

1 **Evaluation of the Luminex ARIES[®] System for the Detection and Quantification of BK**
2 **virus DNA in Plasma and Serum Samples from Kidney Transplant Recipients**

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6
7
8 **Abstract**

9
10 BKV nephropathy is as serious complication in renal transplant recipients due to the need for
11 immunosuppression. Nearly 50% of renal transplant patients with BKV nephropathy experience
12 a significant loss of function of the transplanted kidney. It is routine practice to screen renal
13 transplant recipients regularly for BK viremia. In this study we compared the performance of
14 BKV qPCR analyte specific reagents (ASR) by ELITech, and Luminex for measuring BKV viral
15 load in plasma and serum using the Roche Cobas[®] z480 instrument, and the Luminex ARIES[®]
16 platform respectively. A total of 34 patients previously tested on the z480, with results spanning
17 the test 's linear range, were analyzed on the ARIES[®]. The BKV DNA copy number correlation
18 between the two methods was very good with an R² value of 0.98. The average difference in log
19 copy number between the two methods was -0.3, indicating that the ARIES[®] method may have
20 slightly greater analytical sensitivity. BKV quantification results were closely matched between
21 the two different methods. The workflow with the ARIES[®] System is greatly simplified by
22 elimination of DNA extraction and most hands-on steps. The high degree of automation allows
23 samples to be tested as they arrive into the laboratory, resulting in enhanced patient care due to
24 more rapid turnaround time for results back to the ordering physician.

25

26

27 **Introduction**

28 BK virus (BKV), a member of the human polyomaviruses family, was first discovered in the
29 urine of renal transplant patients with postoperative ureteral stenosis. (6) BKV is ubiquitous; in
30 adults in the United States and Europe, seroprevalence ranges from 60% to 80%.

31 Epidemiological studies indicate that BKV is contracted during childhood; between 3 to 4 years
32 of age (11). Very little is known about the transmission of this virus or about events during
33 primary infection. The majority of primary infections are asymptomatic; however, BKV may
34 cause mild upper respiratory disease in young children (13). A latent infection is established in
35 the kidneys of the infected host by BKV.

36

37 BKV is recognized as a significant cause of nephropathy in renal transplant patients and is a
38 significant cause of graft failure in 1% to 10% of patients (7, 14). Approximately 40% of renal
39 allograft recipients shed BKV in the urine, either transiently or continuously over weeks to
40 months (1, 2). Nephropathy caused by BKV is asymptomatic and associated with increased
41 serum creatinine levels. Nearly 50% of renal transplant patients diagnosed with BKV
42 nephropathy experience a significant loss of function of the transplanted kidney.

43 BKV viruria has also been demonstrated in approximately 10 – 25% of bone marrow transplant
44 (BMT) patients, typically two months post-transplantation (2). The frequency of cystitis is
45 higher in adult allogeneic compared to autologous BMT recipients. BKV can be demonstrated in
46 the peripheral blood of both groups (5). Because detection of BKV in the urine of BMT patients
47 in the absence of disease is common, it is difficult to correlate BKV as the cause of cystitis in
48 these individuals (5).

49

50 Surveillance testing for BKV in the blood and urine of renal transplant recipients has become
51 routine. In 2013, an expert panel recommended the use of either urine cytology or nucleic acid–
52 based testing to screen renal transplant recipients monthly during the first 6 months post-
53 transplantation, and then every 3 months until 2 years after transplantation (14). These
54 guidelines included quantitative cutoffs for BKV loads that indicate additional testing. Urine
55 DNA loads of $>10^7$ copies/mL or plasma DNA loads of $>10^4$ copies/mL that persist for more
56 than three weeks constitute a diagnosis of “presumptive polyomavirus-associated nephropathy”
57 and should be followed up with a renal biopsy (14).

58
59 There are no antiviral agents for the treatment of BKV. Treatment of kidney transplant
60 recipients with BKV infection consists of lowering the dosage of immunosuppressive drugs used
61 to prevent rejection of the transplanted organ to allow the patient’s immune response to clear the
62 infection (3, 10, 12 16). For this reason, rapid turnaround time for quantitative BKV
63 determination is important to initiate such treatment, and then to resume proper
64 immunosuppressive therapy once the infection is cleared.

