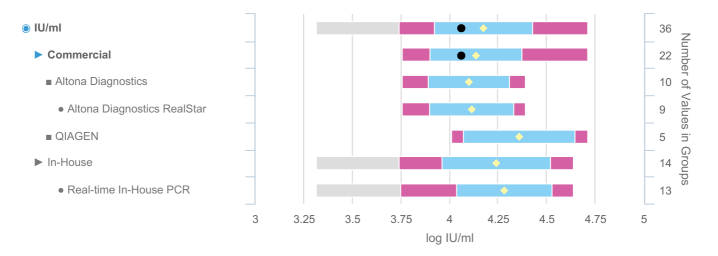
**Dataset:** 460864

Report UID: 2677/460864/3563

Laboratory CZ023

### B19DNA21C2-01 - Quantitative Results Breakdown (IU/ml)

Sample Code	Sample Content	Matrix	Sample Relationships	Detection Frequency	Sample Status	Consensus (IU/ml)		Range
						(Log <sub>10</sub> )	(n)	
B19DNA21C2-01	B19 Virus Type 1	Plasma	DS1_1	Frequently Detected	CORE	4.173	36	3.313 - 4.710



Groups below n=5: AB Analitica (n=2), AB Analitica - AB Analitica REALQUALITY RQ (n=2), Altona Diagnostics - Altona Diagnostics AltoStar (n=1), ELITech Group (n=3), ELITech Group - Elitech Elite Real Time kit (n=3), GeneProof (n=2), GeneProof - GeneProof Real Time PCR kit (n=2), In-House - Conventional In-House PCR (n=1)

Groups Rolled Up: QIAGEN - Qiagen Artus Real Time (n=5)

# QCMD 2021 B19 virus DNA EQA Programme



Catalogue Code: QAV034116

Ref Code: B19DNA21

Challenge: C2 Analysis Type:
Qualitative and Quantitative

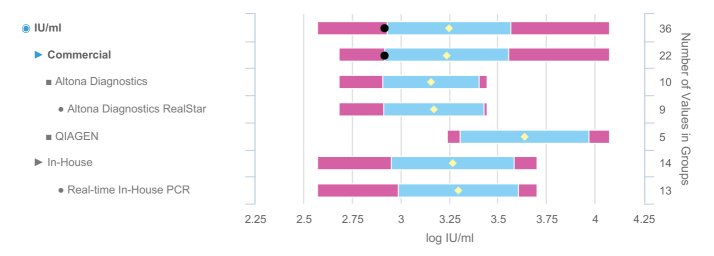
**Dataset:** 460864

Report UID: 2677/460864/3563

Laboratory CZ023

### B19DNA21C2-02 - Quantitative Results Breakdown (IU/ml)

Sample Code	Sample Content	Matrix	Sample Relationships	Detection Frequency	Sample Status	Consensus (IU/ml)		Range
						(Log <sub>10</sub> )	(n)	
B19DNA21C2-02	B19 Virus Type 1	Plasma	DS1_2	Frequently Detected	CORE	3.246	36	2.569 - 4.072



Groups below n=5: AB Analitica (n=2), AB Analitica - AB Analitica REALQUALITY RQ (n=2), Altona Diagnostics - Altona Diagnostics AltoStar (n=1), ELITech Group (n=3), ELITech Group - Elitech Elite Real Time kit (n=3), GeneProof (n=2), GeneProof - GeneProof Real Time PCR kit (n=2), In-House - Conventional In-House PCR (n=1)

Groups Rolled Up: QIAGEN - Qiagen Artus Real Time (n=5)

# QCMD 2021 B19 virus DNA EQA Programme



Catalogue Code: QAV034116

Ref Code: B19DNA21 Challenge: C2 Analysis Type:
Qualitative and Quantitative

**Dataset:** 460864

**Report UID:** 2677/460864/3563

Laboratory CZ023

### B19DNA21C2-01 - Quantitative Results Breakdown (Copies/ml)

Sample Code	Sample Content	Matrix	Sample Relationships	Detection Frequency	Sample Status	Consensus (Copies/ml)		Range
						(Log <sub>10</sub> )	(n)	
B19DNA21C2-01	B19 Virus Type 1	Plasma	DS1_1	Frequently Detected	CORE	4.324	33	3.591 - 5.173