65
66 There are no FDA approved assays commercially available for BKV, so molecular testing is
67 performed using laboratory developed tests; reagents for BKV nucleic acid amplification
68 (NAAT) tests are available as either ASRs or RUOs from several commercial sources. In this
69 paper we describe a rapid, sample-to-answer quantitative BKV quantitative polymerase chain
70 reaction (qPCR) assay based on Luminex MultiCode[®] PCR technology on the ARIES[®] platform
71 (Luminex Corp. Austin TX), which allows for samples to be analyzed as they are received in the
72 laboratory as opposed to batch testing.

73 **Materials and Methods**

74 Clinical Samples

75 All clinical samples used in this study were obtained from kidney transplant recipients as part of
76 routine screening for BKV viremia, submitted to the St. John Hospital and Medical Center
77 Specialized Testing Laboratory for testing. All samples were de-identified prior to testing in this
78 study eliminating the requirement for patient informed consent.

79

80 Specimen processing

81 Plasma or serum samples from kidney transplant recipients were submitted to the St. John
82 Hospital and Medical Center Specialized Testing Laboratory for BK qPCR virus testing. DNA
83 was extracted from 1 mL of each sample using a bioMerieux NucliSENS[®] easyMag[®] nucleic
84 acid extractor (bioMerieux, Durham, NC). The extracted nucleic acids from each sample were
85 eluted into 50 µL of extraction buffer.

86

87 BKV TaqMan qPCR

88 BKV qPCR testing based on TaqMan technology was performed using ELITech (ELITech
89 Group, Logan, UT) 20X MGB Alert[®] BK Virus Primers and 20X MGB Alert[®] BK Virus Probe
90 analyte specific reagents (ASR). The 25 µL reaction consisted of 1X of the primers, probe,
91 MGB Alert[®] Hot Start Master and MGB Alert[®] BK Virus Internal Control primers and probes
92 and 5 uL of the extracted nucleic acid solution. Analysis was performed using the Roche Cobas
93 z480 platform with their proprietary user defined software package (Roche Diagnostics,
94 Indianapolis, IN). Thermocycling conditions were 1 cycle each of 50°C for 2 minutes, 93°C for
95 2 minutes, followed by 45 cycles of 93°C for 15 seconds, 56°C for 30 seconds, and 72°C for 30

96 seconds. The standard curve was established using 5 μ L of dilutions of a quantified synthetic
97 BK virus DNA (ATCC, Manassas, VA) tested in triplicate.

98

99 MultiCode[®] PCR Analysis on the ARIES[®] Platform

100 The MultiCode[®] primer mix (Luminex Corp. Austin, TX) was prepared by adding 2 μ L of
101 MultiCode[®] BK virus ASR primer and 2 μ L of Control Primer (CP3) per sample to a 1.5 mL
102 microcentrifuge tube. A volume of 4 μ L of this primer mix was added to a MultiCode[®] Ready
103 Mix tube (Luminex, Austin, TX). The MultiCode[®] Ready Mix tube contains all of reagents
104 required for the MultiCode[®] PCR reaction to occur. The prepared MultiCode[®] Ready Mix tube
105 is then attached to the appropriate position on the ARIES[®] Extraction Cassette. Two hundred
106 microliters (200 μ L) of plasma or serum is added to the sample chamber of the cassette which is
107 placed into the ARIES[®] cassette holder and loaded into the ARIES[®] instrument. Testing was
108 performed according to standard instrument settings supplied by Luminex using their proprietary
109 SYNCT[™] software and the User Defined Protocol (UDP) application. To generate the standard
110 curve, a 200 μ L sample of each of the above BKV DNA dilutions was added to the sample
111 chamber of the ARIES[®] cassette. Each dilution was tested in duplicate.

112 **Results**

113 Generation of the BKV qPCR Standard Curve:

114 The BKV qPCR standard curve on the Cobas z480 and ARIES[®] platforms was generated using
115 dilutions made from quantified synthetic BK virus DNA. The 6-member panel ranged from
116 1×10^7 to 5×10^2 DNA copies/mL. Each dilution was tested in triplicate on the Cobas z480, and
117 in duplicate on the ARIES[®]. The z480 software automatically generates a standard curve that
118 can be saved and imported for subsequent analyses, meaning that a standard curve need not be

119 generated for each run. The same dilution panel was used to generate the BKV DNA standard
120 curve for the ARIES[®] platform. In this case 200 uL of each panel member was added to
121 ARIES[®] Extraction Cassettes containing the MultiCode[®] BK virus primers. The ARIES[®]
122 software was not designed for quantitative PCR analyses; it is necessary to import the standard
123 curve Ct's and patient data from the ARIES[®] printout into Excel for analysis. The results, shown
124 in Figure 2, panels A and B for the Cobas z480 and ARIES[®] platforms respectfully are very
125 similar. Based on the slopes of both standard curves the efficiency of amplification for the
126 ELITech reagents on the z480, and the MultiCode[®] BKV primers on the ARIES[®] platform were
127 determined to be very close at 97.12% and 99.18% respectively (where a slope of 3.22 = 100%
128 efficiency ([https://www.thermofisher.com/us/en/home/brands/thermo-scientific/molecular-
129 biology/molecular-biology-learning-center/molecular-biology-resource-library/thermo-scientific-
130 web-tools/qpcr-efficiency-calculator.html](https://www.thermofisher.com/us/en/home/brands/thermo-scientific/molecular-biology/molecular-biology-learning-center/molecular-biology-resource-library/thermo-scientific-web-tools/qpcr-efficiency-calculator.html))).

131

132 Determination of the Limit of Detection (LoD):

133 The lower limits of detection (LoD) for both methods were determined by testing dilutions made
134 from the quantified BKV DNA used to generate the standard curve. Each dilution was tested 20
135 times. The lowest dilution resulting in a positivity of $\geq 95\%$ was set as the LoD. The LoD for
136 the ELITech/z480 method was determined to be 500 copies/mL of original sample and 250
137 copies/mL for the MultiCode[®]/ARIES[®] procedure (data not shown).

138

139 Reproducibility of BK virus load testing on the ARIES[®] platform:

140 Five previously tested patient samples were used to determine reproducibility of the BK virus
141 qPCR test on the ARIES[®] platform. Three of these were positive and two were undetectable for

142 BKV DNA. The results shown in Table 1 demonstrates a high degree of reproducibility as
143 demonstrated by the very low percent coefficient of variation between each of the replicate
144 analyses.

145

146 Comparison of testing with clinical samples between the ELITech and ARIES[®] qPCR assays:

147 To compare the performance of both assays with actual clinical samples a panel of 34 plasma or
148 serum samples submitted previously for BKV DNA qPCR testing were analyzed using the
149 ELITech reagents on the z480 and the MultiCode[®] BKV primers on the ARIES[®] Platform. The
150 results are shown in Table 1 and Figure 2. Test results for two different methods are considered
151 equivalent if the log difference is ≤ 0.3 log copies/mL. The overall average Δ log difference
152 between the ARIES[®] and z480 BKV quantification tests was determined to be -0.3 logs. This
153 value decreases to -0.24 if the samples with viral loads <500 copies per mL are removed from
154 the analysis since 500 BKV DNA copies/mL is the lower limit of detection for the ELITech/z480
155 methodology. Linear regression analysis of the data in Table 1 (Figure 2 A) yielded a correlation
156 coefficient (R-squared) value of 0.98, indicating very close agreement of the results obtained by
157 both methods. The residual plot in panel B clearly demonstrates the validity of the linear
158 regression plot in panel A.

159 **Discussion**

160 Due to their highly immunocompromised state, kidney transplant recipients are at high risk of
161 loss of their transplanted organ due to reactivated BK virus infection (1, 2, 7, 14). In the
162 absence of antiviral agents that are active against BKV, “treatment” of BK virus infection in this
163 population is limited to reduction of the use of immunosuppressive drugs to allow the patient’s
164 immune system to clear the infection (10). As a result, the patient is placed at greatly increased

165 risk of immunological rejection of the transplanted kidney (4, 9, 15, 17). Therefore, it is
166 important that BKV viral load testing be performed as expeditiously as possible to determine the
167 need for reduction of immunosuppression, with follow-up monitoring of BKV viral load to
168 determine when appropriate levels of immunosuppressive therapy can be resumed (3, 8, 12, 16).
169 Batch testing of plasma samples for BKV qPCR can result in a delay of testing, especially if the
170 volume of testing is not sufficiently high to permit testing daily. Similarly, sending samples to
171 reference laboratories can result in longer than desirable turn-around times to receiving results.
172 In this study we evaluated the use of the Luminex MultiCode[®] PCR primers on the ARIES[®] real-
173 time PCR platform to determine its applicability for performing quantitative PCR analysis,
174 specifically for quantification of BKV DNA in plasma and serum samples. The ARIES[®] BK
175 virus test, utilizing primers based on MultiCode[®] technology, was compared to our standard
176 method with BKV PCR primers and an MGB probe from ELITech. For the latter, analysis was
177 performed on the Roche Cobas z480 real-time PCR platform. Unlike the z480 platform, the
178 ARIES[®] instrument is not designed for qPCR analysis, so it is necessary to import the standard
179 curve Ct's and patient data from the ARIES[®] printout into Excel for analysis. Since it would be
180 neither convenient nor economically prudent to perform a standard curve each for each BK
181 qPCR test run on the ARIES[®], a standard curve was established using serial dilutions made from
182 commercially sourced quantified BKV DNA, the results of which was stored and applied to
183 successive test runs. The standard curves generated using both the ELITech and ARIES[®]
184 methods were nearly identical in terms of both slope and R-squared values (Figure 1). In
185 addition, the slope of the standard curve generated on the ARIES[®] platform of -3.34
186 demonstrates an amplification efficiency of >99% indicating a high degree of robustness of this