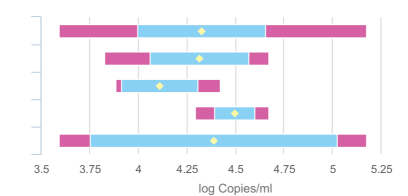


► Commercial

■ ELITech Group

■ bioMerieux

▶ In-House





**Groups below n=5:** AB Analitica (n=3), AB Analitica - AB Analitica REALQUALITY RQ (n=3), Altona Diagnostics (n=3), Altona Diagnostics - Altona Diagnostics RealStar (n=3), PrimerDesign (n=1), PrimerDesign - PrimerDesign Genesig (n=1), QIAGEN (n=1), QIAGEN - Qiagen Artus Real Time (n=1)

**Groups Rolled Up:** ELITech Group - Elitech Elite Real Time kit (n=6), bioMerieux - bioMerieux R-gene Kit (n=14), In-House - Real-time In-House PCR (n=5)

# QCMD 2021 B19 virus DNA EQA Programme



Catalogue Code: QAV034116

Ref Code: B19DNA21

Challenge:

Analysis Type:
Qualitative and Quantitative

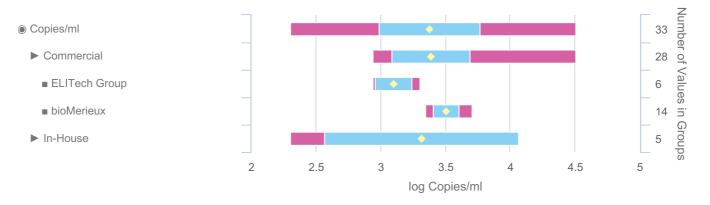
**Dataset:** 460864

Report UID: 2677/460864/3563

Laboratory CZ023

### B19DNA21C2-02 - Quantitative Results Breakdown (Copies/ml)

Sample Code	Sample Content	Matrix	Sample Relationships	Detection Frequency	Sample Status	Consensus (Copies/ml)		Range
						(Log <sub>10</sub> )	(n)	
B19DNA21C2-02	B19 Virus Type 1	Plasma	DS1_2	Frequently Detected	CORE	3.374	33	2.301 - 4.504



**Groups below n=5:** AB Analitica (n=3), AB Analitica - AB Analitica REALQUALITY RQ (n=3), Altona Diagnostics (n=3), Altona Diagnostics - Altona Diagnostics RealStar (n=3), PrimerDesign (n=1), PrimerDesign Genesig (n=1), QIAGEN (n=1), QIAGEN - Qiagen Artus Real Time (n=1)

**Groups Rolled Up:** ELITech Group - Elitech Elite Real Time kit (n=6), bioMerieux - bioMerieux R-gene Kit (n=14), In-House - Real-time In-House PCR (n=5)

### **Individual Panel Member Analysis (Qualitative)**

Qualitative analysis for each panel member is provided in relation to your EQA assessment group. EQA assessment groups are established using the molecular workflow information reported by all participants within this EQA challenge / distribution. The principal level of assessment is at the individual method level which is defined based on your reported "amplification/detection method" and other laboratories using the same or similar amplification/detection methods.

To allow meaningful assessment at the individual method level the EQA assessment group must consist of 5 or more datasets. If there are not sufficient datasets at the individual method level then your results will be included within a higher EQA assessment group based on whether it is a commercial or in house technology/method. The highest level assessment grouping is "All" participant reported qualitative results.

A breakdown of qualitative results reported by participants on each of the panel members within this EQA challenge / distribution is provided below. You can compare your results to those within your EQA assessment group and those obtained within other EQA assessment groups or to the overall consensus for each sample within this EQA challenge / distribution.

# QCMD 2021 B19 virus DNA EQA Programme



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Ref Code: B19DNA21 Challenge:

C2

Analysis Type:
Qualitative and Quantitative

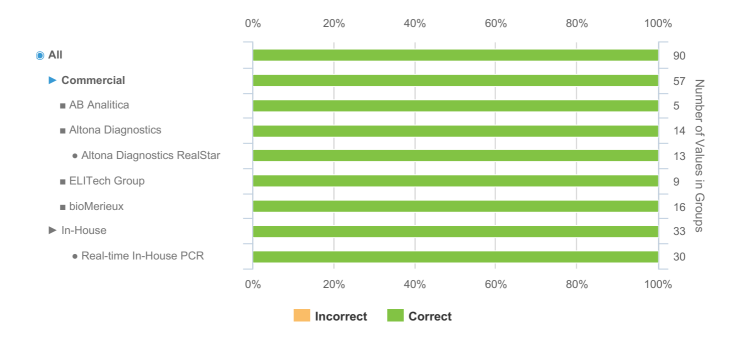
**Dataset:** 460864

**Report UID:** 2677/460864/3563

**Laboratory** CZ023

### B19DNA21C2-01 - Qualitative Results Breakdown

Sample Code	Sample Content	Matrix	Sample Relationships	Detection Frequency	Sample Status	Percentage (	Correct
						(%)	(n)
B19DNA21C2-01	B19 Virus Type 1	Plasma	DS1_1	Frequently Detected	CORE	100.0	90