187 assay design (19). A new ARIES[®] standard curve was generated when there was a lot change in
188 any of the test reagents (ARIES[®] test cartridges or MultiCode[®] primers).

189 The robustness of this test design was clearly demonstrated by the results of the reproducibility
190 experiment shown in Table 1. The log copy numbers of each of the samples in 3 different test
191 runs resulted in low percent coefficients of variation demonstrating very low variability from run
192 to run.

193
194 Comparison of testing with clinical samples between the ELITech and ARIES[®] qPCR assays
195 demonstrated essentially equivalent results between the two methods. The average difference
196 between log copies per mL generated by both methods for each sample was -0.3. Linear
197 regression analysis of the two sets of results (Figure 2) demonstrated 98% correlation between
198 both methods. The high degree of correlation between results was somewhat surprising
199 considering the large number of differences involved in the mechanics of performing both
200 methodologies. The ELITech method on the z480 platform is entirely manual, including DNA
201 extraction and assembly of the qPCR reactions. The ARIES[®] method is completely automated,
202 including the DNA extraction and reaction set-up processes. Another major difference is the
203 qPCR chemistries used for both methods. The ELITech method utilizes TaqMan PCR
204 technology and measures an increase in fluorescence with each PCR cycle. Due to its unique
205 chemistry, MultiCode[®] technology relies on loss of fluorescent signal per amplification cycle
206 (18).

207 The high degree of correlation between both methods is important since it negates the need to
208 re-baseline our kidney transplant population that have been previously tested with the ELITech
209 methodology when switching to the ARIES[®] BKV qPCR test.

210 In summary, our results clearly demonstrate that BKV quantification results were closely
211 matched between the two different methods described above. The workflow with the ARIES®
212 System is greatly simplified due to elimination of DNA extraction and most hands-on steps. The
213 high degree of automation allows samples to be tested as they arrive in the laboratory, resulting
214 in enhanced patient care due to more rapid turnaround time for results back to the ordering
215 physician.

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297 Figure Legends

298

299 Figure 1: Generation of BKV qPCR standard curves

300 Panel A – standard curve generated with serial dilutions of BKV purified DNA
301 performed in triplicate.

302 Panel B – standard curve generated with serial dilutions of BKV purified DNA
303 performed in duplicate.

304

305 Figure 2: ARIES® /z480 Comparison - Quantification of BK DNA in Clinical Samples

306 Panel A – linear regression analysis of data from Table 2.

307 Panel B – residual plot generated from regression analysis in panel A

308

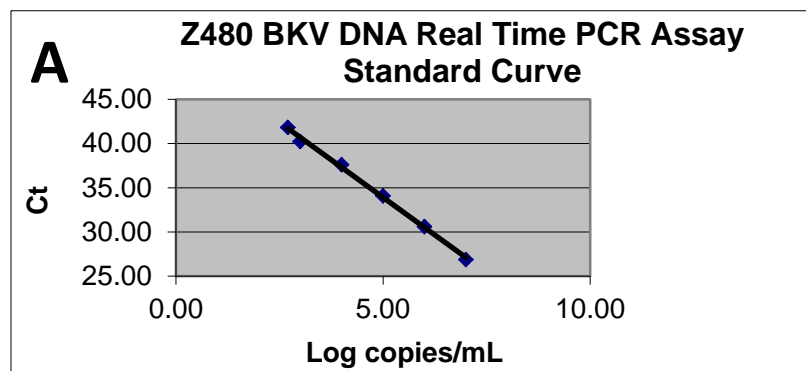
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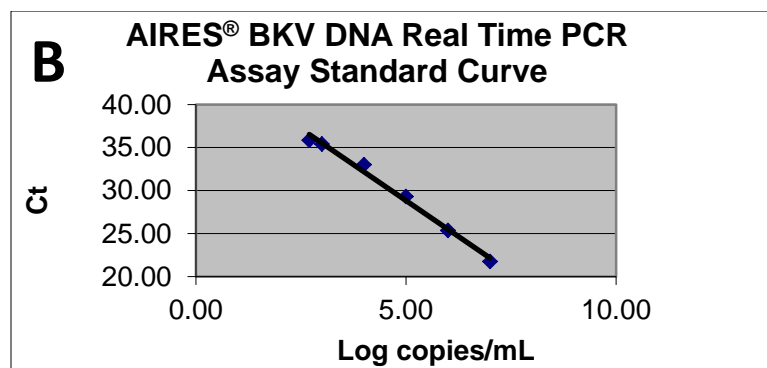
313 **Figure 1: Generation of BKV qPCR standard curves**



314

Slope:	-3.3932
R squared:	0.9975

315



316

Equation	$y = -3.3417x + 45.535$
Slope:	-3.3417
R squared:	0.9905

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320

321

322 **Table1: Reproducibility of BKV qPCR on the ARIES® Platform**

Sample	Replicate 1		Replicate 2		Replicate 3		Avg. log cp/mL	SD	%CV
	Ct	log cp/mL	Ct	log cp/mL	Ct	log cp/mL			
413531	32.2	3.99	31.6	4.17	31.0	4.35	4.17	0.18	4.32
412173	*ND	ND	ND	ND	ND	ND	-	-	-
410089	31.3	4.26	32.0	4.05	31.5	4.20	4.17	0.11	2.59
428857	26.2	5.79	25.8	5.91	25.9	5.88	5.86	0.06	1.07
341582	ND	ND	ND	ND	ND	ND	-	-	-

*ND = Not detected

323

324 **Table 2: Comparison of testing with clinical samples between the ELITech and ARIES®**
 325 **qPCR assays**

Sample	MultiCode®/ARIES		ELITech/z480		Δ log LOG Difference ARIES®/z480
	Copies/mL	Log copies/mL	Copies/mL	Log copies/mL	
313611	39	1.6	787	2.9	-1.3
318856	52	1.72	436	2.64	-0.92
147329	79	1.9	ND	ND	-
416826	111	2.05	473	2.67	-0.62
147329	254	2.4	ND	ND	-
497868	714	2.85	4,410	3.64	-0.79
211229	1,079	3.03	2,427	3.39	-0.36
299629	1,872	3.27	1,309	3.12	0.15
299521	2,149	3.33	9,756	3.99	-0.66
162904	2,831	3.45	8,346	3.92	-0.47
417959	3,729	3.57	5,406	3.73	-0.16
162904	7,428	3.87	8,346	3.92	-0.05
596591	9,785	3.99	44,607	4.65	-0.66
308868	14,795	4.17	25,571	4.41	-0.24
724744	23,966	4.38	27,553	4.44	-0.06
531684	23,966	4.38	52,497	4.72	-0.34
299531	23,966	4.38	54,309	4.73	-0.35
234871	25,675	4.41	52,497	4.72	-0.31
123044	41,590	4.62	71,244	4.85	-0.23
596643	44,557	4.65	94,739	4.98	-0.33
433255	47,735	4.68	93,462	4.97	-0.29
40181	62,884	4.8	33,773	4.53	0.27
240660	125,252	5.1	221,263	5.34	-0.24
355063	286,340	5.46	427,340	5.63	-0.17
597242	654,604	5.82	1,671,599	6.22	-0.4
473995	804,920	5.91	2,237,977	6.35	-0.44
551855	862,338	5.94	1,150,917	6.06	-0.12
40052	923,852	5.97	853,834	5.93	0.04
228172	2,112,027	6.32	787,062	5.9	0.42
149337	40,875,563	7.61	51,103,299	7.71	-0.1
149337	43,791,366	7.64	51,103,299	7.71	-0.07
433174	ND	ND	368	2.57	-
131845	ND	ND	ND	ND	-
293718	ND	ND	ND	ND	-
Average Δ log copies/mL					-0.3

326

327 **Figure 2: ARIES® /z480 Comparison - Quantification of BK DNA in Clinical Samples**