Groups below n=5: Altona Diagnostics - Altona Diagnostics AltoStar (n=1), Anatolia Geneworks (n=2), Anatolia Geneworks - Anatolia Geneworks Bosphore (n=2), GFE Blut (n=1), GFE Blut - GFE Blut VSPK (n=1), GeneProof (n=2), GeneProof - GeneProof Real Time PCR kit (n=2), Grifols (n=1), Grifols - Grifols Procleix (n=1), PrimerDesign (n=2), PrimerDesign Genesig (n=2), Progenie Molecular (n=1), Progenie Molecular - Progenie Molecular RealCycler (n=1), QIAGEN (n=4), QIAGEN - Qiagen Artus Real Time (n=4), In-House - Conventional In-House PCR (n=3)

**Groups Rolled Up:** AB Analitica - AB Analitica REALQUALITY RQ (n=5), ELITech Group - Elitech Elite Real Time kit (n=9), bioMerieux - bioMerieux Regene Kit (n=16)

# QCMD 2021 B19 virus DNA EQA Programme



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Ref Code: B19DNA21

Challenge: C2 Analysis Type:
Qualitative and Quantitative

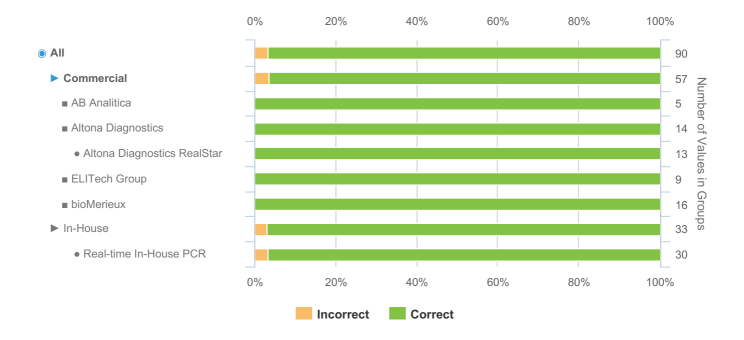
**Dataset:** 460864

**Report UID:** 2677/460864/3563

Laboratory CZ023

### B19DNA21C2-02 - Qualitative Results Breakdown

Sample Code	Sample Content	Matrix	Sample Relationships	Detection Frequency	Sample Status	Percentage (All)	Correct
						(%)	(n)
B19DNA21C2-02	B19 Virus Type 1	Plasma	DS1_2	Frequently Detected	CORE	96.7	90



Groups below n=5: Altona Diagnostics - Altona Diagnostics AltoStar (n=1), Anatolia Geneworks (n=2), Anatolia Geneworks - Anatolia Geneworks Bosphore (n=2), GFE Blut (n=1), GFE Blut - GFE Blut VSPK (n=1), GeneProof (n=2), GeneProof - GeneProof Real Time PCR kit (n=2), Grifols (n=1), Grifols - Grifols Procleix (n=1), PrimerDesign (n=2), PrimerDesign Genesig (n=2), Progenie Molecular (n=1), Progenie Molecular - Progenie Molecular RealCycler (n=1), QIAGEN (n=4), QIAGEN - Qiagen Artus Real Time (n=4), In-House - Conventional In-House PCR (n=3)

**Groups Rolled Up:** AB Analitica - AB Analitica REALQUALITY RQ (n=5), ELITech Group - Elitech Elite Real Time kit (n=9), bioMerieux - bioMerieux Regene Kit (n=16)

# QCMD 2021 B19 virus DNA EQA Programme



Catalogue Code: QAV034116

Ref Code: B19DNA21 Challenge: C2 Analysis Type:
Qualitative and Quantitative

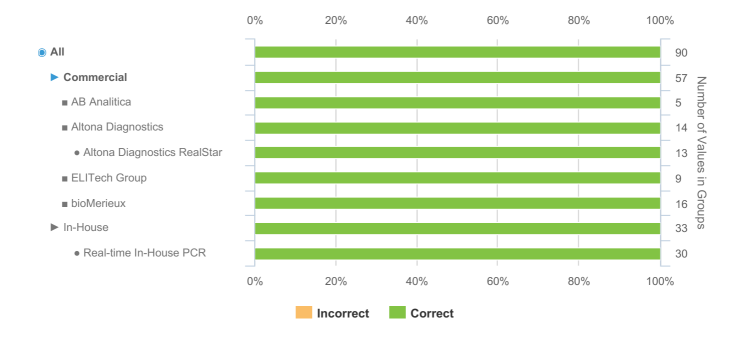
**Dataset:** 460864

**Report UID:** 2677/460864/3563

**Laboratory** CZ023

### B19DNA21C2-03 - Qualitative Results Breakdown

	Sample Code	Sample Content	Matrix	Sample Relationships	Detection Frequency	Sample Status	Percentage Correct (All)	
							(%)	(n)
	B19DNA21C2-03	B19 Virus Negative	Plasma		Negative	CORE	100.0	90



Groups below n=5: Altona Diagnostics - Altona Diagnostics AltoStar (n=1), Anatolia Geneworks (n=2), Anatolia Geneworks - Anatolia Geneworks Bosphore (n=2), GFE Blut (n=1), GFE Blut - GFE Blut VSPK (n=1), GeneProof (n=2), GeneProof - GeneProof Real Time PCR kit (n=2), Grifols (n=1), Grifols - Grifols Procleix (n=1), PrimerDesign (n=2), PrimerDesign Genesig (n=2), Progenie Molecular (n=1), Progenie Molecular - Progenie Molecular RealCycler (n=1), QIAGEN (n=4), QIAGEN - Qiagen Artus Real Time (n=4), In-House - Conventional In-House PCR (n=3)

Groups Rolled Up: AB Analitica - AB Analitica REALQUALITY RQ (n=5), ELITech Group - Elitech Elite Real Time kit (n=9), bioMerieux - bioMerieux Rene Kit (n=16)

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C2

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Qualitative and Quantitative

**Dataset:** 460864

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Laboratory CZ023

### **Additional Educational Samples Information**

The following section has been categorised as shown below:

Educational ► Quantitative ► IU/ml, Copies/ml ► Qualitative

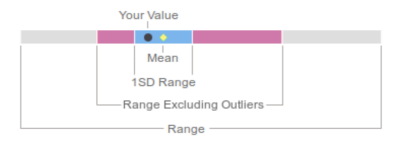
### **Individual Panel Member Analysis (Quantitative)**

Quantitative analysis for each panel member is provided in relation to your EQA assessment group. EQA assessment groups are established using the molecular workflow information reported by all participants within this EQA challenge / distribution. The principal level of assessment is at the individual method level which is defined based on your reported "amplification/detection method" and other laboratories using the same or similar amplification/detection methods.

To allow meaningful assessment at the individual method level the EQA assessment group must consist of 5 or more datasets. If there are not sufficient datasets at the individual method level then your results will be included within a higher EQA assessment group based on whether it is a commercial or in house technology/method. The highest level assessment grouping is all reported results using the same unit of measurement (i.e. Copies/ml or IU/ml).

The results below provide a breakdown of participant reported values on each of the panel members within this EQA challenge / distribution. Your result for each panel member is indicated by "your value". You can compare your value to the "mean" within your EQA assessment group and the overall consensus for each sample within this EQA challenge / distribution.

Key



# QCMD 2021 B19 virus DNA EQA Programme



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Ref Code: B19DNA21 Challenge: C2 Analysis Type:
Qualitative and Quantitative

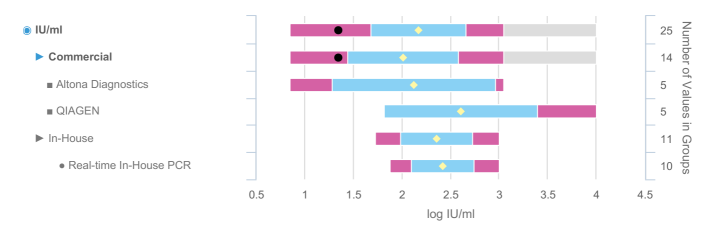
**Dataset:** 460864

Report UID: 2677/460864/3563

Laboratory CZ023

### B19DNA21C2-04 - Quantitative Results Breakdown (IU/ml)





Groups below n=5: AB Analitica (n=2), AB Analitica - AB Analitica REALQUALITY RQ (n=2), Altona Diagnostics - Altona Diagnostics AltoStar (n=1), Altona Diagnostics - Altona Diagnostics RealStar (n=4), GeneProof (n=2), GeneProof - GeneProof Real Time PCR kit (n=2), In-House - Conventional In-House PCR (n=1)

Groups Rolled Up: QIAGEN - Qiagen Artus Real Time (n=5)

### B19DNA21C2-04 - Quantitative Results Breakdown (Copies/ml)

Sample Code	Sample Content	Matrix	Sample Relationships	Detection Frequency	Sample Status	Consensu (Copies/m		Range
						(Log <sub>10</sub> )	(n)	
B19DNA21C2-04	B19 Virus Type 1	Plasma		Detected	EDUCATIONAL	2.412	26	1.041 - 3.158



► Commercial

■ bioMerieux





# QCMD 2021 B19 virus DNA EQA Programme



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Ref Code: B19DNA21 Challenge:

C2

**Analysis Type:** 

**Dataset:** 460864

Report UID: 2677/460864/3563

Laboratory CZ023

Groups below n=5: AB Analitica (n=2), AB Analitica - AB Analitica REALQUALITY RQ (n=2), Altona Diagnostics (n=2), Altona Diagnostics - Altona Diagnostics RealStar (n=2), ELITech Group (n=2), ELITech Group - Elitech Elite Real Time kit (n=2), PrimerDesign (n=1), PrimerDesign - PrimerDesign Genesig (n=1), QIAGEN (n=1), QIAGEN - Qiagen Artus Real Time (n=1), In-House (n=4), In-House - Real-time In-House PCR (n=4)

Qualitative and Quantitative

Groups Rolled Up: bioMerieux - bioMerieux R-gene Kit (n=14)

### **Individual Panel Member Analysis (Qualitative)**

Qualitative analysis for each panel member is provided in relation to your EQA assessment group. EQA assessment groups are established using the molecular workflow information reported by all participants within this EQA challenge / distribution. The principal level of assessment is at the individual method level which is defined based on your reported "amplification/detection method" and other laboratories using the same or similar amplification/detection methods.

To allow meaningful assessment at the individual method level the EQA assessment group must consist of 5 or more datasets. If there are not sufficient datasets at the individual method level then your results will be included within a higher EQA assessment group based on whether it is a commercial or in house technology/method. The highest level assessment grouping is "All" participant reported qualitative results.

A breakdown of qualitative results reported by participants on each of the panel members within this EQA challenge / distribution is provided below. You can compare your results to those within your EQA assessment group and those obtained within other EQA assessment groups or to the overall consensus for each sample within this EQA challenge / distribution.

# QCMD 2021 B19 virus DNA EQA Programme



Catalogue Code: QAV034116

Ref Code: B19DNA21 Challenge:

C2

Analysis Type:
Qualitative and Quantitative

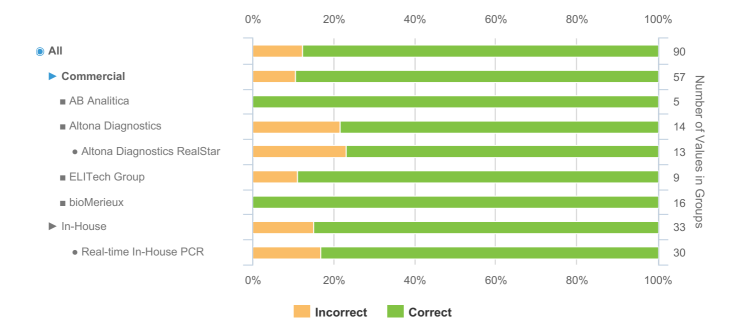
**Dataset:** 460864

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#### B19DNA21C2-04 - Qualitative Results Breakdown

Sample Code	Sample Content	Matrix	Sample Relationships	Detection Frequency	Sample Status	Percentage Correct (All)	
						(%)	(n)
B19DNA21C2-04	B19 Virus Type 1	Plasma		Detected	EDUCATIONAL	87.8	90



Groups below n=5: Altona Diagnostics - Altona Diagnostics AltoStar (n=1), Anatolia Geneworks (n=2), Anatolia Geneworks - Anatolia Geneworks Bosphore (n=2), GFE Blut (n=1), GFE Blut - GFE Blut VSPK (n=1), GeneProof (n=2), GeneProof - GeneProof Real Time PCR kit (n=2), Grifols (n=1), Grifols - Grifols Procleix (n=1), PrimerDesign (n=2), PrimerDesign Genesig (n=2), Progenie Molecular (n=1), Progenie Molecular - Progenie Molecular RealCycler (n=1), QIAGEN (n=4), QIAGEN - Qiagen Artus Real Time (n=4), In-House - Conventional In-House PCR (n=3)

**Groups Rolled Up:** AB Analitica - AB Analitica REALQUALITY RQ (n=5), ELITech Group - Elitech Elite Real Time kit (n=9), bioMerieux - bioMerieux Rene Kit (n=16)

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